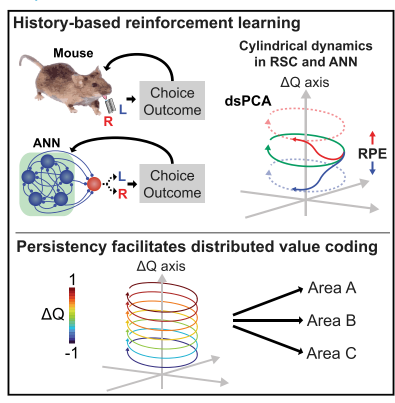
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Context-dependent persistency as a coding mechanism for robust and widely distributed value coding

Graphical abstract



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In brief

Hattori et al. showed that highly persistent value coding emerges in the mouse retrosplenial cortex and artificial recurrent neural network in a learning-and context-dependent manner.

Persistency ensures untangled value representations in neural population dynamics and facilitates distributed value coding across the brain, highlighting the benefits of persistent coding.

Highlights

- Coding persistency in the cortex is learning and context dependent
- Highly persistent value coding emerged in RSC and ANN during reinforcement learning
- Persistency ensures untangled value representation within cylindrical geometry
- Coding persistency facilitates brain-wide distributed information coding

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Context-dependent persistency as a coding mechanism for robust and widely distributed value coding

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SUMMARY

Task-related information is widely distributed across the brain with different coding properties, such as persistency. We found in mice that coding persistency of action history and value was variable across areas, learning phases, and task context, with the highest persistency in the retrosplenial cortex of expert mice performing value-based decisions where history needs to be maintained across trials. Persistent coding also emerged in artificial networks trained to perform mouse-like reinforcement learning. Persistency allows temporally untangled value representations in neuronal manifolds where population activity exhibits cyclic trajectories that transition along the value axis after action outcomes, collectively forming cylindrical dynamics. Simulations indicated that untangled persistency facilitates robust value retrieval by downstream networks. Even leakage of persistently maintained value through non-specific connectivity could contribute to the brain-wide distributed value coding with different levels of persistency. These results reveal that context-dependent, untangled persistency facilitates reliable signal coding and its distribution across the brain.

INTRODUCTION

The parallel distributed processing (PDP) theory (McClelland et al., 1986; Rogers and McClelland, 2014; Rumelhart et al., 1986) highlights computational advantages of distributed information coding in neural networks and has had a profound impact on our understanding of cognition and deep learning. Growing evidence revealed that information coding in the brain is highly distributed across neurons and distinct brain areas (Allen et al., 2019; Hattori et al., 2019; Koay et al., 2020; Musall et al., 2019; Steinmetz et al., 2019; Stringer et al., 2019). Even neurons in the primary sensory cortex, which were classically thought to process only sensory information of a single modality, have been found to encode diverse information such as other sensory modalities (Hattori and Hensch, 2017; Hattori et al., 2017; Iurilli et al., 2012), spontaneous movements (Musall et al., 2019; Stringer et al., 2019), actions (Hattori et al., 2019; Koay et al., 2020; Steinmetz et al., 2019), reward (Hattori et al., 2019; Koay et al., 2020), event history (Hattori et al., 2019; Koay et al., 2020), and value (Hattori et al., 2019; Serences, 2008). Although these signals are widely distributed, activity perturbations of a brain area typically affect only a subset of behavioral outputs that are associated with the information encoded in the area. These results suggest that, although information coding is highly distributed, not all of the information represented in neural activity may be used in each area.

A clue to understand the utility of encoded information may lie in the temporal dynamics of the information coding. In working memory tasks where information is maintained for several seconds in a trial, information can be maintained as either persistent neural activity or sequential transient activity across a neural population that tiles the memory period (Cavanagh et al., 2018; Fuster and Alexander, 1971; Masse et al., 2019; Miller et al., 1996; Murray et al., 2017; Orhan and Ma, 2019; Romo et al., 1999; Zhu et al., 2020). Recently, it was shown that certain brain areas in mice, such as the retrosplenial cortex (RSC) (Hattori et al., 2019) and the medial prefrontal cortex (Bari et al., 2019), encode action values with exceptional persistency during history-dependent, value-based decision-making tasks where values need to be stably maintained across trials. Inactivation of either area impaired the ability to use the action value for their decision making. These results suggest that persistent value coding is critical for animals to exploit value for decision-making when the value needs to be maintained for extended periods of time. Similar persistent coding is prevalent across the brain and species, ranging from coding of motor planning (Guo et al., 2017; Inagaki et al., 2019; Li et al., 2016) and internal states (Allen et al., 2019; Margues et al., 2020) to emotions (Jung et al., 2020; Kennedy et al., 2020), yet the computational advantages of persistent coding has not been fully established quantitatively.

Here we investigated the neural dynamics of action history and value coding in six areas of the mouse cortex and artificial



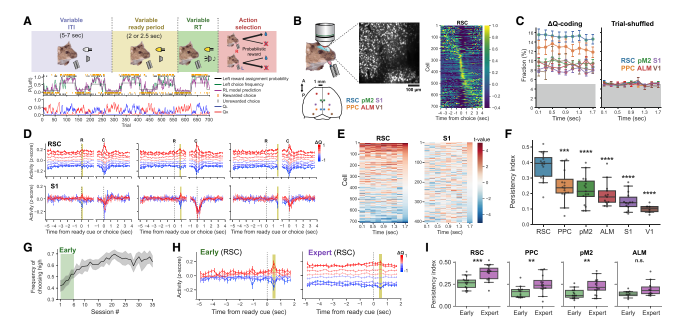


Figure 1. Persistency of action value coding across mouse cortex is area and learning dependent

(A) Schematic of the value-based decision task and an example expert behavior.

(B) Neural activity was recorded from 6 cortical areas. The heatmap is the trial-averaged z-scored deconvolved activity of an example RSC population. The activity of each neuron was normalized to its peak. A half of the recorded trials were used to sort cells by the peak time, and the mean activity of the other half are shown.

