

A reservoir of foraging decision variables in the mouse brain

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In any given situation, the environment can be parsed in different ways to yield decision variables (DVs) defining strategies useful for different tasks. It is generally presumed that the brain only computes a single DV defining the current behavioral strategy. Here to test this assumption, we recorded neural ensembles in the frontal cortex of mice performing a foraging task admitting multiple DVs. Methods developed to uncover the currently employed DV revealed the use of multiple strategies and occasional switches in strategy within sessions. Optogenetic manipulations showed that the secondary motor cortex (M2) is needed for mice to use the different DVs in the task. Surprisingly, we found that regardless of which DV best explained the current behavior, M2 activity concurrently encoded a full basis set of computations defining a reservoir of DVs appropriate for alternative tasks. This form of neural multiplexing may confer considerable advantages for learning and adaptive behavior.

An adaptive strategy to control behavior is to take actions that lead to good outcomes given that the environment is in a particular state. Yet, environmental states are often complex, with manifold sources of potentially relevant information, some that are directly observable and others that can only be revealed through a process of inference. Therefore, an agent typically also faces the problem of selecting the environmental variables on which to base a decision and how much these variables be processed algorithmically to reveal the appropriate ‘decision variable’ (DV). The problem of selecting a DV is likely a more difficult computational problem faced by a decision maker than the decision itself, but how it is accomplished has received scant investigation¹.

A possibility is that an agent need not commit to a particular DV but may entertain several in parallel. The ability to parallelize operations of decision processing, such as temporal integration, would permit adaptation to changes in task contingencies without implementation of new computations, and could therefore potentially speed learning and provide flexibility in combining and switching strategies. However, little is known about the limitations and possibilities for multiplexing the algorithms used to derive DVs from sensory evidence. On the one hand,

behavioral studies in humans suggested that two streams of sensory evidence can only be incorporated into a DV one at a time, necessitating serial processing^{2–4}. On the other hand, it has been shown that there exist neurons integrating evidence about a single sensory event with diverse timescales⁵, and that diverse timescales are present in neurons within local circuits⁶, which could reflect a simple form of algorithmic multiplexing. It thus remains unclear whether various computations can be carried out in parallel on different streams of evidence to form a broad range of simultaneously available DVs.

To study the possibility of multiplexing computations on sequential inputs in the brain, we leveraged a foraging task based on processing a stream of binary outcomes to inform a decision of whether to leave or stay^{7,8}. This task admits multiple strategies for processing the series of outcomes that are associated with different precisely quantifiable DVs. Evaluation of these DVs allows the experimenter to infer the implementation of ‘counterfactual’ strategies, that is, strategies that are potentially applicable, but unused. If such counterfactual strategies could be decoded from the brain, it would be evidence for parallel processing of serial information.

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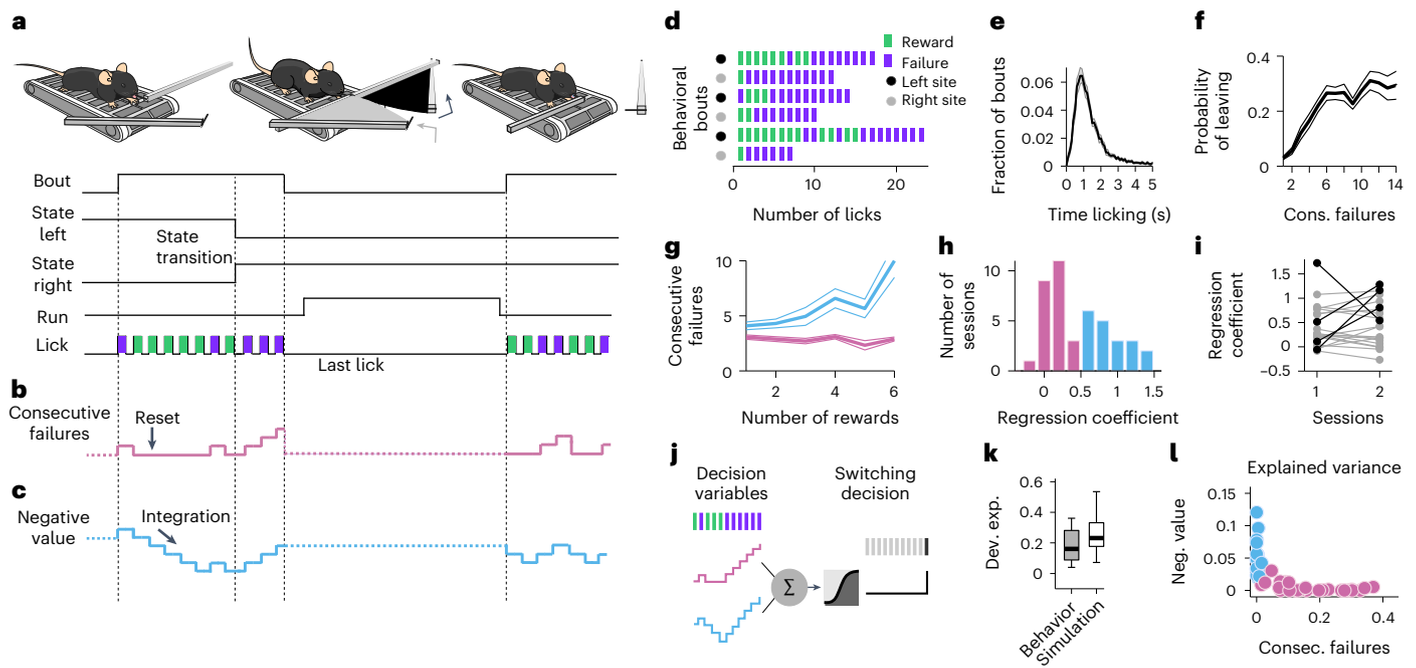


Fig. 1 | Multiple DVs predict foraging behavior. **a**, A head-fixed mouse placed on a treadmill chooses to exploit one of the two foraging sites (two movable arms on each side of the treadmill). A bout of behavior consists of a series of rewarded (1 μ l consumed in a single lick) and unrewarded licks at one of the sites. The switch from active to inactive state only happened once while the mouse was at the site, so if it left the site before the switch, no rewards were delivered at the other site (and it had to return to the original site and restart licking). Independently from state transition, animals can choose to switch between sites at any time by running a set distance on the treadmill. During site-switching, the spout in front moves away and the distal one moves into place. **b**, The DV that the mouse needs to compute to infer the hidden state of the resource site. **c**, Alternative DV supporting a stimulus-bound strategy—the ‘negative value’. **d**, Example sequences of observable events during different behavior bouts. **e**, Histogram of bout duration (mean \pm s.e.m. across sessions; $n = 42$). **f**, Probability of leaving the foraging site as a function of the number of consecutive failures after the last reward (mean \pm s.d. across mice). **g**, Consecutive failures before leaving as a function of reward number (mean \pm s.d.) in example sessions from two different mice. **h**, Distribution of the slope coefficients of an LM that predicted the number of consecutive failures before leaving as a function of

the number of prior rewards. For visualization, pink are the slope coefficients close to zero (coefficient < 0.5 , arbitrary threshold), while blue are sessions with positive slope coefficients. **i**, Slope coefficients from **h** between two consecutive sessions (1 and 2) for different mice. Sessions between which the coefficient values vary by more than 0.5 (arbitrary threshold) are highlighted in black. **j**, Illustration of the logistic regression model for predicting the switching decision of the mouse from the two different DVs. **k**, Deviance explained from the logistic regression that predicts choice behavior based on the DVs (gray box) and from simulated data where the behavior is truly inference-based (white box). On each box, the central mark indicates the median across behavioral sessions ($n = 42$ sessions from 21 mice), and the bottom and top edges of the box indicate the 25th and 75th percentiles, respectively. The whiskers extend to the most extreme data points. **l**, Explained variance from the logistic regression that predicts choice behavior based on the DVs. Sessions where ‘consecutive failures’ are dominant (var. exp. consec. failures $>$ var. exp. neg. value) are labeled in pink, while sessions where ‘negative value’ is dominant are labeled in blue (var. exp. consec. failures $<$ var. exp. neg. value). Var., variance; exp., explained; consec., consecutive; neg., negative.

Here using population recordings and optogenetic silencing in the frontal cortex of mice performing the foraging task, we identified a brain region (the secondary motor cortex M2) where the multiple DVs used by the mice could be decoded simultaneously. Critically, we found that the extent to which each DV was represented in the cortex did not depend on the strategy used by each mouse. These observations suggest that mice use an algorithm for decision-making that relies on the parallel computation of multiple DVs in the frontal cortex.

Results

Multiple DVs predict switching decision

In our task, a head-fixed mouse collected probabilistically delivered rewards at a virtual foraging site by licking from a spout (Fig. 1a and Extended Data Fig. 1). At any time, the mouse could choose to continue licking or give up and explore a new site by starting to run. There were two virtual foraging sites, only one of which was active at a given time and would deliver a reward with a probability of 0.9 after each lick. The active site also had a probability of 0.3 of switching once per bout after each lick⁸. Therefore, the best strategy to time the switching decision was to infer the latent state corresponding to which site was currently active⁸. This inference-based strategy was supported by a particular DV

that consisted of temporally accumulating consecutive failures with a complete reset upon receiving a reward (Fig. 1b). This is because a failure to receive reward provides only partial evidence that the active state had switched, whereas a reward always signaled the active state with certainty. Using this strategy, mice would leave the current site when the ‘consecutive failures’ DV reaches a given threshold⁸. Yet, in principle, mice could time their decision to leave by using any number of alternative strategies based on the sequence of rewarded and unrewarded licks regardless of the true causal structure of the task. In fact, early on during training when learning the task, mice do not appear to calculate the inference-based DV⁸. Their behavior is better described by a strategy that does not contemplate discrete transitions to a fully depleted state, and instead relies on a running estimate of the ‘value’ of the current site based on the difference between recently observed rewards and failures (Fig. 1c). Using this strategy, mice decide to abandon a foraging site when its value is sufficiently low (or its negative value is sufficiently high). We refer to this as a stimulus-bound strategy because it treats observable outcomes (the stimuli) as direct—although probabilistic—reporters of the valence of current environmental states, without further assumptions or models about environmental dynamics. For our present purposes, the essential aspect of these two strategies

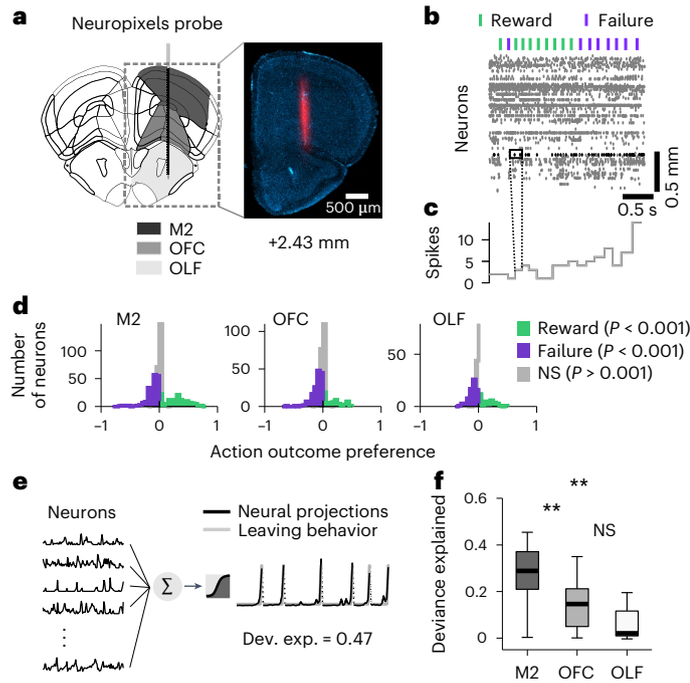


Fig. 2 | Neural activity related to the switching decision. **a**, Schematic target location of probe insertion and an example histology of electrode track (1 of 10 example brain). Vertical insertions were performed within a 1-mm diameter craniotomy centered around +2.5-mm anterior and +1.5-mm lateral from Bregma. **b**, Example raster plot of 140 simultaneously recorded neurons from M2. Lick-outcome times are indicated by the green (reward) and purple (failure) dashes. **c**, Binned response profile of an example neuron. For all analyses, otherwise noted, we averaged for each neuron the number of spikes into bins by considering a 200 ms window centered around each lick. **d**, Histogram of outcome selectivity of all neurons recorded M2 (left), OFC (middle) and OLF (right). We used ROC analysis to assign a preference index to each neuron. In brief, an ideal observer measures how well the modulation of neuronal firing can classify the outcome (reward or failure) on a lick-by-lick basis. We derived the outcome preference from the area under the ROC curve as $PREF_{R,F} = 2(ROC_{AREA}(f_R, f_F) - 0.5)$, where f_R and f_F are the firing rate distributions for trials where outcomes are reward and failure, respectively. This measure ranges from -1 to 1 , where -1 indicates preference for F (failure), 1 means preference for R (reward) and 0 represents no selectivity. The statistical significance of the preference index ($P < 0.001$, one-sided) was assessed via bootstrapping (1,000 iterations). Violet and green bars indicate neurons where the index was significantly different from 0 . In all regions, we found neurons significantly modulated by rewards and failures. **e**, Illustration of the logistic regression method for predicting the switching decision (gray right, that is, the probability that each lick was the last in the bout, $n = 2,533 \pm 1,524$ licks per session; mean \pm s.d. across 11 sessions) from the principal components of neurons (left—M2, $n = 31 \pm 17$; OFC, $n = 29 \pm 9$; OLF, $n = 16 \pm 13$). **f**, Deviance explained from the logistic regression in each region. Two stars indicate a significant difference between regions (two-sided Wilcoxon signed rank test, $P = 0.0068$ between M2 and OFC; $P = 0.0049$ between M2 and OLF; $P = 0$ between OFC and OLF). On each box, the central mark indicates the median across recording sessions ($n = 11$ sessions), and the bottom and top edges of the box indicate the 25th and 75th percentiles, respectively. The whiskers extend to the most extreme data points. NS, not significant; ROC, receiver operator characteristic.

is that they use the same observable outcomes (series of rewarded and unrewarded licks) in qualitatively different ways to update their corresponding DV—a full reset versus a quantitative incremental increase in current value. This allows us to unambiguously identify the two DVs, their behavioral consequences and their neural representations.

After several days of interaction with this setup ($n = 13 \pm 5$ d; mean \pm s.d.), mice ($n = 21$) learned to exploit each site for several seconds (Fig. 1d,e). Considering the last two sessions of training ($n = 42$

sessions total), we examined which strategy mice used to time their leaving decisions. As demonstrated previously⁸, for all mice, the probability of leaving increased with the number of consecutive failures (Fig. 1f). Yet not all mice treated rewards equally. For some mice, the number of previous rewards did not affect the probability of leaving after a set number of failures (Fig. 1g, pink), consistent with the inference-based strategy. In contrast, for some other mice, the number of failed attempts that they tolerated before leaving the site correlated with the number of previous rewards (Fig. 1g, blue), consistent with the stimulus-bound strategy. We quantified these effects using a linear regression model (LM) that predicted the number of consecutive failures before leaving as a function of the number of prior rewards in the current bout (Fig. 1h). We found that the regression coefficient varied strongly within our cohort, consistent with the just-described behavioral heterogeneity across sessions. The distribution across sessions showed signs of bimodality with a dip close to 0.5. Using this criterion, the behavior was more consistent with the inference-based strategy in $n = 23$ sessions (coefficient < 0.5) and more consistent with the stimulus-bound strategy in the remaining $n = 19$ sessions (coefficient > 0.5). To check if the heterogeneity in strategy was due to variability from session-to-session, mouse-to-mouse or both, we examined whether the regression coefficients of each mouse varied across consecutive sessions (Fig. 1i). Overall, we observed that most mice kept the same dominant strategy across consecutive sessions (Fig. 1i, gray; but see also Fig. 7), but some mice ($n = 4$) also switched strategy from one session to the next (Fig. 1i, black).