(C) Fractions of cells with significant ΔQ coding during ready period based on the mean activity within each of the non-overlapping 200 ms bins (Regression, p < 0.05, 2-sided t test). The fractions with filled circles are significantly above the chance fraction of 5% (p < 0.05, one-sided t test). ΔQ values were shuffled across trials for the right panel.

(D) Activity of ΔQ coding neurons that were identified at different time windows (yellow shadings) for example RSC and S1 populations. Trials were binned according to the ΔQ of each trial, and the activity in each trial bin was averaged.

(E) t-values for ΔQ coding at each time bin of ready period for example populations of RSC and S1 (Regression). Neurons were sorted based on the t-values at the

(F) ΔQ coding persistency of each population as quantified by the persistency index (0: chance persistency, 1: maximum-possible persistency, STAR Methods, ***p < 0.001, ****p < 0.0001, one-way ANOVA with Tukey's HSD).

(G) Fraction of trials when mice chose the side with higher reward assignment probability across training sessions (n = 9 mice, mean ± Cl). The first 6 sessions were treated as early sessions.

(H) Activity of ΔQ coding RSC neurons that were identified from the activity within the specified time bin (yellow shadings) in early and late sessions (same RSC population between the 2 sessions) indicating an increase in persistency during learning.

(I) $\Delta\Omega$ coding persistency of each population as quantified by the persistency index for early and expert sessions (**p < 0.01, ***p < 0.001, mixed effects model with population as the fixed intercept).

All error bars are SEM.

recurrent neural network (RNN) agents to understand the computational advantages of persistent coding and its impact on distributed coding.

RESULTS

Learning and context dependence of coding persistency across cortical areas

We first used the neural activity data recorded in mice performing decision-making based on history-dependent action value we reported previously (Hattori et al., 2019). Each trial consisted of a ready period, an answer period, and an intertrial interval (ITI). The duration of each period was variable from trial to trial, making the task more naturalistic than a fixed temporal sequence (Figure 1A). During the ready period (LED cue), mice needed to withhold licking to enter the answer period. This ensured that the neural activity during the ready period was free of lickingrelated motor activity. Mice were allowed to freely choose either left or right lickport after a go cue tone. Different reward probabilities were assigned to the 2 lickports, and the reward probabilities changed every 60-80 trials without cue. Therefore, mice were encouraged to dynamically estimate the underlying reward probabilities of the 2 options on a trial-by-trial basis by forming subjective action values based on their recent choice outcome history using reinforcement learning (RL) (Sutton and Barto, 2018). The action values need to be stably maintained within each trial and updated after each trial based on the action and its outcome. Neural activity was collected with in vivo 2-photon calcium imaging from transgenic mice that express GCaMP6s (Chen et al., 2013) in excitatory neurons (Wekselblatt et al., 2016) (Figure 1B), and the calcium signals were converted to estimated spike rates by non-negative deconvolution (Friedrich **Neuron Article**



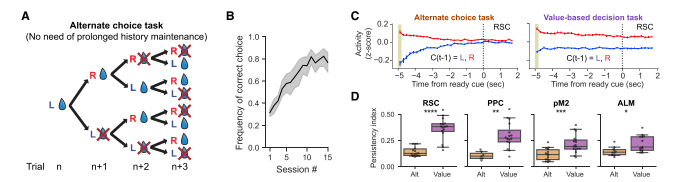


Figure 2. Persistency of history coding is task dependent

- (A) Schematic of the alternate choice task. The choice opposite to the choice in the previous trial was rewarded regardless of reward outcome in the previous trial. (B) Fraction of trials of correctly choosing the side with reward across training sessions (n = 9 mice, mean ± 95% CI).
- (C) Activity of RSC neurons that significantly encoded the action history from previous trial in alternate choice task and value-based decision task. These neurons were identified using the activity within the specified time bin (yellow shadings). The activity of the identified action history coding neurons was separately averaged according to the choice on the previous trial.
- (D) Persistency of action history coding in each population as quantified by the persistency index for the alternate choice task (Alt) and the value-based decision task (Value) (*p < 0.05, **p < 0.01, ***p < 0.001, ***p < 0.0001, mixed effects model with population as the fixed intercept).

et al., 2017; Pachitariu et al., 2018). The recording data were from 6 cortical areas including 2 association (RSC: retrosplenial; PPC: posterior parietal), 2 premotor (pM2: posterior secondary motor; ALM: anterior-lateral motor), and 2 primary sensory (S1: primary somatosensory; V1: primary visual) cortex. We estimated the 2 action values on each trial (QL and QR) by fitting an RL model to the choices of mice, and we focused our analyses on the neural coding of the policy value ($\Delta Q = Q_L - Q_R$: the value difference between the 2 actions) on which animals rely their decisionmaking.

Regression analysis of the activity of individual neurons at different time bins within the ready period identified significant fractions of neurons that encode ΔQ in all 6 areas, with the highest fraction in the RSC (Figure 1C). ΔQ coding in these neurons was independent of upcoming choice directions (Figure S1), and reliably updated at single-trial resolution (Figure S2), indicating that these neurons faithfully encoded ΔQ on a trial-by-trial basis across all 6 areas. Despite the widespread ΔQ coding, the temporal stability of ΔQ coding within the ready period differed across areas. Only in the RSC, the ΔQ -coding neurons identified at different time bins reliably encoded ΔQ throughout the trial and across trials, while the encoding was temporally unstable in the other 5 areas (Figures 1D and S1H). This was because the way individual neurons encoded ΔQ across time differed across areas (Figures 1E and S1I). We quantified the temporal stability of ΔQ coding by defining the persistency index which reflects the coding persistency relative to the chance level (STAR Methods). The analysis revealed RSC as the area with the highest ΔQ coding persistency (Figure 1F).

We next examined whether the coding persistency is a fixed property of individual areas or changes with learning. We analyzed the population activity from RSC, PPC, pM2 and ALM during early stages of training (< 1 week from training start, Figure 1G). We compared their value coding persistency between early and expert sessions. We found that the ΔQ coding persistency significantly increases in RSC, PPC and pM2 during

training (Figures 1H and 1I), indicating that coding persistency can change during task learning.

The coding persistency may have increased during learning because the value-based decision task requires stable value maintenance for an extended period of time across trials. Therefore, we tested whether coding persistency differs in another task that does not require long maintenance of value. We trained 9 mice in the alternate choice task in which a reward was given when mice made a choice that was the opposite to the previous action (Figure 2A). Thus, the correct action depended only on the immediately preceding trial, in contrast to the value task in which history from multiple past trials was informative. All other task conditions were identical between the 2 tasks. camk2-tTA:: tetO-GCaMP6s transgenic mice were trained in the alternate choice task for at least 2 weeks to achieve a plateau-level performance (~80% correct) (Figure 2B). We then performed 2-photon calcium imaging of 8,524 RSC cells, 3,186 PPC cells, 7,915 pM2 cells and 4,911 ALM cells (RSC: 14 populations, 608.9 ± 18.1 cells, PPC: 7 populations, 455.1 ± 25.1 cells, pM2: 14 populations, 565.4 ± 34.6 cells, ALM: 10 populations, 491.1 ± 36.8 cells, mean \pm s.e.m per population). The coding persistency of action history in the alternate choice task was significantly weaker than in value-based task for the 4 imaged areas (Figures 2C and 2D). These results indicate that the coding persistency in the cortex is context dependent.

Persistent value coding in RSC forms cylindrical

In the value-based task, ΔQ coding in RSC is temporally stable within each trial. However, this does not necessarily mean that RSC population activity is static during these periods. In fact, individual neurons in RSC showed heterogeneous and rather dynamic activity patterns (Figure 1B). To investigate how the coding of different information temporally interacts, we sought to decompose population activity into the demixed neural subspaces where different task-related signals are separated into



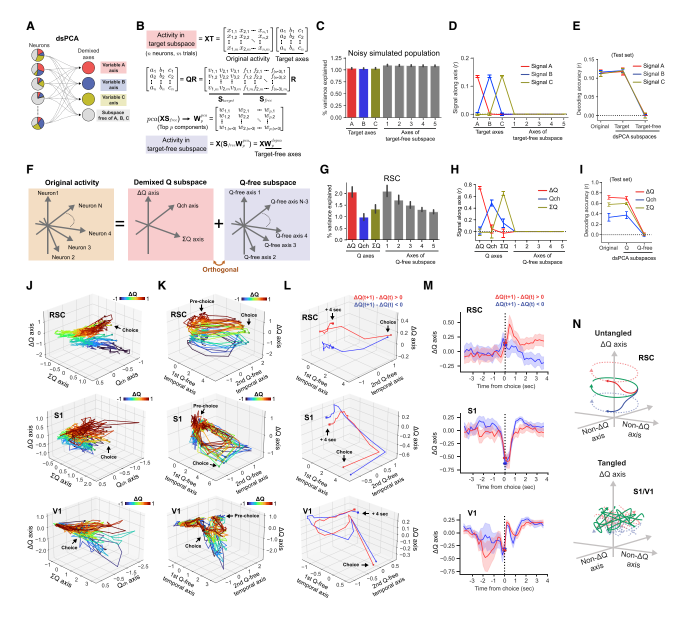


Figure 3. dsPCA reveals cylindrical dynamics with untangled value representation in RSC

(A) dsPCA decomposes the activity of a population of individual neurons that exhibit mixed selectivity for multiple variables into demixed dimensions and the remaining subspace that is free of the targeted signals.

- (B) Matrix operations to identify the target-free axes. Full QR decomposition of a matrix with target axes (T) identifies a set of basis vectors that spans the targetfree subspace (S_{free}). These target-free axes are realigned based on the principal component vectors (W_p^{pca}, matrix with top p PCA loadings) of the activity in the target-free subspace. The target-free axes in the original n-dimensional space are the columns of $\mathbf{W}_{n}^{\mathsf{dspca}} = \mathbf{S}_{\mathsf{free}} \mathbf{W}_{n}^{\mathsf{pca}}$.
- (C) Fraction of activity variance along each target axis and the top 5 PC axes from the target-free subspace. dsPCA was performed on noisy simulated data with target signals A, B, and C (10 repeated simulations). The amount of variance is similar between the 5 target-free axes because only Gaussian noise remained in the target-free subspace.
- (D) Signals A, B, and C along each dimension identified with dsPCA for the simulated data. Pearson correlations between the projected activity and each signal are shown.
- (E) Decoding accuracy of target signals from original population activity, activity in the target subspace (3 dimensions), and activity in the target-free subspace (n-3 dimensions). 50,000 and 10,000 trials for training and test sets.
- (F) We applied dsPCA to decompose the original population activity into the demixed Q subspace that consists of ΔQ , Qch, and ΣQ dimensions, and the Q-free subspace which is orthogonal to the Q subspace.
- (G) Fraction of activity variance along each Q-related axis and the top 5 PC axes from the Q-free subspace for RSC populations. Unlike simulated data (C), the amount of variance between axes of the Q-free subspace differ, indicating that non-targeted correlated signals exist in the Q-free subspace.
- (H) Q-related signals along each dsPCA dimension for RSC populations. Pearson correlations between the projected activity and each signal are shown.
- (I) Decoding accuracy of Q signals from the original RSC population activity, activity in the Q subspace (3 dimensions), and activity in the target-free subspace.