These observations indicate that mice vary in their foraging strategies across individuals and sessions but do not directly indicate how well the mice’s behavior is described by the DVs. Therefore, we next quantified how well the different DVs could predict the precise moment (lick) when an individual mouse would switch sites on a given trial. Specifically, we used regularized logistic regression to model the probability that each lick ($n = 2,882 \pm 1,631$ licks per session; mean \pm s.d. across 42 sessions) was the last one in the bout, considering simultaneously the two DVs as predictors (Fig. 1j; Methods). We estimated the goodness of fit of the two models using the ‘deviance explained’, where ‘0’ meant chance level and ‘1’ meant perfect predictions. We found a median deviance explained of 0.16, a value significantly better than chance level for all mice (Fig. 1k, gray box; Wilcoxon rank test, $P < 10^{-6}$). To provide a reference for the meaning of deviance of this magnitude, we used the same logistic regression model to predict the leaving decisions of a simulated agent in which the ‘ground truth’ was known. For this, we simulated behavioral sessions of an agent making decisions using a logistic function and the DV of the inference-based strategy with equal numbers of bouts as in the real sessions. We found that the model recovered the ground truth parameters with high accuracy (Extended Data Fig. 2a–d) and performed better than a model attempting to fit the same data using the stimulus-bound DV, which is distinct but correlated with the DV of the inference strategy (Extended data Fig. 2e). Furthermore, the deviance explained by the simulated data (median = 0.25; Extended Data Fig. 2f,g) was only slightly greater than that of the real data (Fig. 1k), indicating that the model with DVs performed close to the maximum that could be expected given the statistical nature of the task. This multivariate approach also confirmed that the two DVs were used to different extents across sessions (Fig. 1l) and, compared to the univariate regression (Fig. 1h), provided even clearer indication of changes in dominant strategy across sessions (Fig. 1l and Extended data Fig. 2h). Finally, the bias term of the model (or intercept) reflected the baseline probability to leave the site (the larger the bias the more impulsive the switching decision) but did not correlate with the use of either DV (Pearson correlation between bias term and explained variance of consecutive failures: $r = -0.12$, $P = 0.44$; negative value $r = -0.18$, $P = 0.25$).

The logistic regression confirmed that the two DVs describe the switching decision relatively well. Yet, alternative strategies not directly

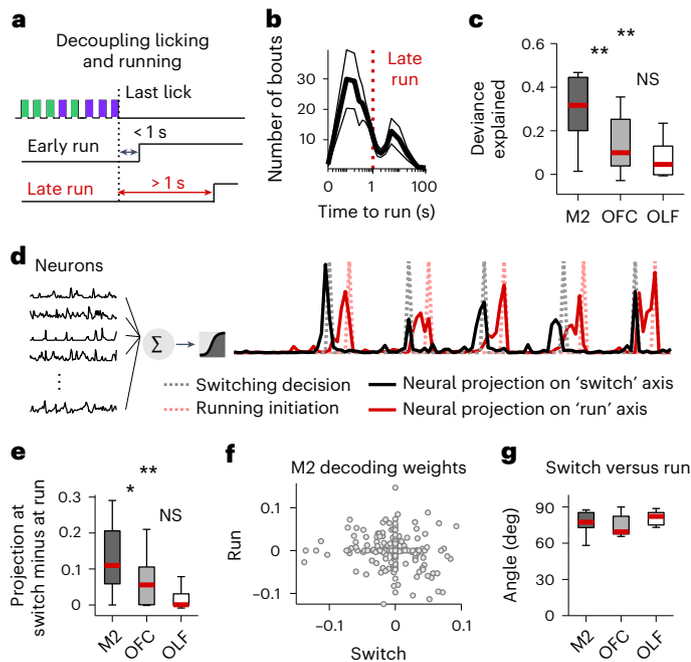


Fig. 3 | Switching decision and running initiation are dissociable. **a**, Last lick always precedes running initiation. Running initiation may occur immediately after the last lick of a bout (<1 s; 'early run') or mice may remain still for several seconds after the last lick and before running initiation (>1 s; 'late run') in record. **b**, Bimodal distribution of time between last lick and running initiation (mean \pm s.e.m. across recording sessions). **c**, The deviance explained from models that predict the switching decision (last lick) from the neural activity from M2 (dark gray), OFC (light gray) and OLF (white), in 'late run' conditions when the last lick is fully decoupled from running initiation. Two stars indicate a significant difference between regions (two-sided Wilcoxon signed rank test, $P = 0.002$ between M2 and OFC; $P = 0.002$ between M2 and OLF). **d**, Illustration of the logistic regression method for predicting the switching decision (gray dash line) and the running initiation (red dash line) using neural activity from first lick to running initiation (black, left) in bouts when running occurred at least 1 s after the last lick. Red and black solid lines are examples of neural projections onto the two different axes. **e**, Difference in values of the neural projection onto the switch axis at the time of switching and the time of running. The larger the difference, the more dissociable the two events. Two stars indicate a significant difference between regions (two-sided Wilcoxon signed rank test, $P = 0.027$ between M2 and OFC; $P = 0.002$ between M2 and OLF; $P = 0$ between OFC and OLF). **f**, Decoding weights of each M2 neuron (gray dots) for the two different axes. **g**, Angles between the two different axes. In all regions, the angle is close to 90° indicating that the two axes are close to orthogonal. On each box of **e**, **e** and **g**, the central mark indicates the median across recording sessions ($n = 11$ sessions), and the bottom and top edges of the box indicate the 25th and 75th percentiles, respectively. The whiskers extend to the most extreme data points.

relying on combinations of action outcomes could also explain well the mice's behavior. Thus, we used the logistic regression model to further explore the space of strategies beyond the two main DVs (Extended Data Fig. 3a). We found that whereas alternative strategies explained some of the behavioral variances, the 'consecutive failures' and 'negative value' DVs still best predicted the switching decision in most sessions (Extended Data Fig. 3b,c). Although we cannot rule out that mice use other unexplored strategies, these results indicate that the inference-based and stimulus-bound strategies are the best predictors of the switching decision among different classes of foraging strategies.

Neural activity related to the switching decision

To examine the neural basis of DVs underlying the switching decision, we first had to identify brain regions that predicted the switching decision. We used Neuropixels 1.0 (ref. 9), which are single shank probes with

hundreds of recording sites that allow registering the activity of large ensembles of neurons ($n = 151 \pm 59$ neurons per session; mean \pm s.d.) in multiple regions of the frontal cortex during the task. We targeted M2 ($n = 66 \pm 37$ neurons per session; mean \pm s.d.), thought to be important for timing self-initiated actions¹⁰, planning licking behavior¹¹ and predicting changes in behavioral strategy¹², and the orbitofrontal cortex (OFC; $n = 55 \pm 24$ neurons per session; mean \pm s.d.), whose inactivation impacted the performance of inference-based decision-making in freely moving mice in the foraging task⁸. We also recorded in the olfactory cortex (OLF; $n = 31 \pm 23$ neurons per session; mean \pm s.d.), which is directly ventral to the OFC (Fig. 2a,b and Extended Data Fig. 4), but which would not be expected to be specifically involved in this task.

To examine neural responses during the evidence accumulation process, we considered the momentary response patterns of isolated neurons in small time windows (Fig. 2c; Methods). Because we observed heterogeneous task-related activity in many single neurons in all regions (Fig. 2d), we focused on how population activity from each single region predicted the switching decision of mice ($n = 11$ recording sessions, one recording session per mouse except one mouse with two recording sessions). Using cross-validated and regularized logistic regressions, we decoded the switching decision from population responses around each lick (200 ms window) in each session (Fig. 2e). To allow for a fair comparison between brain regions, we controlled for the different number of recorded neurons in each region by using as predictors only the first n principal components of neural activity, which predicted up to 95% of its total variance (Methods for additional control analyses). We found that the switching decision could be better decoded using population activity from neurons in M2 than in OFC or OLF (Fig. 2f). This suggests that, unlike OFC, which has been shown to be important for the inference process⁸, M2 may be directly involved in the instantaneous timing of action selection.

Switching decision and running initiation are dissociable

To test that the neural activity predictive of a switching decision does not simply reflect running initiation, we decoded the switching decision on a subset of behavioral bouts where the last lick and running initiation were clearly decoupled (Fig. 3a,b). We found that the last lick could still be decoded with high accuracy, especially in M2 (Fig. 3c), suggesting that M2 activity encodes the intention to switch sites rather than just reflecting the initiation of running behavior.

To further test whether the switching decision and running initiation are dissociable in M2, we used neural activity up to the point of running initiation to simultaneously decode the switching decision and the decision to initiate running, again using only bouts where licking and running were decoupled in time (Fig. 3d). The neural activity projected onto the two decoding axes (switching and running) peaked at different times (Fig. 3d,e), and the two axes were close to orthogonal (Fig. 3f,g), consistent with previous studies showing that M2 populations encode preparatory activity for upcoming actions^{11,13}. These results indicate that M2 simultaneously encodes, in a separable format, the relevant DVs used to guide an action, as well as a signal associated with the time of initiation of the action itself.

M2 is involved in the switching decision

The above results point to M2 as a key region for timing the switching decision by relying on specific DVs. To further test the contribution of M2 to the implementation of DVs, we partially silenced M2 using an optogenetic strategy (as in ref. 8; Fig. 4a). Specifically, we examined 43 sessions from six mice, four of which were ChR2-expressing and two of which were control wild-type littermates that express no inhibitory opsin implanted and stimulated in the same manner. M2 silencing caused no gross changes in action timing (Extended data Fig. 5), but only a slight decrease in licking rate (Extended data Fig. 5c), and perhaps a trend for increasing the time spent licking (Extended data Fig. 5d). Because M2 inactivation did not substantially impair the

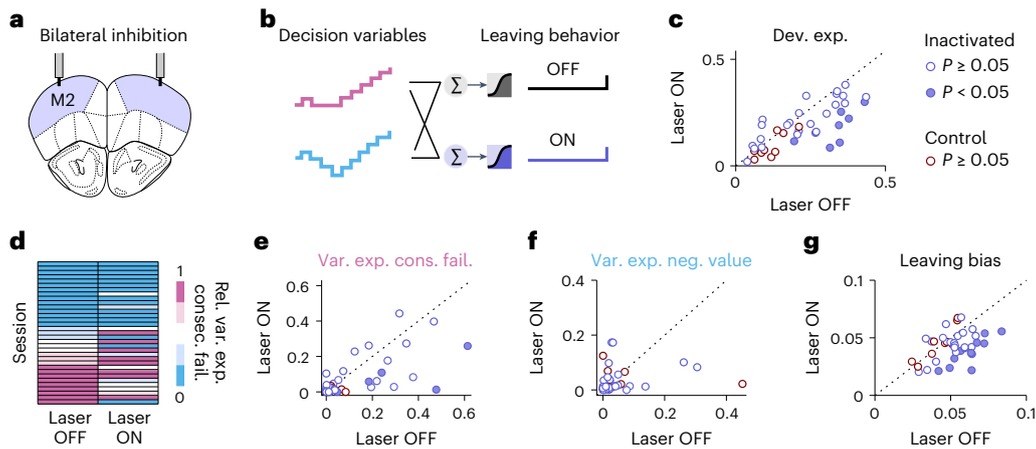


Fig. 4 | M2 is involved in the switching decision. **a**, Schematic target of optic fibers placement in the brain of VGAT-ChR2 mice, which express the excitatory opsin channelrhodopsin-2 in inhibitory GABAergic neurons. Bilateral photostimulation to silence M2 (5 mW power per fiber, 10 ms pulses at 75 Hz) was triggered by the first lick in 30% of randomly selected bouts and lasted until the last lick of the bout. **b**, Illustration of the logistic regression models for independently predicting the switching decision of the mouse based on the DVs during photostimulation (Laser ON) and control bouts (Laser OFF) in each session ($n = 43$). **c**, Deviance explained from the models in **b** for each session (dots) of inactivated mice (violet) and control mice (red). Dots below the identity indicate the sessions where the model performed worse during photostimulation of M2. Fixed effect of stimulation ('Laser' predictor)—inactivated: -0.04 ± 0.02 , $P = 0.021$; control: -0.03 ± 0.014 , $P = 0.054$. **d**, Relative variance explained of the DVs for predicting the switching decision in 'Laser OFF' versus 'Laser ON' condition. Because both DVs are used as regressors, their relative variances explained sum to 1. Larger values of the relative variance explained of the 'consecutive failures' are colored in pink and indicate that the mouse mainly uses

the inference-based strategy. Conversely, lower values of the relative variance explained of 'consecutive failures' are equivalent to larger values of relative variance explained of 'negative value' (colored in blue), indicating the mouse mainly uses the stimulus-bound strategy. **e**, Variance explained of 'consecutive failures' in 'Laser OFF' versus 'Laser ON' condition. Fixed effect of stimulation—inactivated: -0.054 ± 0.025 , $P = 0.032$; control: -0.012 ± 0.009 , $P = 0.22$. **f**, Variance explained of 'negative value' in 'Laser OFF' versus 'Laser ON' condition. Fixed effect of stimulation—inactivated: -0.011 ± 0.012 , $P = 0.35$; control: -0.032 ± 0.045 , $P = 0.49$. **g**, Bias term of the logistic regression (intercept) in 'Laser OFF' versus 'Laser ON' condition. Fixed effect of stimulation—inactivated: -0.45 ± 0.078 , $P < 10^{-6}$; control: 0.092 ± 0.075 , $P = 0.24$. **c, e–g**, Filled dots indicate that the effect of photostimulation is significant within single sessions ($P < 0.05$, two-sided, estimated using bootstrapping; Methods). To estimate the effect of photostimulation on the deviance explained across mice and session, we used the following mixed model (Methods): Dev.Exp. $\sim 1 + \text{Laser} + (1 + \text{Laser} | \text{Mouse}) + (1 + \text{Laser} | \text{Session})$. Dev. Exp., Deviance explained.

motor behavior, we tested if silencing M2 affected the use of the DVs to time the leaving decision (Fig. 4b). We found that the inactivation of M2 substantially decreased the predictive power of the DVs (Fig. 4c, violet). The same protocol applied to control mice had no significant effect on this behavior (Fig. 4c, red). The photostimulation decreased the use of the DV 'consecutive failures' (Fig. 4d–f), as well as the leaving bias (Fig. 4g), making animals less inference-based and less impulsive. These results suggest that M2 is part of the neural pathway through which the DVs shape the behavior of the mice.