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distinct dimensions. Specifically, we sought to define 3 demixed axes each encoding $\Delta Q,\,Q_{ch}$ (value of selected action [e.g., Q_L on left choice trial]), or ΣQ (sum of 2 values), and the remaining Q-free subspace that retains all the activity variance that is not explained by the 3 Q-related axes. A previous study reported demixed principal component analysis (dPCA) (Kobak et al., 2016) as a method to decompose population activity into demixed target-dependent and independent dimensions. However, dPCA is only designed to identify dimensions for discrete variables and cannot be applied for continuous variables such as Q-related signals. In addition, dPCA splits each targeted signal into multiple linear axes, which makes the signal interpretation difficult. To overcome these limitations, we developed a novel dimensionality reduction method that is more generally applicable, which we term demixed subspace principal component analysis (dsPCA) (Figure 3A). dsPCA identifies demixed dimensions for targeted signals and dimensions for target-independent activity, similarly to dPCA. However, unlike dPCA, it groups each of the target signals along a single linear coding dimension and can identify such dimensions for both discrete and continuous target variables. The first step of dsPCA identifies the best demixed linear axes for the target variables using a regression-based approach, similarly to (Mante et al., 2013). This step involves fitting a multiple linear regression model of the form $x(trial) = \beta_A A(trial) + \beta_B B(trial) + \beta_C C(trial) + \beta_0$ to the activity of individual neurons for the targeted variables, A, B, and C. The regression coefficients, β_A , β_B , and β_C are the partial derivatives of the neural activity by each target variable, and the vectors that consist of the coefficients from all neurons are the linearly demixed coding directions of the neural population for the 3 targeted variables. We defined the targeted coding axes as the unit vectors of these coding directions. By definition, these demixed coding vectors capture all linear information of targeted variables in a population. Next, dsPCA identifies the remaining target-free subspace that is orthogonal to these targeted axes and captures all the remaining activity variance. The target-free orthogonal subspace is identified by performing full QR decomposition of the matrix with the coding axis vectors. Then the axes of the target-free subspace are further realigned based on the principal components of the activity within the target-free subspace to define axes that contain large fractions of remaining variance. (Figure 3B). Therefore, dsPCA can be viewed as a general extension of PCA by combining the regression-based supervised target axis identifications and the PCA-based unsupervised dimensionality reduction of the target-free population dynamics.

We evaluated the demixing performance of dsPCA using noisy simulated neural populations (200 neurons/population with Gaussian noise) where graded signals A, B, and C are linearly encoded in 20% of the neurons. Each target signal was uniquely encoded only along the single target axis (Figures 3C and 3D), and linear decoders failed to decode any A, B, or C signals in the remaining target-free subspace (Figure 3E). We next applied dsPCA on the cortical population activity time-averaged over the ready period to identify demixed coding axes for ΔQ , Q_{ch} , and ΣQ , and the remaining, Q-free subspace (Figure 3F). For all 6 areas, most of the targeted information was confined to each of the coding axes, and the remaining subspace completely lacked any of the targeted information even though this subspace contained the highest activity variance (Figures 3G, 3H, 3I, and S3). Although we detected some Qch signal along the ΣQ axis (Figures 3H and S3B), this is expected because Q_{ch} is a component of ΣQ ($\Sigma Q = Q_{ch}$ + unchosen Q). However, note that ΣQ signal is not detectable along the Q_{ch} axis, indicating that the demixing of activity variance worked correctly. Thus, dsPCA successfully identified demixed coding axes for Q-related variables and the remaining Q-free subspace.

With dsPCA, we examined how ΔQ coding temporally interacts with other dynamics. The activity dynamics around the choices (between ± 4 s from the choice) was visualized in the neuronal manifold consisting of the ΔQ coding axis and the other value-related axes (Figure 3J), or the manifold consisting of the ΔQ coding axis and the 2 largest temporal activity variance axes within the Q-free subspace (Figures 3K). We found in both manifolds that activity trajectories in RSC from trials with different ΔQ values do not cross with each other across time. In the manifold with the largest temporal dynamics (Figures 3K, S4 and S5), the RSC population remained in the initial positions linearly segregated along ΔQ axis according to ΔQ of the trial ('Pre-choice' in the figures). Around the go cue time, the RSC population diverged from these initial positions and drew rotational dynamics. After a choice, the population returned toward the initial positions following a circular geometry. The return geometry was warped along the ΔQ axis, reflecting the reward prediction error (RPE) on each trial depending on the choice and its outcome, which updates the ΔQ representation in the population (Figures 3L and 3M). The RPE-dependent, bidirectional transition of the activity state ensures that the neural population closely represents and updates the ΔQ coding online in each trial. In contrast, the dynamics in S1 and V1 were highly tangled over time, and similar ΔQ values could accompany different activity states at different times. Therefore, although ΔQ coding is widely distributed across the cortex, the different levels of persistency confer different levels of tangling in ΔQ coding (Figure 3N). The exceptionally high ΔQ coding persistency in RSC allows a temporally untangled value representation with the within-trial cyclic dynamics that transitions along the value axis

All error bars are 95% Cl.

⁽J–K) Example RSC, S1, and V1 population activity dynamics in neuronal manifolds where ΔQ axis is paired with Qch and ΣQ axes (J), or axes that reflect major within-trial temporal activity variance of Q-free subspace (K). dsPCA was applied on the activity between -2 and -1 s from choice, and the activity between ± 4 s from choice was projected onto the identified axes. Circles indicate the choice time. Projected activity was temporally downsampled to non-overlapping

⁽L) Activity state transitions along ΔQ axis according to the updated action values in RSC, whereas S1 and V1 activity draw complex trajectories that lead to tangling in the geometry. Post-action selection trajectory was separately averaged according to the sign of ΔQ update.

⁽M) Activity state transitions in (L) shown along the ΔQ axis.

⁽N) Population activity in RSC forms cylindrical dynamics where within-trial cyclic dynamics can transition along ΔQ axis across trials according to the RPE, while in the other areas ΔQ representation is tangled.



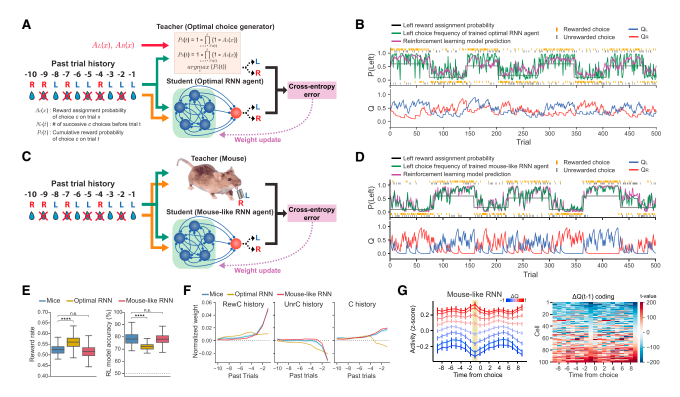


Figure 4. Untangled persistency emerges in the artificial RNNs trained to perform "mouse-like" RL

(A) Optimal RNN agent was trained by updating its synaptic weights to minimize the discrepancy in decisions (cross-entropy error) between the teacher (optimal choice generator) and the student (RNN).