Neural representation of DVs

The inactivation experiments suggest that one might be able to read out the DV used by the mouse from M2 neural activity and that M2 might represent this DV better than other cortical regions that afford less accurate predictions of foraging decisions. To test these ideas, we used regression-based generalized linear models (GLM; Methods) to decode the instantaneous magnitude of the DV associated with the behaviorally dominant strategy (that is, the DV most predictive of behavior; Fig. 5a,b). The example data from Fig. 5a,b, which are from a single recording session during which the dominant strategy of the mouse was the inference (var. exp. consec. fail. = 0.164 versus var. exp. neg. value = 0.004), show that the related DV 'consecutive failures' could be decoded with high accuracy from M2 activity. In fact, the dominant DV could be well decoded from M2 activity in all sessions ($n = 11$) from the different mice (Fig. 5c, black). The decodability of dominant DVs was substantially lower in other cortical regions (Fig. 5c, gray and white), consistent with the poorer decoding of leaving time in other areas (Fig. 2f).

Because we have shown that different mice can rely on different DVs and individual mice can change decision strategies across sessions

(Fig. 1), we next asked whether session-by-session heterogeneity in decision strategy could be explained by the degree to which M2 neurons reflected the DVs in each session. Here we used the GLM to compare the decoding of the dominant and the alternative DVs from M2 neurons in each recording session (Fig. 5a,d). Contrary to our expectation, we found that decoding was similar between the dominant and alternative decision strategies. For instance, in the example session of Fig. 5a,b,d, despite the selectivity of the behavior for inference-based decisions, the DV supporting the stimulus-bound strategy could also be well decoded from M2. This finding was consistent across our experiments—in all sessions, the DVs could both be read out from M2 activity (Fig. 5e and Extended Data Fig. 6). On average, the 'consecutive failures' DV was somewhat better represented than the 'negative value' (Fig. 5e). This average difference could stem from the fact that the majority of mice (8 of 11) used the inference-based strategy that relies on the 'consecutive failures'. Thus, to test whether the DV that was most predictive of the switching decision was also the one that was better decoded from M2 on a session-by-session basis, we predicted the decision to switch sites from each DV (Fig. 4f) and compared the accuracy of this prediction to the accuracy of the neural representations of the DVs (Fig. 5g). There was no correlation between how M2 represented each DV in a session and how well the DV predicted behavior in the same session ($P = 0.9$). Together these analyses suggest that whereas M2 neural activity is important to the execution of a decision strategy (Fig. 4), the pattern of neural activity in M2 is not adapted to represent specifically the DV executed by the mouse, and instead reflects a broader range of decision strategies even when they are not currently used.

To further characterize the multiplexing of DVs in M2, we asked whether different variables are supported by distinct or overlapping populations. We examined the weights assigned to each neuron when

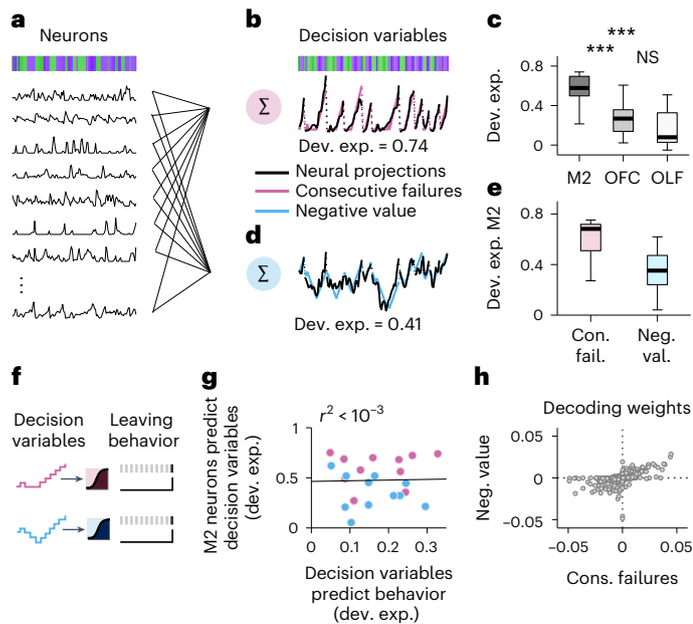


Fig. 5 | Neural representation of DVs. **a**, The regression models take as predictors the activity of simultaneously recorded neurons (black traces) and derive a set of decoding weights to predict the DV. The method is applied during each bout ($n = 223 \pm 119$ bouts per session; mean \pm s.d.) on a lick-by-lick basis—from the responses of neurons (the putative single units) in each recorded brain region. **b**, Predictions of the model (black trace is the weighted sums of neural activity) overlaid onto the ‘consecutive failures’ DV (pink trace). **c**, Deviance explained across sessions ($n = 11$ sessions, median \pm 25th and 75th percentiles, the whiskers extend to the most extreme data points) from the model in **a** and **b** in each cortical region. The stars indicate the significance of two-sided Wilcoxon signed rank tests ($P = 0.00098$ between M2 and OFC; $P = 0.00098$ between M2 and OLF; $P = 0$ between OFC and OLF). **d**, Predictions of the model (black trace is the weighted sums of neural activity) overlaid onto the ‘negative value’ DV (blue trace). **e**, Deviance explained across sessions ($n = 11$ sessions, median \pm 25th and 75th percentiles, the whiskers extend to the most extreme data points) predicted from M2 neurons for each DV. Two-sided Wilcoxon signed rank test: $P = 0.00098$. **f**, Illustration of the logistic regression methods for predicting the switching decision of the mouse from each DV separately. **g**, Correlation between the neural representations of different DVs (color-coded as in **b** and **d**) and how well each DV predicts behavior. Each dot corresponds to a particular DV from a given recording session. The linear regression is reported in black. **h**, Decoding weights of each M2 neuron (gray dots; total across recording $n = 778$) for the two different DVs. Pairwise linear correlation—Pearson coefficient = 0.56 , $P < 10^{-4}$.

decoding the two DVs (Fig. 5h). We found that decoding weights for both DVs were strongly correlated, indicating a considerable overlap between the populations of M2 neurons that supported each DV, as opposed to compartmentalization into distinct populations for each variable.

Independent representations of DVs

A possible concern with the interpretation that M2 multiplexes used and unused DVs is that alternative DVs might be decodable only by virtue of being like the one reflected behaviorally. Although the computations underlying the two DVs are different, for the sequences of rewards and failures experienced by the mice, the DVs themselves are correlated overall (Pearson coefficient: 0.79 ± 0.15 ; mean \pm s.d.).

As a first strategy to overcome this limitation, we took advantage of the fact that the two different DVs differ in the way that they treat rewards: while the ‘negative value’ requires negative integration of rewards, the ‘consecutive failures’ requires a complete reset by a single reward (Fig. 6a). Analysis of subsets of sequences that consist of multiple consecutive rewards should therefore reveal the

differences between the two DVs (Fig. 6b). To test this, we sub-selected lick sequences and sorted them according to the relative number of rewards and failures. This produced subsequences with varying degrees of correlation between the two decision variables (Fig. 6c). We then ran the same decoding analyses as before on these subsequences of M2 activity. We found that the ability to decode the subsequences was independent of their degree of correlation (Fig. 6d). Our second approach was to investigate whether we could decode the component of each DV that is uncorrelated with the other one, that is, its residual. Indeed, we could decode the residuals from both DVs from the activity of M2 populations (Fig. 6e,f). Together, these results establish that the ability to decode an alternative DV does not arise from the correlation of that variable with the dominant DV. Interestingly, this approach revealed that OFC better represented the ‘consecutive failures’, consistent with previous work suggesting that OFC is important for the inference-based strategy⁸.

Using only the sequences of trials in which the DVs were fully decorrelated (Pearson correlation between DVs: 0.03 ± 0.02 ; median \pm MAD across session), we again tested the possibility that the DVs that were best decoded from M2 were the most predictive of behavior (as in Fig. 5e,g,h). Here the ‘consecutive failures’ remained better represented than the ‘negative value’ (Fig. 6h). Like the results with the intact DVs, there was no correlation between how M2 represented each decorrelated subset of DVs and how well the DV predicted behavior (Fig. 6i). This was the case even if the populations of M2 neurons that supported each decorrelated subset of DVs were nearly orthogonal, as indicated by the small correlation between decoding weights (Fig. 6j).

DV multiplexing does not reflect strategy switching

While one interpretation of multiplexing is true simultaneous representation of multiple DVs, our interpretation is relying on decoding analyses carried out over entire sessions of behavior. Could it be that multiplexing of DVs results from sequential switching between the two strategies within a session? To investigate this, we first examined whether there was any evidence that mice switched strategies within a session using a framework based on hidden Markov models (HMM) combined with LM (Methods; ref. 14). The resulting ‘LM-HMM’ framework modeled the number of consecutive failures that the animal bears before switching sites using the following two inputs: (1) the total number of rewards, which allows distinguishing between inference-based (that is reward independent) and stimulus-bound (that is reward dependent) strategies, as in Fig. 1g, and (2) a constant bias, which reflects the level of impulsivity of the animal. Each hidden state in the model captures a specific dependence of consecutive failures on the total rewards and the bias, characterizing a particular decision-making strategy.

A model with three states best described the switching decision and yielded interpretable and persistent states (Fig. 7a and Extended Data Fig. 7a). One of the states had a large weight on the number of rewards, indicative of a stimulus-bound strategy, while the other two had negligible weights on rewards, consistent with the inference (Fig. 7b and Extended Data Fig. 7b,c). To visualize the temporal structure of the foraging decision within a session, we computed the posterior probability over the latent states across all behavioral bouts (Fig. 7c,d), which revealed that mice mostly remained in discrete states (average probability of the dominant strategies over all bouts: 0.91 ± 0.06 ; median \pm MAD across 11 sessions) for many bouts in a row (average duration of states: 56 ± 41 bouts; median \pm MAD across sessions), but tended to switch states at least once per session (state transition in 8 of 11 sessions; Extended Data Fig. 7d).

Because mice alternated between states of inference-based and stimulus-bound strategies within the course of their recording session, we examined whether we could decode better from M2 activity the ‘consecutive failures’ DV during the inference-based states than during the stimulus-bound states (Fig. 7e, pink dots), and vice versa for ‘negative

value' DV (Fig. 7e, blue dots). Consistent with the whole-session analysis (Fig. 5g), there were no significant differences between how well a given DV could be decoded when the mice's behavior relied on it or when it did not. The residual signals after the DVs, which are orthogonalized, were also decodable in their respective alternate states (Fig. 7f). These analyses suggest that multiplexing of strategy is not due to the switch of strategies within a session.

M2 represents foraging algorithms

Given that M2 appears to multiplex different DVs, we wondered whether this might reflect a generic capacity to represent any signal with similar temporal characteristics as the DVs in the task, as predicted by the reservoir computing framework¹⁵⁻¹⁷. Decoding analyses of artificial signals with matched temporal statistics revealed this not to be the case (Extended Data Fig. 8). Therefore, we next considered that the space of signals encoded in M2 might be restricted to potentially meaningful variables generated from a common set of essential computations. Here the two DVs we have been considering could both be conceptualized

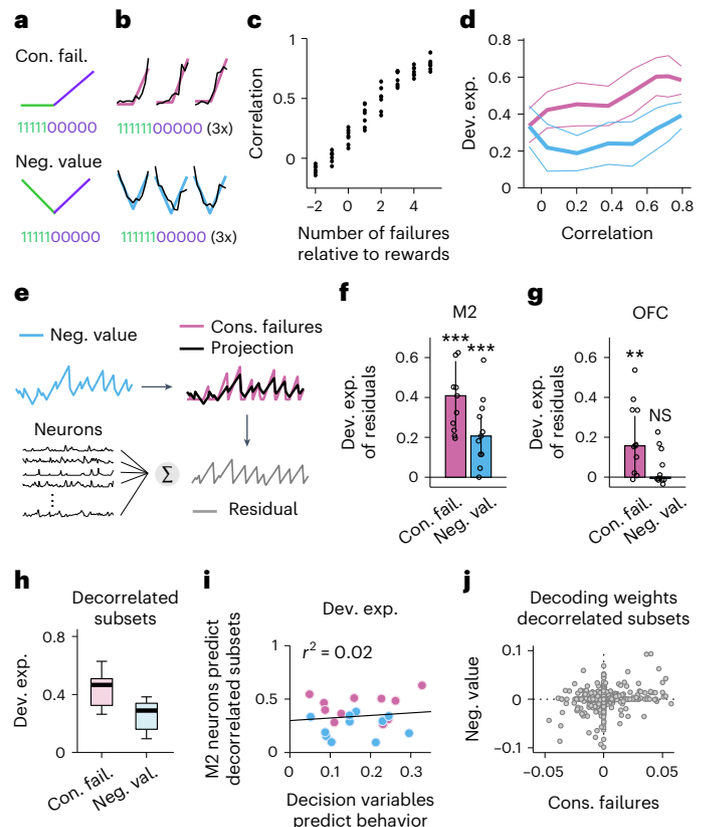
as an adaptive, outcome-dependent feedback gain on a running count. For instance, if we refer to the running count after the t -th lick as x_t and to the outcome of the next lick as o_{t+1} (equal 1 or 0 if the outcome is a reward or a failure, respectively), then we can write the update rule compactly as

$$x_{t+1} = g(o_{t+1})x_t + c(o_{t+1})$$

with $g(o_{t+1} = 1) = 0$, $g(o_{t+1} = 0) = 1$ and $c(o_{t+1} = 1) = c(o_{t+1} = 0) = 1$ for the inference-based DV, and $g(o_{t+1} = 1) = g(o_{t+1} = 0) = 1$ and $c(o_{t+1} = 0) = -c(o_{t+1} = 1) = 1$ for the stimulus-bound DV. This realization suggests that a common generative model, which we named the 'INTEGRATE-AND-RESET model', can produce these two different DVs by adjusting certain model parameters (Fig. 8a). The INTEGRATE-AND-RESET model describes, within a single algorithmic framework, the computations necessary to generate, not only the two DVs considered so far but also other DVs relevant for a variety of other commonly studied behavioral tasks. For instance, a 'global count' (accumulated number of outcomes) DV is related to counting or timing tasks^{18,19}. Similarly, matching tasks involving randomly timed cached rewards are optimally solved by integrating the difference between rewards and failures with an exponential decay²⁰. Sequential foraging in patchy environments is also solved by integrating the difference between rewards and failures, equivalent to tracking the relative 'negative value' of a foraging site²¹. Other integration tasks, like the 'Poisson clicks' task²², require perfect integration of two variables. Thus, the space of DVs generated by the INTEGRATE-AND-RESET model covers a large space of tasks that have been studied in the lab and might be useful in different behavioral contexts.