- (B) Behaviors of the trained optimal RNN agent in an example session. The agent ran the task by itself using its recurrent activity dynamics to implement RL. The left choice probability of the RNN agent was taken from its output neuron activity. Left (QL) and right (QB) action values were estimated by fitting a RL model to the behaviors.
- (C) Mouse-like RNN agent was trained by updating its synaptic weights to minimize the discrepancy in decisions (cross-entropy error) between the teacher (expert mice) and the student (RNN).
- (D) Behaviors of the trained mouse-like RNN agent in an example session.
- (E) Frequency of rewarded trials (left) and choice predictability by a RL model optimized to describe expert mouse behaviors (right, 5-fold cross-validation). n = 82 sessions for mice, 500 sessions (5 trained networks, each ran 100 sessions of 500 trials/session) each for the optimal and mouse-like RNN agents.
- (F) Decision dependence on history from past 10 trials, quantified by a regression model (STAR Methods). RewC: rewarded choice, UnrC: unrewarded choice, C: outcome-independent choice history. n = 82 sessions for mice, 5 sessions (5 trained networks, each ran 10,000 trials) each for the optimal and mouse-like RNN agents. The regression weights were normalized by the model accuracy. Error bars are 95% CI.
- (G) Activity of ΔQ coding neurons that were identified using the activity at the highlighted time bin (yellow shading, -1 time step before choice) in the recurrent layer of a trained mouse-like RNN agent (left), and the t-values of ΔQ after choice for the activity in each time bin (right). Each trial had 10 time steps, and 0 corresponds to the choice time. t-values were sorted based on the last time step (+9). The t-values in RNNs are higher than in mice due to smaller amount of activity noise. Error bars are SEM.

to reflect value updates. These dynamics across trials collectively form cylindrical dynamics during task performance.

Untangled, persistent value coding emerges in the RNN trained with the mouse RL strategy

The persistent and untangled ΔQ coding in the RSC, together with our previous observation that RSC inactivation impairs value-based decision (Hattori et al., 2019), raises the possibility that persistent value coding is advantageous in the task. We investigated this possibility by training artificial RNN agents to perform RL in the same task and subsequently examining the ΔQ coding scheme in the trained network. The training of RNNs was done without constraining the activity dynamics. We reasoned that, if persistent coding is advantageous, trained RNN agents may use persistent coding to perform the task.

First, we trained RNNs to perform RL optimally by teaching them the ideal choices of each trial based on the reward assignment rule. In this task, once a reward is assigned to a choice, the reward remains assigned until the choice is selected. As a result, the actual reward probability of a choice cumulatively increases if the choice is not selected in the recent trials. Therefore, an optimal choice would depend on the current reward assignment probabilities, which are unknown to mice and RNN agents, and past choice history. By using the optimal choices as the teacher, we trained synaptic weights of RNNs such that the RNNs use only history of choice and reward to make near-optimal decisions (Figures 4A and 4B). The durations between decisions were made variable, similarly to the task structure in mice. The RNNs receive action outcome information only at the time step after choice and need to maintain the information through

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recurrent connectivity across time steps and trials. These optimally trained networks ("optimal RNN agents") achieved higher reward rate than expert mice (Figure 4E). Furthermore, the choice patterns of optimal RNN agents diverged from the RL model that has been optimized to describe the behavior of expert mice (Figure 4E), indicating that the optimal RNN acquired a RL strategy that is distinct from mice. Accordingly, a regression analysis showed that the dependence of optimal RNN agents on choice and reward history differed from that of expert mice

To obtain a network model that better mimics the mouse strategy, we trained RNNs to imitate expert mouse behaviors using behavioral cloning, a form of imitation learning (Osa et al., 2018). We used 50,472 decision making trials of expert mice as the teaching labels to train the synaptic weights of the RNN. The goal of this training was for the RNN to make the same decisions as expert mice with its recurrent activity dynamics based on the same history of choice and outcome in the past trials (Figure 4C). The trained RNNs ("mouse-like RNN agents") performed RL using their recurrent activity (Figure 4D), and the reward rate and the RL model accuracy were equivalent to those of expert mice (Figure 4E). Furthermore, the mouse-like RNN agents used history from previous trials for its decisions in a similar way as expert mice (Figure 4F). Therefore, the RL strategy of expert mice was successfully transferred to the synaptic weights of the trained RNN agents, and the trained RNNs could implement mouse-like RL using its recurrent activity dynamics without updating synaptic weights from trial to trial.

We then examined how the mouse-like RNN agents encoded ΔQ . We found that RSC-like persistent ΔQ coding emerged in their recurrent activity (Figures 4G). This observation is significant as the training procedure did not impose a priori constraints on the coding scheme of the RNN. We also examined how the population activity dynamics evolved during training. We had RNN agents at 3 stages of training run the task (before training, intermediate (after 1 epoch of training), and fully trained) and analyzed their recurrent activity during the task performance. dsPCA revealed that untrained networks with random connectivity exhibit highly tangled ΔQ coding, while training gradually shaped the networks to form stacked circular dynamics (Figure 5A). Unlike RSC that formed cylindrical dynamics (Figure 3K), the diameter of rotational trajectory varied across different ΔQ states in the trained networks, suggesting that additional biological constraints that were not considered for RNN training may have imposed a constant diameter in the mouse brain. In addition to the analysis of ΔQ estimates from an RL model fit to behaviors, we examined the coding persistency of the ground truth ΔQ which is available as the activity of the action output neuron in each RNN agent. We confirmed that the ground truth ΔQ was also persistently encoded in both optimal and mouse-like RNN agents (Figure S6).

Persistency facilitates reliable and robust value retrieval by downstream neural networks

The emergence of ΔQ coding persistency in RNN agents suggests that persistent coding is a preferred solution in the task. What would be the advantage of persistent coding? One possibility is that untangled persistency may allow a more reliable signal retrieval by the downstream network to guide the action selection. We tested this possibility by training artificial RNNs to retrieve the ΔQ signal from different temporal patterns of simulated population activity (Figure 6A). For this purpose, RNNs are biologically relevant as they receive time-varying inputs sequentially, as opposed to other decoder models (e.g., regression models).

We created artificial population activity encoding ΔQ in 4 different patterns: persistent, and 3 types of non-persistent coding (Figure 6B). In persistent coding, 20% of cells encode ΔQ as rate coding persistently. The slope of the ΔQ tuning curve for each neuron was taken from its distribution among RSC neurons (Figure S7). For the first 2 types of nonpersistent coding, the cellular identity of the persistent coding pattern was shuffled independently at each time bin to alter the ΔQ persistency of each neuron without altering the population-level ΔQ signal in each time bin. Nonpersistent 1 allowed each neuron to encode ΔQ in multiple time points, while nonpersistent 2 was constrained that each neuron encodes ΔQ in only one of the 5 time points. In the third nonpersistent coding scheme, binary signals (active or inactive) at each time bin were used to encode ΔQ by activating distinct sequences of neurons across time for different values of ΔQ . We prepared 10 different sequences for 10 bins of ΔQ values.

Using these activity patterns as inputs, we trained RNNs to retrieve ΔQ . Various levels of noise were added to the input activity to test a range of signal-to-noise ratio (SNR). The RNN trained with the persistent ΔQ codes was able to retrieve ΔQ better than those trained with nonpersistent codes, especially when the input activity noise was high (Figure 6C-D). This indicates that persistent coding facilitates reliable information retrieval by downstream circuits. Furthermore, the RNNs that were trained to retrieve ΔQ from persistent coding were more robust to changes in the synaptic weights, loss of synapses, and cells (Figure 6E).

To investigate the impact of persistency in the brain activity, we next examined how ΔQ could be retrieved from the neural activity with different levels of persistency recorded from the 6 cortical areas (Figure 6F). In addition to the original recorded activity ('Raw'), we artificially increased or decreased ΔQ coding persistency by temporally sorting ('Sorted') or shuffling ('Shuffled') the cell identity in each area. These persistency manipulations simply changed the neuron ID of activity and thus did not alter the total amount of ΔQ signal in each time bin. Using these sets of neural activity as inputs, we trained RNNs to retrieve ΔQ . There was a general trend that an increase in persistency (sorted activity) improved retrieval accuracy, while a decrease in persistency (shuffled activity) impaired retrieval accuracy (Figure 6G). However, the effect size differed across different cortical areas. We found that the increase in retrieval accuracy by sorting was larger when the original persistency in the population was lower, and the decrease in retrieval accuracy by shuffling was larger when the original persistency was higher (Figures 6H and 6I). These results further support the notion that coding persistency is a critical determinant that enhances the accuracy of information retrieval by the downstream network.

The results above indicate that persistent codes can be read out by the downstream more effectively than nonpersistent codes when the artificial neural network is allowed to train its



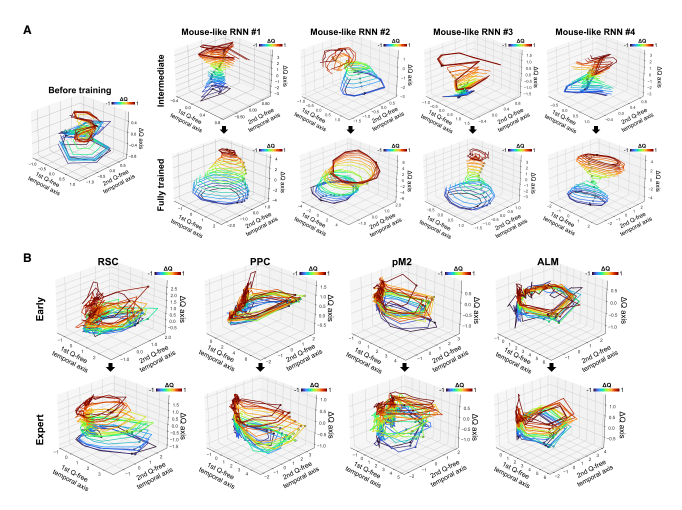


Figure 5. Cylindrical dynamics emerges in mouse-like RNN agents and mice during training

(A) Population activity dynamics of the recurrent layer of mouse-like RNN agents in neuronal manifolds where ΔQ axis is paired with axes that reflect major withintrial temporal activity variance in Q-free subspace. Agents at each training stage ran the task for 10,000 trials. dsPCA was applied on the activity averaged between -5 and -1 time steps from choice, and the population activity between ± 5 time steps from choice was projected onto the identified axes. 4 independently trained mouse-like RNN agents are shown. Circles indicate the choice time.

(B) Population activity dynamics of example RSC, PPC, pM2, and ALM populations in early and expert sessions. The same population of neurons was longitudinally compared for each area. dsPCA was applied on the activity averaged between -2 and -1 s from choice, and the population activity between ± 4 s is visualized. Circles indicate the choice time.