All nontrivial time series produced by the INTEGRATE-AND-RESET model can be expressed as linear combinations of four basis sequences (Fig. 8a; Methods). The two sequences involving reset describe integration of failures and reset by rewards ('consecutive failures') and vice versa ('consecutive rewards'). The two sequences for accumulation

Fig. 6 | Independent representations of DVs. **a**, Two different sequences relying on different computations involving reset (top) and accumulations (bottom) of rewards. **b**, Three example bouts (columns) of population activity (black traces) projected onto the dimensions that best predict the trajectory of the different sequences (color traces). Only subsequences of consecutive rewards followed by consecutive failures were selected to visualize the different computations in **a** (~5% of bouts). **c**, Selecting subsets of action outcomes where the total number of failures changes relative to the number of rewards (abscissa) alters the correlation between sequences generated with the computations in **a** (ordinates). Black dots for each value of the number of failures represent a recording session. **d**, How well the sequences relying on the two different computations can be decoded from M2 (ordinates) as a function of the correlation between them (median \pm median absolute deviation or MAD, across sessions, one-way ANOVA for each sequence across correlation values followed by multiple pairwise comparison tests, all P values > 0.05). Pink are sequences that accumulate failures and reset with rewards (equivalent to 'consecutive failures'). Blue are sequences that accumulate failures upward and rewards downward (equivalent to 'negative value'). **e**, Schematic description of our strategy to linearly regress each of the two DVs on the other. This approach allowed us to express DV1 (for example, 'consecutive failures') as the sum of a time series proportional to DV2 (for example, 'negative value') plus a time series orthogonal (uncorrelated) to DV2, which we denote as its residual. Here the 'consecutive failures residual' (gray) is orthogonal to the 'negative value' (blue). The same procedure was used to generate the 'negative value residual' orthogonal to the 'consecutive failures'. Both residuals were then fit by M2 neurons. **f**, Deviance explained across sessions ($n = 11$ sessions, median \pm MAD) of the model in **e**. Pink, residual consecutive failures; blue, residual negative value. The residuals relative to each DV were both significantly represented in M2 (two-sided Wilcoxon rank sum test, $P = 0.00098$ for both, indicated by the stars). The size of the pink bar measures how well one can decode the part of 'consecutive failure' orthogonal to 'negative value' (residual consecutive failures) and the size of the blue bar measures how well one can decode the part of 'negative value' orthogonal to 'consecutive failure (residual negative value)'. If only 'consecutive failures' were represented, the residual consecutive failures should be represented, but the residual negative value would not be represented. On the other hand, if both DVs are represented, both residuals should be decodable, as shown here in **M2, g**. Same as in **f** but with OFC neurons. The residuals 'consecutive failures' were decodable from OFC ensembles (pink; two-sided Wilcoxon rank sum test, $P = 0.0029$), but the residuals 'negative values' were not (pink; Wilcoxon rank sum test, $P = 0.52$). **h**, Deviance explained across sessions ($n = 11$ sessions, median \pm 25th and 75th percentiles, the whiskers extend to the most extreme data points) predicted from M2 neurons for each decorrelated subsets of DVs (two-sided Wilcoxon signed rank test: $P = 0.00098$). **i**, Correlation between the neural representations of decorrelated subsets of DVs (color-coded as in **b, d** and **e**) and how well each DV predicts behavior. Each dot corresponds to a particular DV subset from a given recording session. The linear regression is reported in black ($r^2 = 0.02$, $P = 0.6$). **j**, Decoding weights of each M2 neuron (gray dots; total across recording $n = 778$) for the decorrelated different subsets of DVs (two-sided Pearson coefficient between decoding weights = 0.20, $P < 10^{-7}$).



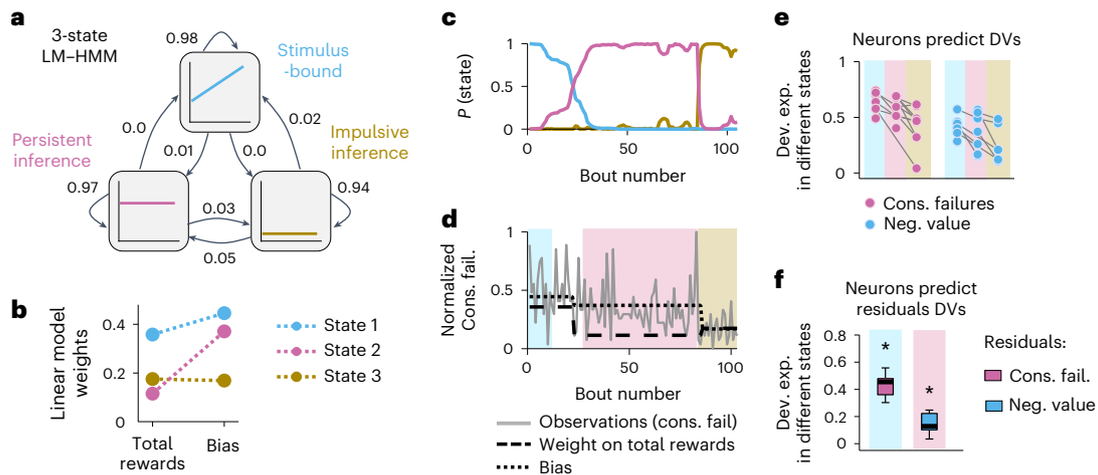


Fig. 7 | Simultaneous representations of DVs. a, Illustration of the LM-HMM, with three different states corresponding to different decision-making strategies (labeled ‘stimulus-bound’, ‘persistent inference’ and ‘impulsive inference’). The high self-transition probabilities of 0.94, 0.97 and 0.98 indicate that states typically persist for many consecutive bouts. The transition probabilities are indicated by the arrows between states. **b**, LM weights for the three-state model fit to all sessions simultaneously. **c**, Posterior state probabilities (computed with Gaussian prior on the weights and Dirichlet prior on transition probabilities) for an example session showing that states typically persisted for many consecutive bouts with high model confidence but transitioned once or twice over the course of a session. **d**, Behavioral data and model parameters of the example session. The gray line indicates the number of consecutive failures (that is, observations of LMs). The shaded color background indicates the high confidence state ($P(\text{state}) > 0.75$). Dash black lines indicate the LMs bias and weights in each state.

e, Deviance explained from models that fit M2 neurons to the DVs (pink dots, consecutive failures; blue dots, negative value) in different states (high model confidence, indicated by the color background). High confidence intervals were defined as $P(\text{state}) > 0.75$ for at least 25 consecutive bouts. Each dot is a recording session. Comparisons between pairs of states were made with two-sided Wilcoxon rank sum test ($P > 0.05$ for all comparisons). **f**, Deviance explained across sessions ($n = 11$ sessions, median ± 25 th and 75th percentiles, the whiskers extend to the most extreme data points) from models that fit M2 neurons to the residual DVs in their respective alternate states of high certainty. Left is the residual consecutive failures (the signal that is orthogonal to the negative value DV) in the stimulus-bound state. Right is the residual negative value (the signal that is orthogonal to the consecutive failure DV) in the inference-based states. Stars indicate that the deviance explained is significantly different from zero (two-sided Wilcoxon rank sum test; left, $P = 0.031$; right, $P = 0.016$).

without reset are upwards integration of both rewards and failures (equivalent to ‘count’) and integration upwards of rewards and downwards of failures (equivalent to ‘negative value’). We already know that M2 simultaneously represents two of these basis elements (‘consecutive failures’ and ‘negative value’). Thus, we tested whether M2 also represented the two additional basis sequences. We found that, indeed, ‘consecutive reward’ and ‘count’ could be decoded from the M2 population (Fig. 8b) and remained decodable from the M2 population when using the subsequences that decorrelate the variables (Fig. 8c).

The INTEGRATE-AND-RESET model can be extended, through analog values of ‘g’, to produce sequences with different dynamics and various time constants (Fig. 8d, left). Note that adjusting analog parameter values can directly relate the INTEGRATE-AND-RESET model to frameworks of reinforcement learning with differential learning, where the ‘reset’ is equivalent to a very large negative rate of decay. Therefore, we further tested the richness of the actual INTEGRATE-AND-RESET model family instantiated by M2 by decoding sequences generated with analog ‘g’. We found that M2 could also represent leaky integration of rewards and failures, and even amplification with small positive feedback ($g(o_{t+1}) < 1.2$ (Fig. 8d, right). Comparing across this parameter space (Fig. 8e), we observed that M2 had a preferred mode of integration that consisted of mostly perfect integration of failures ($0.85 \leq g(o_{t+1} = 0) \leq 1$) and integration of rewards with a variety of time constants ($g(o_{t+1} = 1) \leq 1$). Altogether, our results show that M2 simultaneously represents a relatively large repertoire of computations that embody a variety of foraging DVs, potentially spanning a set of optimal strategies for environments with different dynamics for the latent state.

Discussion

We explored the capacity of several regions of the cortex to deploy different algorithms for generating a diversity of DVs. We studied this in the context of a foraging task whose solution required mice to process

streams of successful and unsuccessful foraging attempts executed over several seconds. We found that mice could use not one but a set of discrete processing strategies to time their decision to switch between foraging sites, and the LM-HMM framework revealed that mice often change strategies within a session. All the decision strategies could be well read out from populations of neurons in M2. Moreover, we found the set of potentially relevant DVs was implemented in parallel within the same neural populations in M2. Conversely, OFC did not appear to multiplex DVs, consistent with the idea that it may be specifically involved in the computations of the inference-based strategy⁸.

While ‘causal’ manipulations of M2 using optogenetic inactivation showed that M2 was important to the deployment of the inference-based strategy, we found that the neural availability of alternative DVs was nearly independent of the actual behaviorally deployed DV. Functionally, the ability of M2 to multiplex the computation of several DVs could allow the mice to rapidly explore and adapt behavior to dynamically changing environmental contingencies by simply modifying linear readouts of M2 neural populations^{23,24} without the need to implement new computations.

The different DVs in M2 were ‘mixed’ but could be recovered through linear decoding. Although multiplexed neural codes have been observed previously in other cortical regions^{15,25–28}, our results establish that the kind of information that is multiplexed is not limited to representations of instantaneously observable events in premotor regions but also includes temporally extended computations spanning several seconds. While the observation of multiplexed DVs is reminiscent of the framework of ‘reservoir’ computing^{15–17,29}, we found that M2’s coding capacity was not universal, and instead implemented a substantial but circumscribed pool of potentially meaningful computations. One computation is accumulation of evidence, which, through its intimate relationship with posterior beliefs^{30,31}, constitutes an essential computation for statistical inference and

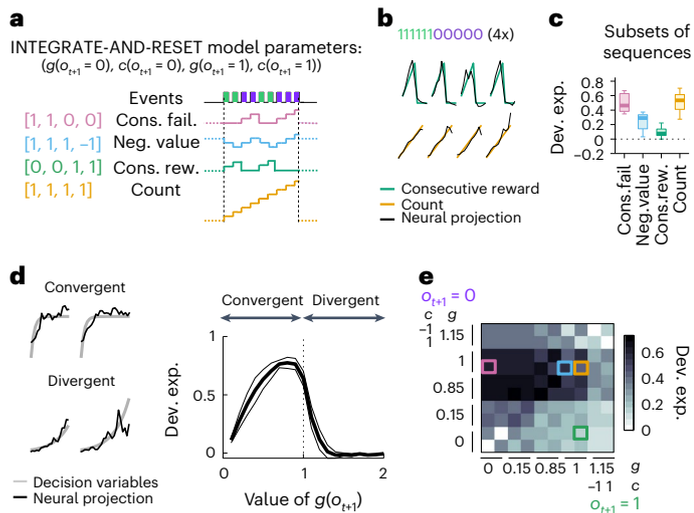


Fig. 8 | M2 represents foraging algorithms. **a**, The INTEGRATE-AND-RESET model with four parameters generates different time series by accumulating, resetting or ignoring each possible event (reward or failure). In the simplest instantiation of this model, the two outcome-dependent parameters are discrete: one is a gain factor (g) that specifies whether the running count should be reset or accumulated by each outcome—a nonlinear operation—and the other (c) specifies how each outcome linearly contributes to the resulting running count, which in general could be positive, negative or zero (leaving it unaffected; see Methods for more details). Each specification of these two discrete parameters leads to a different DV example set of parameters yielding example DVs on the right. **b**, Four example bouts (columns) of population activity (black traces) projected onto the dimensions that best predict the trajectory of the 'consecutive rewards' (green) and 'count' (yellow). Only subsequences of consecutive rewards followed by consecutive failures were selected to highlight the computations underlying the different variables. **c**, Deviance explained across sessions ($n = 11$ sessions, median \pm 25th and 75th percentiles, the whiskers extend to the most extreme data points) of the four basis sequences decoded from M2 population activity. The sequences were decorrelated using the same method as in Fig. 6c,d. Two-sided Wilcoxon signed rank test: $P = 0.002$ for 'consecutive reward' and $P = 0.00098$ for 'count'. **d**, Left, example sequences (gray) produced by analog parameters (convergent: $c(o_{t+1} = 0) = c(o_{t+1} = 1) = 1$ and $g(o_{t+1} = 0) = g(o_{t+1} = 1) = 0.5$; divergent: $c(o_{t+1} = 0) = c(o_{t+1} = 1) = 1$ and $g(o_{t+1} = 0) = g(o_{t+1} = 1) = 1.15$). Black traces are the neural projection from M2 population activity. Right, deviance explained from decoding convergent and divergent integrations by M2 population activity ($n = 11$ sessions, median \pm MAD). Here we show an example where the parameters of the INTEGRATE-AND-RESET model are as follows: $c(o_{t+1} = 0) = c(o_{t+1} = 1) = 1$ and $g(o_{t+1} = 0) = g(o_{t+1} = 1) = 1$. **e**, Matrix of deviance explained from decoding sequences with different time constants (corresponding to different values of g) of integrations of rewards (columns) and failures (rows) with M2 population activity. The basis sequences are indicated by the color-coded squares.

has therefore been implicated in a variety of decision-making and reasoning tasks^{22,32–36}. Accumulation (possibly temporally discounted) of action outcomes also underlies several reinforcement-learning algorithms^{37–40}. Although less attention has been devoted to reset-like computations (but see ref. 41), they are also essential for inference when certain observations specify a state unambiguously⁸.

The two strategies that we describe in the context of foraging represent a particular example of a more general phenomenon. In complex environments, agents can adapt their behavior in different ways depending on how accurately they can infer and specify the relevant causal structure⁴², a process that can be described as finding the correct 'task representation'. Even if unable to apprehend the true causal model, agents can display reasonably well-adapted behavior by leveraging the predictive power of salient environmental events. However, because the task representation is not correct, the association between these events and outcomes will necessarily be more probabilistic from the point of view of the agent. Such agents incorrectly model outcome

variance arising from incomplete task representations as unexplainable, and often resort to exploratory strategies that are adaptive in what they construe as highly volatile environments^{43–45}. Our results suggest that, at least in the case of foraging, the computations necessary to implement strategies lying along this continuum are computed simultaneously and available, which might facilitate the process of 'insight' necessary to switch between them.

Our finding also speaks to the debate on the nature of serial processing limitations in the brain. While it has been shown that limitations apply in some kinds of evidence accumulation tasks^{2,4,46}, here we show in a different, but ethologically important, setting that some forms of evidence accumulation can proceed in parallel. An important difference between our task and standard behavioral paradigms that study cognitive bottlenecks is that our mice do not need to simultaneously compute two DVs to perform the task successfully. Nevertheless, we show that neural populations in the premotor cortex of mice using a strategy where a single reward resets a counter of failures, reveal both this reset and simultaneously the updating of a reward counter. Our findings are thus consistent with proposals favoring parallel integration^{47,48} and with models that place serial constraints on behavior close to the specification of the timing of action^{47,49}.

Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41593-023-01305-8>.

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Methods

Data and processing

Animal subjects. A total of 27 adult male and female mice (24 C57BL/6J and 6 VGAT, 2–9 months old) were used in this study. All experimental procedures were approved and performed in accordance with the Champalimaud Centre for the Unknown Ethics Committee guidelines and by the Portuguese Veterinary General Board (Direco-Geral de Veterinaria, approval 0421/000/000/2016). Mice were housed in individually ventilated cages under a normal 12 h light/12 h dark cycle, temperature was maintained between 19 °C and 23 °C and humidity between 50% and 65%. During training and recording, mice were water-restricted (starting 5–10 d after head-bar implantation), and sucrose water (10%) was available to them only during the task. Mice were given 1 ml of water or 1 g of hydrogel (clear H₂O) on days when no training or recording occurred or if they did not receive enough water during the task.

Surgery and head fixation. All surgeries used standard aseptic procedures. Mice were deeply anesthetized with 4% isoflurane (by volume in O₂) and mounted in a stereotaxic apparatus (Kopf Instruments). Mice were kept on a heating pad and their eyes were covered with eye ointment (Vitaminofalmina A). During the surgery, the anesthesia levels were adjusted between 1% and 2% to achieve 1 s⁻¹ breathing rate. The scalp was shaved and disinfected with 70% ethanol and betadine. Carprofen (nonsteroidal anti-inflammatory and analgesic drug, 5 mg kg⁻¹) was injected subcutaneously. A flap of skin (less than 1 cm²) was removed from the dorsal skull with a single cut and the skull was cleaned and dried with sterile cotton swabs. The bone was scraped with a delicate bone scraper tool and covered with a thin layer of cement (C&B Super-Bond). Four small craniotomies were drilled (HM1 005 Meisinger tungsten) between Bregma and Lambda (around -0.5 and -1 AP; ±1 ml) and four small screws (Antrin Miniature Specialities, 000-120 × 1/16) previously soaked in 90% ethanol, were inserted in the craniotomies to stabilize the implant. The head bar (stainless steel, 19.1 × 3.2 mm), previously soaked in 90% ethanol, was positioned directly on top of the screws. Dental cement (Tab 2000 Kerr) was added to fix the head bar in position and to form a well around the frontal bone (from the head bar to the coronal suture). Finally, an external ground for electrophysiological recording (a male pin whose one extremity touched the skull) was cemented onto the head bar.

Behavioral apparatus. Head-fixed mice were placed on a linear treadmill with a 3D-printed plastic base and a conveyor belt made of Lego small tread links. The running speed on the treadmill was monitored with a microcontroller (Arduino Mega 2560), which acquired the trace of an analog rotary encoder (MAE3 Absolute Magnetic Kit Encoder) embedded in the treadmill. The treadmill could activate two movable arms via coupling with two motors (Digital Servo motor Hitec HS-5625-MG). A lick-port, made of a cut and polished 18G needle, was glued at the extremity of each arm. Water flowed to the lick-port by gravity through water tubing and was controlled by calibrated solenoid valves (Lee Company). Licks were detected in real-time with a camera (Sony PlayStation 3 Eye Camera or FLIR Chameleon-USB3) located on the side of the treadmill. Using BONSAI⁵⁰, an open-source visual programming language, a small, squared region of interest was defined around the tongue. To detect the licks, a threshold was applied to the signal within the region of interest. The behavioral apparatus was controlled by microcontrollers (Arduino Mega 2560) and scientific boards (Champalimaud Hardware platform), which were responsible for recording the time of the licks and the running speed on the treadmill, and for controlling water-reward delivery and reward depletion according to the statistics of the task.

Task design. In the foraging task, two reward sites, materialized by two movable arms, could be exploited. Mice licked at a given site to obtain liquid reward and decided when to leave the current site to explore the

other one. Each site could be in one of the following two states: 'ACTIVE', which is delivering probabilistic reward, or 'INACTIVE', which is not delivering any reward. If one of the sites was 'ACTIVE', the other one was automatically 'INACTIVE'. Each lick at the site in the 'ACTIVE' state yielded reward with a probability of 90% and could cause the state to transition to 'INACTIVE' with a probability of 30%. Licks could trigger the state of the exploited site to transition from 'ACTIVE' to 'INACTIVE', but never the other way around. Notably, this transition was hidden from the animal. Therefore, mice had to infer the hidden state of the exploited site from the history of rewarded and unrewarded licks (that is, rewards and failures). We defined 'behavioral bout' as the sequence of consecutive licks at one spout. A tone (150 ms, 10 kHz) was played when one of the arms moved into place (that is, in front of the mouse) to signal that a bout could start. At the tone, the closed loop between the motors and the treadmill decoupled during 1.5 s or until the first valid lick was detected. During this time, mice had to 'STOP', that is, decrease their running speed for more than 250 ms below a threshold for movement (6 cm s⁻¹). Licks were considered invalid if they happened before 'STOP' or at any moment after 'STOP' if the speed was above the threshold. If a mouse failed to 'STOP', 'LEAVE' was triggered by reactivating the closed loop after 1.5 s, which activated the movement of the arms (the one in front moved away and the other moved into place). Mice typically took around 200 ms to 'STOP' and initiate valid licking. During the licking periods, each lick was rewarded in a probabilistic fashion by a small drop of water (1 µl). The small reward size ensured that there was no strong difference in licking rate between rewarded and unrewarded licks. To 'LEAVE', mice had to restart running above the threshold for movement for more than 150 ms and travel a fixed distance on the treadmill (around 16 cm) to reach the other arm. We defined as correct bouts the ones in which mice stopped licking after the states transitioned from 'ACTIVE' to 'INACTIVE'. Error bouts were ones in which mice stopped licking before the state transition occurred. In this case, mice had to travel double the distance to get back to the arm in 'ACTIVE' state. Missed bouts were ones in which mice alternated between arms without any valid lick. These 'missed bouts' were excluded from our analysis.

Mouse training. Mice were handled by the experimenter from 3 to 7 d, starting from the beginning of the water restriction and before the first training session. At the beginning of the training, mice were acclimated to the head fixation and to the arm movement and received liquid reward simply by licking at the lick-port. The position of the lick-ports relative to the snout of the mouse had an important effect on behavioral performances. Thus, to ensure that the position of the lick-ports remained unchanged across experimental sessions, it was carefully adjusted on the first session and calibrated before the beginning of every other session. There were no explicit cues that allow discriminating between the two arms, and it was not even necessary that the animal be fully aware of the two different arms to perform the task. After mice learned to lick for water reward (typically after one or two sessions), the next sessions consisted of an easier version of the task (with a low probability of state transition, typically 5% or 10%, and high probability of reward delivery, 90%), and both arms in 'ACTIVE' state. That way, if mice alternated between arms before the states of the sites transitioned, the other arm would still deliver reward and animals would not receive the travel penalty. Occasionally, during the early phase of training, manual water delivery was necessary to motivate the mice to lick or stop running. Alternatively, it was sometimes necessary to gently touch the tail of the animals, such that they started to run and gradually associated running with the movement of the arms. The difficulty of the following sessions was progressively increased by increasing the probability of state transition if the performance improved. Performance improvement was indicated by an increase in the number of bouts and licking rate, and by a decrease in the average time of different events within a bout. Mice were then trained for at least five consecutive days on the

final task (90% reward delivery, 30% chance of state transition) before the recording sessions. The reason for choosing these statistics is that they correspond to a level of environmental uncertainty that is relatively low. This allows the mice to learn the task faster than at a high level of uncertainty and to remain highly motivated during the recording sessions, thus yielding a large number of behavioral bouts.

Electrophysiology. Recordings were made using electrode arrays with 374 recording sites (Neuropixels 'Phase3A'). The Neuropixels probes were mounted on a custom 3D-printed piece attached to a stereotaxic apparatus (Kopf Instruments). Before each recording session, the shank of the probe was stained with red-fluorescent dye (Dil Vybrant; Thermo Fisher, V22885) to allow later track localization. Mice were habituated to the recording setup for a few days before the first recording session. Before the first recording session, mice were briefly anesthetized with isoflurane and administered a nonsteroidal analgesic (carprofen) before drilling one small craniotomy (1 mm diameter) over the secondary motor cortex. The craniotomy was cleaned with a sterile solution and covered with silicone sealant (Kwik-Sil, World Precision Instruments). Mice were allowed to recover in their home cages for several hours before the recording. After head fixation, the silicone sealant was removed, and the shank of the probe was advanced through the dura and slowly lowered to its final position. The craniotomies and the ground pin were covered with a sterile cortex buffer. The probe was allowed to settle for 10–20 min before starting recording. Recordings were acquired with SpikeGLX Neural recording system (<https://billkarsh.github.io/SpikeGLX/>) using the external reference setting and a gain of 500 for the AP band (300 Hz high pass filter). Recordings were made from either hemisphere. The target location of the probe corresponded to the coordinates of the anterior lateral motor cortex; a region of the secondary motor cortex important for motor planning of licking behavior¹¹. The probe simultaneously traversed the OFC, directly ventral to the secondary motor cortex and the probe tip ended in the OLF, ventral to the OFC.

Histology and probe localization. After the recording session, mice were deeply anesthetized with ketamine/xylazine and perfused with 4% paraformaldehyde. The brain was extracted and fixed for 24 h in paraformaldehyde at 4 °C, and then washed with 1% phosphate-buffered saline. The brain was sectioned at 50 μm , mounted on glass slides and stained with DAPI. Images were taken at $\times 5$ magnifications for each section using a Zeiss AxioImager at two different wavelengths (one for DAPI and one for Dil). To determine the trajectory of the probe and approximate the location of the recording sites, we used SHARP-Track⁵¹, an open-source tool for analyzing electrode tracks from slice histology. First, an initial visual guess was made to find the coordinates from the Allen Mouse Brain Atlas (3D Allen CCF, http://download.alleninstitute.org/informatics-archive/current-release/mouse_ccf/annotation/) for each Dil mark along the track by comparing structural aspects of the histological slice with features in the atlas. Once the coordinates were identified, slice images were registered to the atlas using manual input and a line was fitted to the Dil track 3D coordinates. As a result, the atlas labels along the probe track were extracted and aligned to the recording sites based on their location on the shank. Finally, we also used characteristic physiological features to refine the alignment procedure (that is, clusters of similar spike amplitude across cortical layers, low spike rate between frontal and olfactory cortical boundaries or LFP signatures in deep olfactory areas).

Optogenetic stimulation. To optically stimulate ChR2-expressing VGAT-expressing GABAergic interneurons, we used blue light from a 473-nm laser (LRS-0473-PFF-00800-03, Laserglow Technologies, Toronto, Canada, or DHOM-M-473-200, UltraLasers, Newmarket, Canada). Light was emitted from the laser through an optical fiber patch-cord (200 μm , 0.22 NA, Doric lenses), connected to a second

fiber patch-cord with a rotatory joint (FRJ 1 \times 1, Doric lenses), which in turn was connected to the chronically implanted optic fiber cannulas (M3 connector, Doric lenses). The cannulas were inserted bilaterally inside small craniotomies performed on top of M2 (+2.5 mm anterior and ± 1.5 mm lateral of bregma) and barely touched the dura (as to avoid damaging superficial cortical layers). Structural glue (Super-bond C&B kit) was used to fix the fiber to the skull. The power of the laser was calibrated before every session using an optical power meter kit (Digital Console with Slim Photodiode Sensor, PM100D, Thorlabs). During the foraging task, the optical stimulation (10-ms pulses, 75 s⁻¹, 5 mW) was turned on during 30% of randomly interleaved bouts. Light delivery started when the first lick was detected and was interrupted if the animal did not lick for 500 ms (which was in 98% of bouts after the last lick of the bouts).

Preprocessing neural data. Neural data were preprocessed as described previously⁵². Briefly, the neural data were first automatically spike-sorted with Kilosort2 (<https://github.com/MouseLand/Kilosort>) using MATLAB (MathWork). To remove the baseline offset of the extracellular voltage traces, the median activity of each channel was subtracted. Then, to remove artifacts, traces were 'common average referenced' by subtracting the median activity across all channels at each time point. Second, the data were manually curated using an open-source neurophysiological data analysis package (Phy: <https://github.com/kwikteam/phy>). This step consisted in categorizing each cluster of events detected by a particular Kilosort template into a good unit or an artifact. There were several criteria to judge a cluster as noise (nonphysiological waveform shape or pattern of activity across channels, spikes with inconsistent waveform shapes within the same cluster, very low-amplitude spikes and high contamination of the refractory period). Units labeled as artifacts were discarded in further analyses. Additionally, each unit was compared to spatially neighboring units with similar waveforms to determine whether they should be merged, based on cross-correlogram features and/or drift patterns. Units passing all these criteria were labeled as good and considered to reflect the spiking activity of a single neuron. For all analyses, otherwise noted, we averaged for each neuron the number of spikes into bins by considering a 200 ms window centered around each lick. The bin vectors were then z-scored. Because the interval between each lick was on average around 150 ms, there was little overlap between two consecutive bins and each bin typically contained the number of spikes associated with only one lick.

Statistics and reproducibility

No statistical method was used to predetermine the sample size. The number of mice ($n = 27$ total) was chosen based on the current standard used for mice in neuroscience for sufficient statistical power. Randomization was used for stimulus presentation in all the experiments. The experimenter was blind to the genotype of the mice when performing optic fiber implantations and running the optogenetics experiments. For other experiments, no comparison across groups was made, therefore blinding was not necessary.

Statistical analysis of optogenetic manipulations. The statistical analysis of optogenetics was performed using generalized linear mixed-effect models, allowing us to pool different sessions of different mice in the same model. Our N is thus the number of mice multiplied by the number of sessions and conditions (Laser OFF/ON). The different groups (control versus inactivated) had different numbers of mice and sessions, which are reported in the results section. For each group, we fitted models with fixed effects of stimulation and random intercepts and effects of stimulation depending on mouse identity and session. For each mixed model, we report the coefficient of the fixed effect of the stimulation predictor (Laser) \pm s.d. of the estimate. We also report the P value that corresponds to the t -statistic for a hypothesis test that the coefficient of the 'Laser' predictor is equal to 0.

To describe mixed models, we use the Wilkinson notation, with $|$ denoting random effects. For example, the formula:

$$\text{TimeLicking} \sim 1 + \text{Laser} + (1 + \text{Laser}|\text{Mouse}) + (1 + \text{Laser}|\text{Session}),$$

uses as predictors for the time spent licking at a foraging site a constant intercept, a coefficient for 'Laser ON' condition that is different from 'Laser OFF' condition, which is considered as baseline, a random intercept across mice and a random intercept across sessions.

To test the strength of the effect of stimulation on the DVs in each single session, we generated 1,000 resamples of behavioral bouts in each 'Laser OFF' versus 'Laser ON' condition and used independent GLMs to predict the switching decision from the DVs for each resample. We compared the deviance explained of the models and the explained variance by each DV in 'Laser OFF' versus 'Laser ON' condition and estimated the significance of the differences. In the plot in Fig. 3, the sessions where P value < 0.05 are indicated by filled dots.

Predicting choice from DVs. All data analyses were performed with custom-written software using MATLAB. We used logistic regression⁵³ to estimate how DVs predicted the choice of the animal (that is, the probability that the current lick is the last in the bout). Using *Glmnet* for Matlab (http://www.stanford.edu/~hastie/glmnet_matlab/) with binomial distribution, model fits were performed with DVs as predictors. We used fivefold nested cross-validation and elastic net regularization ($\alpha = 0.5$). To assess a metric of model fit, we calculated the deviance explained (as implemented by the *devianceTest* function in Matlab). The deviance explained is a global measure of fit that is a generalization of the determination coefficient (r squared) for GLM. It is calculated as

$$\text{Deviance explained} = 1 - \frac{\text{residual deviance}}{\text{null deviance}}.$$

The residual deviance is defined as twice the difference between the log-likelihoods of the perfect fit (that is, the saturated model) and the fitted model. The null deviance is the residual deviance of the worst fit (that is, the model that only contains an intercept). The log-likelihood of the fitted model is always smaller than the log-likelihood of the saturated model, and always larger than the log-likelihood of the null model. Therefore, if the fitted model does better than the null model at predicting choice, the resulting deviance explained should be between 0 and 1. When the fitted model does not predict much better than the null model, the deviance explained is close to zero.