synaptic weights by minimizing the difference between its output and the target (ΔQ) as supervised learning. However, in the real brain, such an explicit supervised target label to guide the shaping of network connectivity is rarely available. Another approach to shape the connectivity to retrieve particular information is unsupervised learning where errors are computed using information readily available to the local network such as the input itself (Lillicrap et al., 2020). Therefore, we next considered the possibility that coding persistency may also affect signal retrieval processes that do not necessitate a supervised target label for each information. It has been suggested that the brain may implement unsupervised learning in a similar way to autoencoder networks in which the target is the input itself (Lillicrap et al., 2020). Autoencoders extract the most dominant signals from the input activity and represent them in the activity of a small number of neurons in the coding layer. The networks shape their connectivity by reconstructing the input activity from the coding layer and minimizing the reconstruction error between the input and the reconstructed activity. To examine what information in the input population activity can be extracted in an unsupervised manner by downstream recurrent networks, we used a recurrent denoising autoencoder (RDAE) (Maas et al., 2012; Vincent et al., 2010) that sequentially processes input activity and extracts the latent representations embedded in the input activity sequence, which are sufficient to reconstruct the original population activity sequence with noise robustness (Figure 7B; STAR Methods). When the RDAE was trained on RSC population activity, ΔQ was extracted in the most dominant dimensions of neural activity in the coding layer (Figure 7A). The ΔQ representation in the coding layer was independent of upcoming choice directions, indicating that the dimensions reflect value and not motor plans. Other task-related signals were not represented





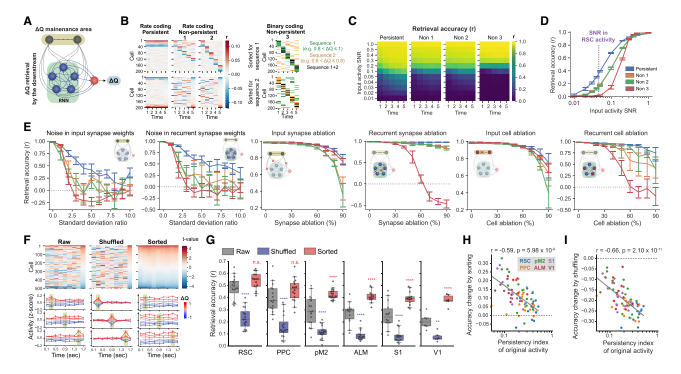


Figure 6. Persistency in value coding facilitates reliable and robust value retrieval by downstream neural networks

(A) RNN (40 recurrent units) was trained to retrieve ΔQ from the input population activity sequence with either persistent or non-persistent ΔQ coding. (B) Artificial population activity with either persistent or non-persistent AQ coding in the 200-cell sequence. 3 types of non-persistent mode were considered (2 rate coding, 1 binary coding; STAR Methods). In the rate coding populations, the color indicates the Pearson correlation between the activity and ΔQ (20% of neurons at each bin encode ΔQ). Example populations were visualized by either clustering ΔQ-coding neurons at each time bin (top) or sorting neurons based on the correlation at the last time bin (bottom). In the binary coding population, ΔQ is encoded by a unique activity sequence across time for each bin of ΔQ values (ten evenly spaced bins between ± 1). 20% of neurons at each time bin participate in each sequence. In the example, cells are sorted for either sequence 1 or 2. Time bins that are active in both sequences are colored black.

- (C) Mean ΔQ retrieval accuracy by the downstream RNNs from populations with different coding modes and varying SNR (10 simulations for each).
- (D) The ΔQ retrieval accuracy at the 5th time step with different SNR in the input activity. The purple dashed line indicates the median SNR of ΔQ coding in imaged
- (E) Robustness of trained RNNs. Simulations were performed using artificial population activity with SNR of 1. Noise to synaptic weights was given by Gaussian noise with the standard deviation relative to the standard deviation of the weight distribution of each connection type. Error bars in (D) and (E) are 95% CI.
- (F) Artificial manipulations of ΔQ coding persistency illustrated in an example PPC population during ready period. Error bars are SEM.
- (G) ΔQ retrieval accuracy before and after the persistency manipulations (subsampled 240 cells were used, **p < 0.01, ****p < 0.0001, one-way ANOVA with Tukey's HSD).
- (H) Gain in retrieval accuracy by sorting correlates with the original ΔQ coding persistency.
- (I) Loss in retrieval accuracy by shuffling correlates with the original ΔQ coding persistency.

as the dominant signals in the coding layer (Figure S8). Similar results were observed in the activity dynamics of the mouse-like RNN agent but not in S1. Systematic comparisons among 6 cortical areas revealed that extracted ΔQ in the coding layer was especially high from RSC, and the amount of extracted ΔQ showed a high correlation with the ΔQ coding persistency in the input population activity (Figures 7B-7D). To directly test the effect of persistency, we artificially manipulated the persistency of ΔQ coding in RSC without changing the total amount of ΔQ signals in the population. We found that artificial increases in the persistency by sorting the cell identity improved the ΔQ extraction, while artificial decreases in the persistency by shuffling the cell identity worsened the ΔQ extraction (Figure 7E). These results indicate that high persistency in the input activity can allow ΔQ retrieval by the downstream network even without supervised learning.

Taken together, these analyses indicate that the persistency of value coding facilitates a robust and accurate readout of value by downstream networks.

Signal leakage can contribute to distributed value coding with varying levels of persistency

The results so far indicate computational advantages of persistent coding. However, in the mouse brain, ΔQ coding was widely distributed across the 6 cortical areas with different levels of persistency (Figures 1C-1F). We asked whether anatomical connectivity among cortical areas relates to the persistency levels of value coding. We analyzed the connectivity among imaged areas using the dataset from the Allen Mouse Brain Connectivity Atlas (Oh et al., 2014). Focusing on the projections from each of the 3 areas with high ΔQ persistency (RSC, PPC, pM2), we quantified their axon projection density in each of the other 5 imaged areas



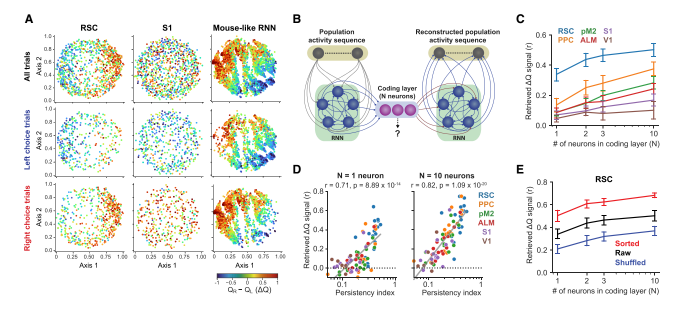


Figure 7. Persistency in value coding also facilitates unsupervised value retrieval by downstream neural networks