Simulated behavior sessions. To test the logistic regression model, we simulated behavioral sessions of an agent making decisions using a logistic function and the DV of the inference strategy (consecutive failures). For each simulated session, the slope and the intercept of the logistic regression in the ground truth model were chosen to fit the distribution of the total number of licks in each bout from the real data. To estimate the parameters of the ground truth model (slope and intercept), we then fit a logistic regression model to predict the leaving decisions of this simulated agent using the consecutive failures DVs.

Predicting DVs from neural population. We used a generalized LM for Poisson response⁵⁴ to predict each DV given the activity of the neural population (or facial motion, or both). Specifically, we predicted the DV A given the neural activity x , by learning a model with parameters, β , such as $A = \exp(\beta_0 + \beta x)$. The Poisson regression with log-link is appropriate to model count data like the DVs studied here. To enforce the positivity of the count responses, we shifted all the DVs to have a minimum value of one. Model fits were performed on each session separately. We employed elastic net regularization with parameter $\alpha = 0.5$. Additionally, we performed a cross-validation implemented by *cvglmnet* using the *lambda_min* option to select the hyperparameter

that minimizes prediction error. To assess the predictive power of the model, we also implemented nested cross-validation. Specifically, the model coefficients and hyperparameters were sequentially fit using a training set consisting of four-fifths of the data, and the prediction was evaluated on the testing set consisting of the remaining one-fifth. The method was implemented until all the data had been used both for training and testing. The deviance explained reported as a metric of the goodness of fit was calculated from the cross-validated results. The final β coefficients were estimated using the full dataset.

Comparison between brain regions. To ensure a fair comparison between brain regions with different numbers of recorded neurons, we excluded regions with very low numbers of recorded neurons (that is less than 20 neurons, $n = 2$ recordings in OLF excluded) and used multiple approaches to match the data from each region. One approach was to run the principal component analysis of the neural data from each region and select the principal components of neural activity that predicted up to 95% of the total variance (as reported in Fig. 2). A second approach was to select a subset of the original data to match the lowest number of neurons per region in each recording (subsampling with replacement, 100 repetitions). Both approaches yielded qualitatively similar results.

Predicting choice from neural population. We used logistic regression⁵³ to estimate how the weighted sum of neural activity (that is, the neural projections onto the weights that best predict the various DVs) predicted the probability that the current lick is the last in the bout. The model fits each recording session separately as described above using the *glmnet* package in MATLAB and implementing elastic net regularization with $\alpha = 0.5$ and a nested fivefold cross-validation to estimate the deviance explained.

Models

Integrate-and-reset. We developed a unified theory of integration in the setting of nonsensory decision-making tasks. In a wide variety of tasks, animals need to keep track of quickly evolving external quantities. Here we considered tasks where the feedback that the animal receives is binary (for example reward or failure). We considered an integrator given by $x_{t+1} = g(o_{t+1} = 1) \cdot x_t + c(o_{t+1} = 1)$, if the attempt is rewarded, and $x_{t+1} = g(o_{t+1} = 0) \cdot x_t + c(o_{t+1} = 0)$, otherwise. The parameters of the integrator $g(o_{t+1} = 0)$ and $g(o_{t+1} = 1)$ represent the computations and are bound between zero and one ($g = 1$ for an accumulation, $g = 0$ for a reset). The parameters $c(o_{t+1} = 1)$, $c(o_{t+1} = 0)$ add linearly and can be negative, positive or null.

We consider different scenarios involving a combination of computations but where the optimal solution only involves a one-dimensional integration. For instance, counting tasks can be solved by a linear integration, that is, $g(o_{t+1} = 0) = g(o_{t+1} = 1) = c(o_{t+1} = 0) = c(o_{t+1} = 1) = 1$, where the integrated value increases by one after each attempt regardless of the outcome. In a two-alternative forced choice and more generally in an n -armed bandit task, each arm would have an integrator that increases with rewards that is, $g(o_{t+1} = 0) = g(o_{t+1} = 1) = 1$, $c(o_{t+1} = 0) = 0$ and $c(o_{t+1} = 1) = 1$, and decays with failures, that is, $g(o_{t+1} = 0) = g(o_{t+1} = 1) = 1$, $c(o_{t+1} = 0) = -1$ and $c(o_{t+1} = 1) = 0$. Even in cognitively more complex tasks, involving inference over hidden states, such as reversal tasks or foraging under uncertainty, a single integrator is often sufficient. Specifically, in the foraging task studied here, the optimal solution is to integrate failures but not rewards, that is, $g(o_{t+1} = 0) = c(o_{t+1} = 0) = 1$, and $g(o_{t+1} = 1) = c(o_{t+1} = 1) = 0$.

More generally, the model produces sequences that ramp up with failures (that is, $g(o_{t+1} = 0) = c(o_{t+1} = 0) = 1$; such as the consecutive failures), and the mirror images that ramp down (that is, $g(o_{t+1} = 0) = 1$, $c(o_{t+1} = 0) = -1$). Similarly, the model can produce sequences that ramp up or down with rewards (that is, $g(o_{t+1} = 1) = 1$, $c(o_{t+1} = 1) = \pm 1$). The model also generates sequences that accumulate one type of event and

persist at a constant level with the other type (that is, $g(o_{t+1} = x) = 1$, $c(o_{t+1} = x) = \pm 1$, $g(o_{t+1} = y) = 1$, $c(o_{t+1} = y) = 0$), such as the cumulative reward integrator or its mirror image. Finally, many sequences generated by the model (where $g(o_{t+1} = 0) = g(o_{t+1} = 1) = 0$) track the outcomes (that is, reward versus failure).

There are 36 different values that the parameters of the model can take ($g(o_{t+1} = 0)$ and $g(o_{t+1} = 1)$ could take the values of 0 or 1 and $c(o_{t+1} = 0)$ and $c(o_{t+1} = 1)$ could take the values of -1, 0 or 1). In principle, each of these defines a different model which generates a time series when fed with sequences of binary action outcomes. The 8 of them for which $c(o_{t+1} = 0) = c(o_{t+1} = 1) = 0$ are trivial (constant). Of the remaining 28, not all are linearly independent. For instance, the time series generated by the model that computes ‘count’ ($g(o_{t+1} = 0) = g(o_{t+1} = 1) = c(o_{t+1} = 0) = c(o_{t+1} = 1) = 1$) is equal to the sum of the time series generated by the model that accumulates reward and is insensitive to failures ($g(o_{t+1} = 0) = g(o_{t+1} = 1) = 1$; $c(o_{t+1} = 0) = 0$; $c(o_{t+1} = 1) = 1$) and the time series generated by the model that accumulates failures and is insensitive to rewards ($g(o_{t+1} = 0) = g(o_{t+1} = 1) = 1$; $c(o_{t+1} = 0) = 1$; $c(o_{t+1} = 1) = 0$). Thus, the rank of the space of time series is 8 (two dimensions for the linear component (c) of the model for each of the four possible combinations of the g parameters, which specify the ‘computation’ the model is performing). Of these 8 dimensions, 4 come from models that are less interesting. Two of these are the two ‘outcome’ time series ($g(o_{t+1} = 0) = g(o_{t+1} = 1) = 0$), which are ‘observable’. We also only consider one time series for each of the two models, because the value of the linear component associated with the outcome that is reset makes very little difference to the overall shape of the time series. For instance, the time series generated by the two models $g(o_{t+1} = 0) = 1$; $g(o_{t+1} = 1) = 0$; $c(o_{t+1} = 0) = 1$; $c(o_{t+1} = 1) = 0$ and $g(o_{t+1} = 0) = 1$; $g(o_{t+1} = 1) = 0$; $c(o_{t+1} = 0) = 1$; $c(o_{t+1} = 1) = 1$ are linearly independent but almost identical for the type of outcome sequences of interest. The remaining 4 dimensions after these ‘trivial’ models are removed are spanned by the 4 basis elements that we focus on in the main text (Fig. 8). Finally, the effective dimensionality of the space of time series also depends on the temporal statistics of the outcome sequences. For the particular outcome sequences experienced by the mice (which are a function of the reward and state-transition probabilities) the effective dimensionality was low, which motivated us to focus on particular subsets of outcome sequences in Fig. 8 where the time series generated by the 4 basis elements are clearly distinct.

LM-HMM. To test the hypothesis that animals switch between discrete decision-making strategies within single sessions, we developed a new HMM with input-driven Gaussian observations modeling a time-varying linear dependence $\hat{F}_t = w^{(k)}\hat{R}_t + b^{(k)}$ of normalized consecutive failures \hat{F}_t (observations) on normalized total rewards \hat{R}_t (inputs) across bouts $t = 1, \dots, T$; ϵ_t is i.i.d. Gaussian noise with mean zero and variance $\sigma^{(k)}$. For each session m , the normalized values $\hat{F}_t = F_t/F_m^{\max}$ and $\hat{R}_t = R_t/R_m^{\max}$ were obtained by min-maxing the raw values F_t, R_t on their within-session $\max F_m^{\max}, R_m^{\max}$. This procedure allowed us to fit a single model to all sessions where both inputs and observations were bounded between zero and one. In this LM-HMM, the slope $w^{(k)}$, intercept $b^{(k)}$ and noise variance $\sigma^{(k)}$ depend on the hidden state $k = 1, \dots, K$, each state representing a different decision-making strategy. For example, states with $w^{(k)} = 0$ or $w^{(k)} > 0$ represent inference-based and stimulus-bound strategies, respectively. Large (small) values of the bias $b^{(k)}$ represent persistent (impulsive) behavior, respectively. Other model parameters include transition probabilities A_{kl} between hidden states and the initial state probabilities $\pi^{(k)}$. We fit an LM-HMM to bouts from all mice using the Expectation–Maximization (EM) algorithm to maximize the log-posterior and obtain the optimized parameters $\Theta = \{w^{(k)}, b^{(k)}, \sigma^{(k)}, A_{kl}, \pi^{(k)}\}$. Model selection for the number of states was performed using threefold cross-validation by concatenating all bouts from all sessions. A model was fit to the training set, and the log-posterior of the test set was estimated (normalized by the number of bouts per test set). Because the EM may lead to local maxima of the

log-posterior, for each choice of number of states, the EM algorithm was performed five times starting from random initial conditions. We performed model selection using maximum a posteriori (MAP, including Gaussian prior on the weights with variance equals to 2, and Dirichlet prior on transition probabilities with $\alpha = 2$; see ref. 14 for details on the procedure). The best number of states was chosen at the plateau of the maximum of the test MAP, leading to three states. We then fit a single model to the normalized observations and inputs \hat{F}_t, \hat{R}_t concatenating all bouts from all sessions, optimizing the model parameters Θ using MLE. Single-session values of weights and biases $w_m^{(k)}, b_m^{(k)}$ were then obtained from these normalized parameters $w^{(k)}, b^{(k)}$ as $w_m^{(k)} = w^{(k)} F_m^{\max}/R_m^{\max}$, $b_m^{(k)} = b^{(k)} F_m^{\max}$. The model was developed using Python 3.8.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

The preprocessed electrophysiological and behavioral data collected for this study are publicly and can be accessed at: <https://doi.org/10.6084/m9.figshare.20449089>.

Raw electrophysiological data are too large to be shared on a publicly available repository and are therefore available from the authors upon reasonable request.

The Allen Mouse Brain Atlas used in this study is publicly available: https://alleninstitute.github.io/AllenSDK/reference_space.html.

Code availability

All analyses were performed using custom code written in MATLAB and available upon request.

The code used for the central GLM analyses is publicly available at: https://hastie.su.domains/glmnet_matlab/.

The code developed for the LM-HMM can be accessed at: [https://github.com/mazzulab/ssm/blob/master/notebooks/2c%20Input-driven%20linear%20model%20\(LM-HMM\).ipynb](https://github.com/mazzulab/ssm/blob/master/notebooks/2c%20Input-driven%20linear%20model%20(LM-HMM).ipynb).

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Author contributions

F.C. and Z.F.M. conceived the project. F.C. and M.M. designed and performed behavioral experiments. J.P.M. and E.A. helped with surgery and behavioral training. F.C. designed and performed electrophysiological experiments. F.C. curated the data. F.C. and A.R. designed and performed the analyses. L.M. designed the LM-HMM. F.C., A.R. and Z.F.M. wrote the manuscript. All authors reviewed the manuscript.

Competing interests

All authors declare no competing interests.

Additional information

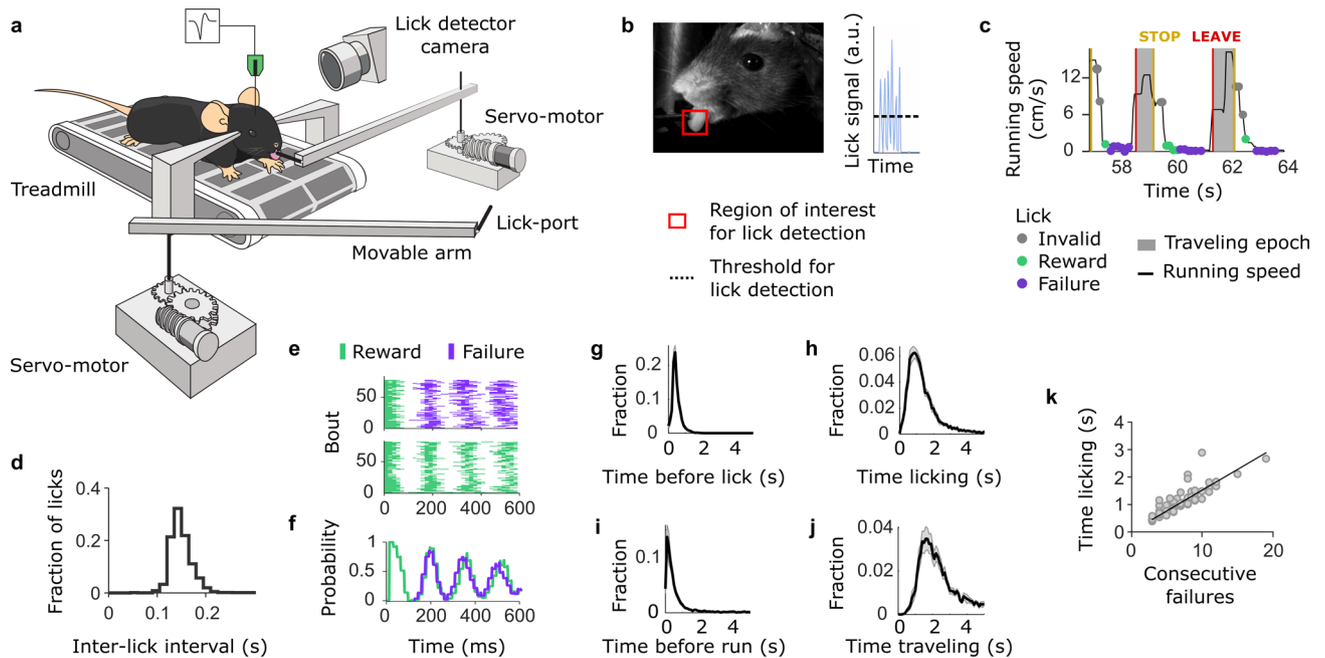
Extended data is available for this paper at <https://doi.org/10.1038/s41593-023-01305-8>.

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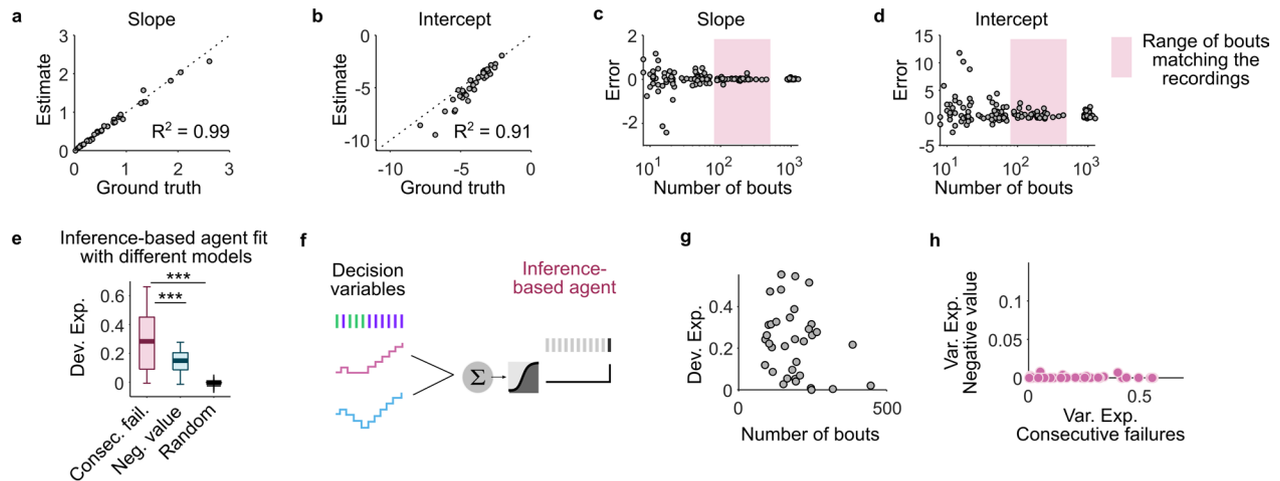
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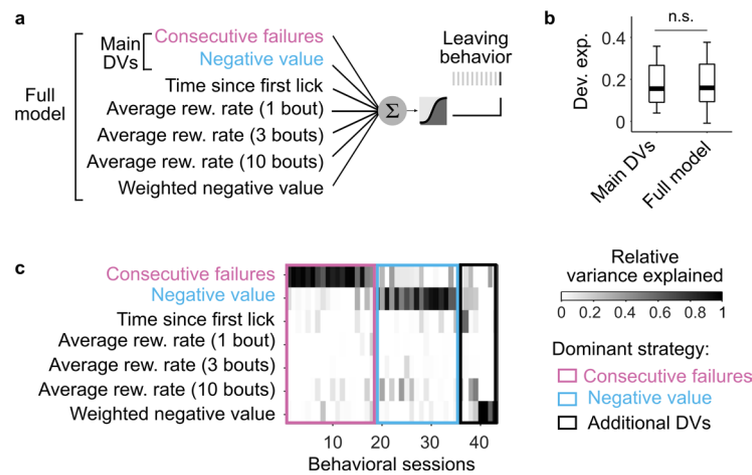
Extended Data Fig. 1 | Task apparatus and behavioral properties. (a) The behavioral apparatus consists of a treadmill, coupled to two motors. Rotating the treadmill activates in a closed-loop fashion the movement of the arms via the motors. A mouse placed on the treadmill with its head fixed can lick at the spout from the arm in front. A camera placed on the side of the animal allows on-line video detection of the licks. (b) View from the lick detector camera. A region of interest is defined around the tongue of the animal. To detect the licks a threshold is applied to the signal within the region of interest. (c) The task consists of behavioral bouts and traveling epochs. Within a behavioral bout, the outcomes of the licks are classified into three types: reward, failure and invalid. Rewards and failures occur when the mouse slows down its running speed below an arbitrary threshold after the 'STOP event'. The 'STOP event' is signaled by an auditory tone when an arm comes into place. Any lick above the running threshold is considered as invalid and always unrewarded. The traveling epoch starts after the 'LEAVE event' when the mouse initiates the run. (d, e, f) The licking

behavior of the animals is stereotyped. (d) Histogram of the time between each lick. (e) Examples of lick raster of consecutive failures (top) and consecutive rewards (bottom). Licks are aligned at the onset of a rewarded lick and sorted based on the following events. (f) The licking frequency that corresponds to the two different examples in (e) (series of consecutive rewards in green and series of consecutive failures in purple). (g, h, i, j) Time distributions of different behavioral events (mean \pm s.e.m.; $n = 21$ mice). The time spent licking was much greater than the time to initiate licking (between STOP event and first lick) or the time to initiate running (between the last lick and LEAVE event). Notably, engaged mice took less than half a second after the last licks to leave the site in most bouts (Median time to run = 0.46 s). The running time is comparable to the licking time. (k) Monotonic relationship between the number of consecutive failures after the last reward and the time licking after the last reward (each dot represents the means across bouts for each session).



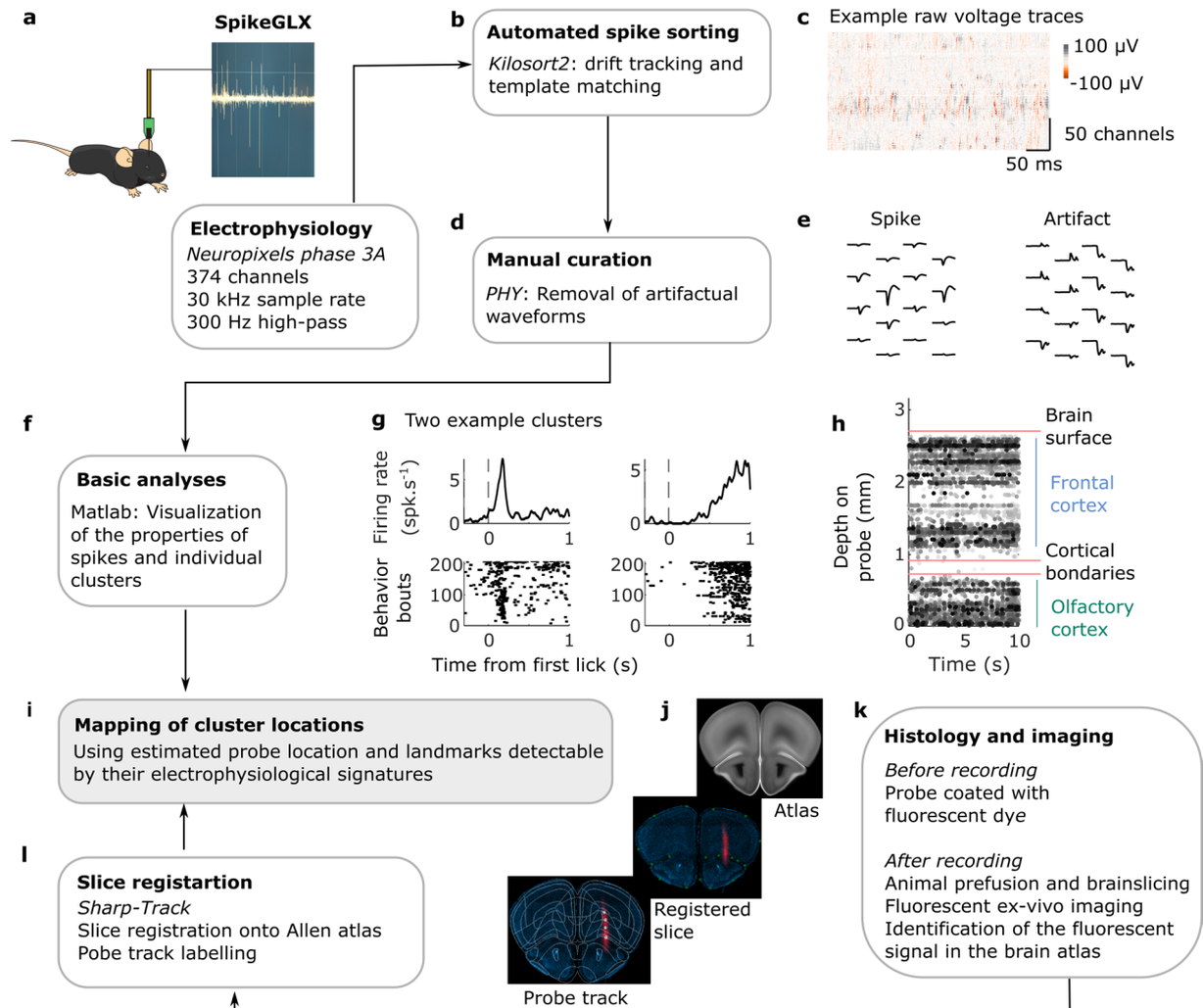
Extended Data Fig. 2 | Ground truth model. (a, b) The slope (a) and intercept (b) estimates as a function of the ground truth for simulated sessions where the number of bouts matched that of real sessions. The ground truth can be recovered ($R^2 = 0.99$ for the slope; $R^2 = 0.91$ for the intercept) from the logistic regression. (c, d) The slope (c) and intercept (d) estimates as a function of the ground truth for simulated sessions with varying number of bouts. Overall, the ground truth can be precisely recovered for sessions with more than 100 bouts. (e) Deviance explained from a logistic regression model that fits simulated sessions of an inference-based agent using the correct model ('Consecutive failures'), a wrong but correlated model ('Negative value') and a random model (where both rewards and failures are arbitrarily accumulated or reset). The deviance explained by the consecutive failures represents the upper-bound of the model performance. The deviance explained by the consecutive failures being smaller than 1 indicates that, although the ground truth can be recovered, the switching decision is not deterministic and involves some stochasticity (here

the variability was matched to that of the data). However, the deviance explained by the consecutive failures is significantly greater than the deviance explained by the correlated model and the random model (two-sided Wilcoxon signed rank test, 3 stars indicate $p < 10^{-3}$, $p = 0.00005$ between Consec. fail. and Neg. value; $p < 10^{-7}$ between Consec. fail. and Random). On each box the central mark indicates the median across simulated sessions ($n = 42$ sessions), and the bottom and top edges of the box indicate the 25th and 75th percentiles, respectively. The whiskers extend to the most extreme data points. (f) Illustration of a logistic regression model for predicting the switching decision of an inference-based simulated agent from the two different DVs ('Consecutive failures' and 'Negative value') simultaneously. (g) Deviance explained from the model in (f) as a function of the number of bouts in each session. (h) For all simulated sessions in (e), the variance explained by the 'consecutive failures' DV was greater than the variance explained by the 'negative value' DV, indicating that the model inferred the true DV.



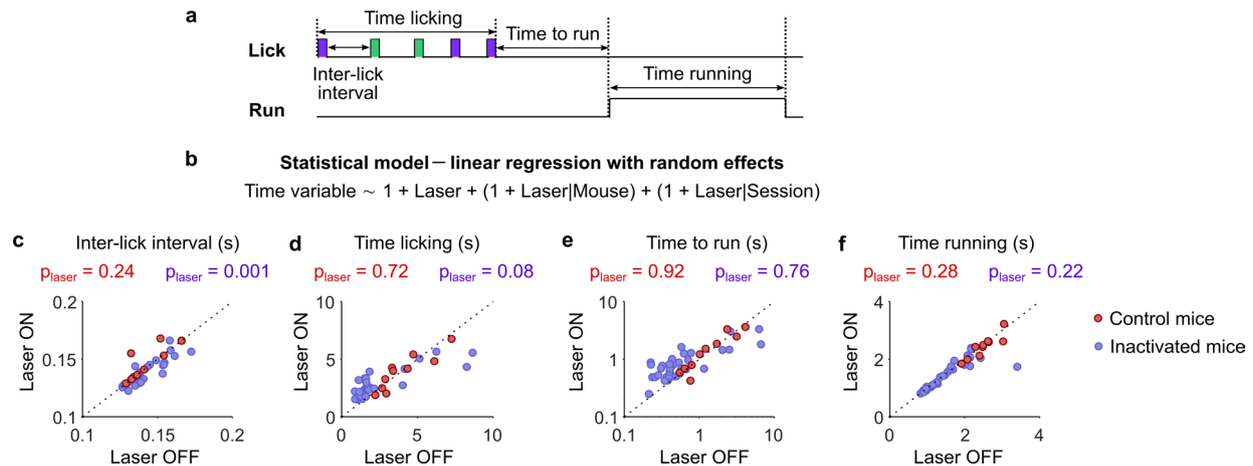
Extended Data Fig. 3 | Testing alternative foraging strategies. (a) Illustration of the logistic regression model for predicting the switching decision of mice using a combination of the two main DVs, ‘Consecutive failures’ and ‘Negative value’, as well as additional DVs. Specifically, we tested 3 classes of additional DVs: 1) those relying on absolute time, 2) those relying on average reward rates, and 3) those that weigh recent evidence more strongly. The design matrix of the model thus consisted of the two main DVs, the time of each lick relative to the first lick of each bout (class 1), the average reward rate over 1, 3 and 10 previous bouts (class 2) and a version of the negative value DV that weighs recent evidence more heavily than the past ones (for class 3), such as: $x_{t+1} = (1 - \alpha) \cdot g(o_{t+1}) \cdot x_t + \alpha \cdot c(o_{t+1})$, with $\alpha = 0.8$. (b) Deviance explained from a logistic regression model that predicts choice behavior based only on the 2 main DVs (left) and from the full model that also includes the additional DVs in (a). The central mark indicates

the median across behavioral sessions ($n = 42$ sessions), and the bottom and top edges of the box indicate the 25th and 75th percentiles, respectively. The whiskers extend to the most extreme data points. There was no significant difference between the deviance explained of the two models (two-sided Wilcoxon signed rank test: $p = 0.22$), indicating that the additional DVs do not improve the performance of the model. (c) Relative variance explained by each predictor of the full model for each behavioral session ($n = 42$ sessions across 21 mice, 2 sessions per mice). The dominant DV (the one with the maximum relative variance explained) was most often the ‘Consecutive failures’ (18 sessions), followed by the ‘Negative value’ (17 sessions), and finally the additional DVs (2 session for the absolute time, 2 sessions for average reward rate, 3 sessions for the weighted negative value).



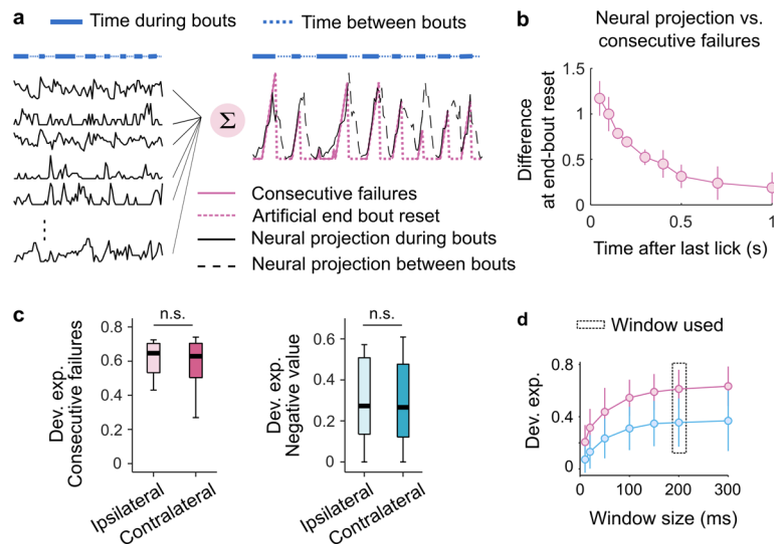
Extended Data Fig. 4 | Pipeline for extracellular electrophysiology, data processing and cluster mapping. (a) Data collection from the Neuropixels probe. (b) Kilosort2 is used to automatically match spike templates to raw data. (c) Example of voltage data input to Kilosort2. Prior to the automatic sorting, the raw data is preprocessed with offset subtraction, median subtraction, and whitening steps. (d) Manual quality control is done on the outputs of Kilosort2 using PHY to remove units with nonphysiological waveforms (e), contaminated refractory periods, low amplitude (less than 50 μV) or low spiking units (less than 0.5 spike $\cdot\text{s}^{-1}$). (f) For further quality control, visualization of peri-event spike

histograms (g, top; examples histogram aligned to first lick) or scatter plots (g, bottom; example scatter plot aligned to first lick) of single neurons are made with custom-written script in MATLAB. (h, i) Example scatter plot of all neurons recorded simultaneously along the shank of the probe. This visualization helps delimitate landmarks based on electrophysiological signatures to map cluster locations. (j, k, l) Landmarks derived from electrophysiological responses are validated with estimates from histology using an open-source software (SHARP-Track).



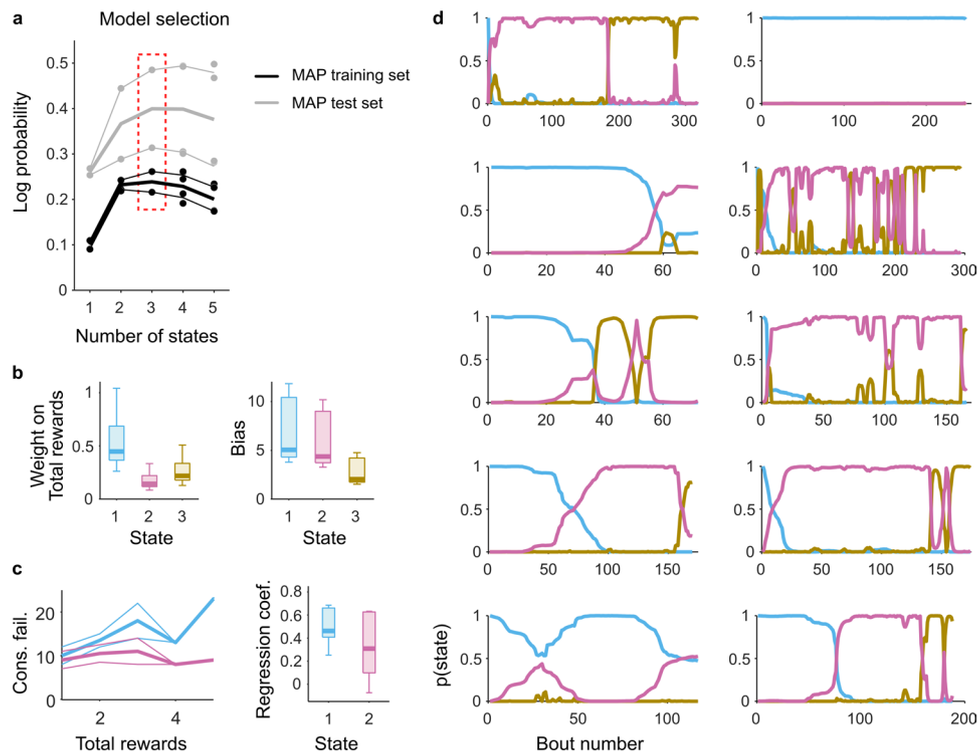
Extended Data Fig. 5 | Optogenetic effect on action timing. (a) Illustration of the different action timing during a behavioral bout. (b) We used generalized linear mixed effect models to evaluate the effect of stimulation ('Laser' predictor) on each action timing (see Methods). The models were fit separately for inactivated and control mice (number of observations: Inactivated = 68; Control = 20). (c–f) Median timing across bouts in Laser OFF vs. Laser ON condition for each session (dots) of inactivated mice (violet) and control mice (red) mice. The p-value corresponding to the t -statistic for a two-sided null hypothesis test that the coefficient of the 'Laser' predictor is equal to 0 (p_{laser}) is reported for each

group of mice (color coded). (c) Fixed effect of stimulation ('Laser' predictor) on the inter-lick interval: Inactivated: -0.003 ± 0.0009 , $p = 0.001$; Control: 0.005 ± 0.004 , $p = 0.24$. (d) Fixed effect of stimulation ('Laser' predictor) on the time licking: Inactivated: 0.45 ± 0.26 , $p = 0.08$; Control: -0.078 ± 0.22 , $p = 0.72$. (e) Fixed effect of stimulation ('Laser' predictor) on the time to run: Inactivated: -0.075 ± 0.25 , $p = 0.76$; Control: 0.014 ± 0.14 , $p = 0.92$. (f) Fixed effect of stimulation ('Laser' predictor) on the time running: Inactivated: -0.079 ± 0.063 , $p = 0.22$; Control: -0.061 ± 0.052 , $p = 0.28$.



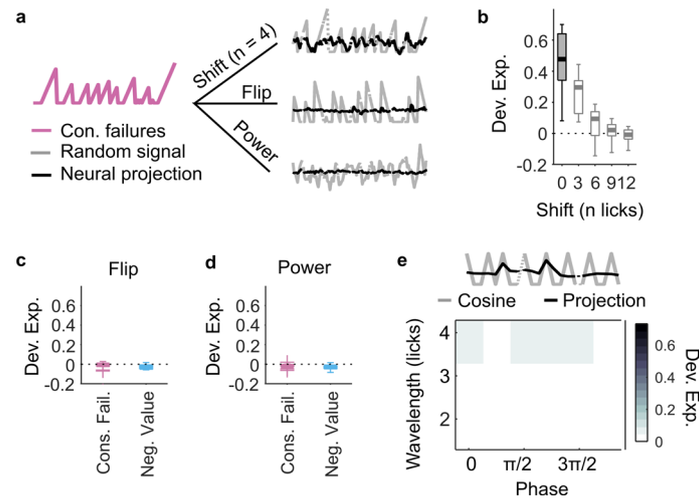
Extended Data Fig. 6 | Properties of decision variables in M2. (a) Illustration of a model to estimate the time constant of the reset at the end of the bout from M2 neurons. Example consecutive failures (pink) and neural projections (black right) of the neural activity (left, example neural traces) including the activity during 2 s after the end of each bout (dashed line). The projection of the neural activity on the decoding weights for the consecutive failure slowly ramps down until the beginning of the next bout. (b) To quantify the time constant of the reset at the end of the bout, the consecutive failures with an additional reset at the end of the bout were decoded from the neural activity. We considered the decoding projection at different times after the end of the last lick of bout 'n' and before the start of bout 'n + 1' and plotted the difference between the number of the consecutive failures (dashed pink) and the neural projection (dashed black) at the end of each bout across recording sessions (median \pm MAD; $n = 11$) as a function of the time after the last lick. The neural activity can reset at the end of the bouts with a time constant of around 200 ms. (c) Deviance explained across sessions ($n = 11$ sessions, median \pm 25th and 75th percentiles, the whiskers extend to the most extreme data points) predicted from M2 neurons for 'Consecutive failures' (left) and 'Negative value' (right) on ipsilateral vs. contralateral bouts. If the recording is performed in the right hemisphere, ipsilateral bouts are those when mice exploit the right foraging site (the right

motorized arm), while contralateral bouts are those when mice exploit the left foraging site (and vice versa for recordings in the left hemisphere). We observed no significant differences in the model performance as a function of the side of the DVs (two-sided Wilcoxon signed rank test; $p > 0.05$). (d) This panel shows the deviance explained across sessions ($n = 11$ sessions, median \pm MAD) for DVs (Pink: 'consecutive failures'; Blue: 'negative value') as a function of window sizes. In all previous analyses, the window used to count the spikes was 200 ms centered around each lick (indicated by the black rectangle), which was a good tradeoff for including a significant number of spikes while mainly considering signals related to a single lick (since the average time between each lick was around 150 ms; Fig. 2b & Extended Data Fig. 1d). Yet, a few spikes linked to the preceding or the following events could still be included in the 200 ms window, making it more difficult to evaluate the contribution of momentary evidence. Therefore, we tested whether both DVs remained decodable in M2 even when we strictly excluded all spikes from neighboring events by using smaller analysis windows. We found that the decodability of the DVs in M2 did not depend on the size of the window for widths larger than 20 ms (one-way ANOVA followed by multiple pairwise comparison tests, all p -values > 0.05 for windows size > 20 ms, both for 'consecutive failures' and 'negative value'), indicating that the results are not overly sensitive to the choice of parameters.



Extended Data Fig. 7 | LM-HMM analysis of switch decision. (a) To determine the number of states that best capture the decision-making of mice, we fit the LM-HMM with a varying number of states and then performed model comparison using cross-validation (see Methods for details). Training and test sets maximum a posteriori (MAP, with gaussian prior on the weights and Dirichlet prior on transition probabilities) are reported in units of bits per bout (median \pm MAD). The dash-line rectangle highlights the log probability for the three-state model, which we used for all subsequent analyses. A single model was fit to all mice, where for each session the consecutive failures and prior rewards were min-maxed (that is, divided by their max F_m^{max} , R_m^{max}), obtaining normalized weights $w^{(k)}$ and biases $b^{(k)}$. Single-sessions weights and biases were then obtained from these normalized parameters as $w_m^{(k)} = w^{(k)} F_m^{max} / R_m^{max}$, $b_m^{(k)} = b^{(k)} F_m^{max}$. (b) Weights $w_m^{(k)}$ on total reward (left) and biases $b_m^{(k)}$ (right) across sessions m ($n = 11$

sessions, median \pm 25th and 75th percentiles, the whiskers extend to the most extreme data points) in the different states $k = 1, 2, 3$. (c) Consecutive failures before leaving as a function of total reward number across behavioral bouts (median \pm MAD) in an example session from two different states (state 1, blue; state 2, pink). The slope coefficients of a linear regression model that predicted the number of consecutive failures before leaving as a function of the number of prior rewards in each state are shown on the right ($n = 6$ sessions for state 1, $n = 7$ sessions for state 2, median \pm 25th and 75th percentiles across sessions, the whiskers extend to the most extreme data points). This result is consistent with the classification of stimulus-bound and inference-based strategies used in Fig. 1. (d) Posterior state probabilities for each recording session. Mice often start off the session with the stimulus-bound strategy and later switch to the inference-based strategies (in 6 out of 11 sessions).



Extended Data Fig. 8 | M2 does not represent arbitrary sequences. (a) A ‘near universal’ representational capacity is a feature of a computational framework known as ‘reservoir computing’ that exploits a potential functional capacity of recurrent networks to represent combinations of current inputs with previous evidence, even arbitrary ones. Thus, to test whether M2 also represented arbitrary signals, we examined whether sequences with similar temporal structure as the DVs but with no obvious relevance to the task could be decoded from M2. Here are examples of random sequences (gray) generated from one of the DVs (pink, here consecutive failures). The DV can lead to a shifted version (top right), a flipped version (middle right) or a random signal with equal power spectra. Each random signal is then decoded from M2 population activity (black traces). (b) Deviance explained (ordinate) by M2 neurons from decoding the DVs shifted by a given number of licks (abscissa). On each box, the central mark indicates the median across recording sessions (n = 11 sessions), and the bottom and top edges of the box indicate the 25th and 75th percentiles, respectively. The whiskers extend to the most extreme data points. The dash black line

indicates chance level (Dev. Exp. = 0). Shifting the DVs by a delay greater than their temporal autocorrelation greatly impaired their decodability (one-way ANOVA, $F = 62.81$, $p < 10^{-26}$). (c) Same as in (b) but for DVs flipped across sessions. None of the flipped signals were decodable from M2 population activity. (d) Same as in (c) but for random signals with power spectra that match each DV. None of the random signals were decodable from M2 population activity. (e) Since any signal can be approximated by sums of periodic functions (Fourier analysis), we also probed the capacity of M2 to represent arbitrary temporal sequences by testing whether we could decode from M2 a basis set of cosine functions with wavelengths in the dynamic range of what we observed with integration and reset of rewards (example top gray trace; wavelength = 4 licks, phase = 0 rad). Overall, the decoding quality of the periodic function (example neural projection, top trace in black, Dev. Exp. = -0.002) was close to chance level (Dev. Exp. = 0.024 ± 0.028 , median \pm MAD) as seen in the matrix of deviance explained from decoding sequences with different wavelengths and phases with M2 population activity.

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Data analysis	MATLAB 2019b Python 3.8 Kilosort2 (https://github.com/MouseLand/Kilosort/releases/tag/v2.0) Phy (https://github.com/cortex-lab/phy) SHARP-Track (https://github.com/cortex-lab/allenCCF) Custom codes developed in the study (https://github.com/mazzulab/ssm/blob/master/notebooks/2c%20Input-driven%20linear%20model%20(LM-HMM).ipynb) Glmnet in Matlab (https://hastie.su.domains/glmnet_matlab/)

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Preprocessed electrophysiological and behavioral data have been deposited to Figshare and can be accessed at: <https://doi.org/10.6084/m9.figshare.20449089>
Raw electrophysiological data are too large (hundreds of GBs) to be shared on a publicly available repository and are therefore available from the authors upon reasonable request.

The Allen Mouse Brain Atlas used in this study is publicly available: https://alleninstitute.github.io/AllenSDK/reference_space.html

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Life sciences study design

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Sample size

Sample sizes were not predetermined. A group of animals (n = 11) was first highly trained over weeks to study the effect of behavior. A second group (n = 10) of animals was then trained for the electrophysiology and also included in the behavioral datasets. A third group of animal (n=6) was used for the optogenetics experiments. The number of mice (n= 27 total) was chosen based on the on the current standard used for mice in neuroscience for sufficient statistical power.

Data exclusions

For the electrophysiological datasets, sessions were included only if all the Neuropixels probe was located in the target regions by post-hoc histological analysis. Single units (neurons) spike clustering quality were assessed manually using Phy. Units were only included if firing rate did not drift over the recording session, and spikes did not violate absolute refractory period (see Method).

Replication

All relevant behavioral effects were replicated in 3 sets of independently trained animals (n = 11, n = 10 and n = 6). Effects reported in neural data were consistent across animals.

Randomization

Randomization was used for stimulus presentation. The task statistics were similar across sessions but probabilistic. Thus, the sequences of events during behavior bouts were naturally randomized by the probabilities and the length of behavior bouts depended on individual behavioral responses. Therefore, each session had a unique sequence of behavior bouts.

Blinding

The experimenter was blind to the genotype of the mice when performing optic fiber implantations and running the optogenetics experiments. For other experiments, no comparison across groups was made, therefore blinding was not necessary.

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Laboratory animals

Male and female C57BL/6J and VGAT mice (2-9 months old) were used in this study. Mice were housed in individually ventilated cages under a normal 12 hour light/dark cycle, temperature was maintained between 19-23°C and humidity between 50-65%.

Wild animals

This study did not involve wild animals.

Reporting on sex

Both male and female mice were used in this study.

Field-collected samples

This study did not involve field-collected samples.

Ethics oversight

All experimental procedures were approved and performed in accordance with the Champalimaud Centre for the Unknown Ethics Committee guidelines and by the Portuguese Veterinary General Board (Direco-Geral de Veterinaria, approval 0421/000/000/2016).

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