(A) Representation of input population activity in the coding layer of denoising recurrent autoencoder networks (RDAE). Each network was trained to extract major signals from example populations of RSC, S1, or a trained mouse-like RNN agent (5,000 trials). Population activity sequence during ready period was used as the input. Each data point corresponds to a trial, with the colors indicating the ΔQ of the trial. Trials were separated according to the choice directions in the upcoming answer period in the bottom 2 rows. The dominant signals extracted in the activity of coding neurons (10 neurons) were visualized in 2 dimensions by multidimensional scaling

- (B) RDAEs extract major signals of the input population activity into the activity of N neurons in the coding layer by unsupervised learning.
- (C) Decoding accuracy of ΔQ from the activity of N neurons in the coding layer. A simple feedforward neural network (N neurons in the coding layer are connected to a single output neuron with tanh activation function) was used to decode from the coding layer. Input populations were subsampled 240 cells.
- (D) Decoding accuracy of ΔQ from the activity of neurons in the coding layer (N = 1 and 10) positively correlates with the ΔQ coding persistency of the input population activity.
- (E) Artificial manipulations of ΔQ coding persistency in the input RSC population bi-directionally alter the amount of extracted ΔQ signal in the coding layer. All error bars are SEM.

(Figure 8A). We found that RSC, PPC, and pM2 predominantly project to each other, with smaller amounts of direct projections to ALM, S1, and V1 (Figures 8B, 8C and S9). Thus, 3 areas with persistent and strong ΔQ coding densely connect with each other, while they send less direct projections to the other 3 areas with weaker and less persistent ΔQ coding. Based on this observation, we hypothesized that the weak ΔQ persistency in ALM, S1 and V1 could result from a signal leakage from the areas that maintain ΔQ as persistent activity. To test this hypothesis, we built RNNs with multiple recurrent layers that receive RSC activity through nonspecific synaptic connectivity and examined how ΔQ coding changes along the downstream hierarchy of layers (Figure 8D). We found that the fractions of neurons with ΔQ coding gradually decreased as the signal leaked through layers of recurrent connectivity (Figures 8E and 8F). Concurrently, ΔQ coding became increasingly less persistent (Figure 8G), and the temporal tangling of ΔQ coding in neuronal manifolds gradually increased in the downstream (Figure 8H). Furthermore, artificial manipulations of ΔQ coding persistency in the input RSC activity revealed that persistency in ΔQ coding can affect the robust distribution of ΔQ coding with graded levels of persistency across the downstream layers (Figures 8F and 8G). We obtained similar results using PPC and pM2 as the input activity (Figure S10A–S10H), and the decreases in the ΔQ coding

neurons and the ΔQ coding persistency in the downstream layers were more dramatic when the direct neural projections from layer to layer were sparse (Figures S10I-S10K). These results indicate that, even without specific connectivity to selectively route particular information, persistently encoded information can propagate thorough layers of nonspecific connectivity to lead to a wide distribution of the information encoded with lower levels of persistency in downstream areas.

DISCUSSION

Brain-wide distribution of task-related information has emerged as a common principle in recent years. In many cases, such as what we observed for ΔQ coding (Figure S1), task-related signals are encoded by a heterogeneous population with some cells increasing but others decreasing their activity. Such information coding may not be identified with classical large-scale recording techniques such as fMRI, EEG and ECoG that quantify population average responses. Even though information coding is widespread, the way by which information is encoded differs across areas (Hattori et al., 2019). In the present study, the big data of > 100k mouse decisions and the activity from > 100k neurons in 2 behavioral tasks allowed us to investigate the potential origin of the distributed information coding and the computational

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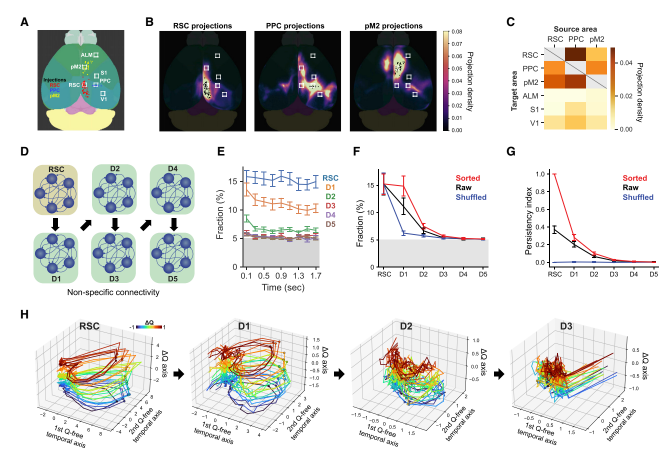


Figure 8. Non-specific signal leakage can contribute to widely distributed value coding with graded persistency

(A) Injection coordinates for anterograde tracing virus. RSC (red, n = 60 experiments), PPC (blue, n = 9), and pM2 (yellow, n = 33). Experiments with left hemisphere injections were mirrored horizontally. Experiments with both WT mice and Cre-transgenic mice were included (See Figure S9 for WT only). White squares indicate the imaging FOVs used for our neural activity analyses.

- (B) Mean projection density of axons from each source area. Black dots indicate the injection coordinates.
- (C) Connectivity matrix with the mean projection density from each source area to the 6 target areas that we used for our neural activity analyses (500 µm × 500 μm white squares).
- (D) RSC population activity sequences were processed through 5 recurrent layers with non-specific connectivity. Connection probability from layer to layer was set to 20% (Other probabilities in Figure S10).
- (E) Fractions of ΔQ coding neurons at each of the 200 ms time bins during ready period (Regression, p < 0.05, 2-sided t test). Error bars are SEM.
- (F) Mean fractions of ΔQ coding neurons at each layer during ready period. Fractions of time bins within the ready period were averaged for each population. Artificial manipulations of ΔQ coding persistency in RSC does not affect the fractions of ΔQ coding neurons in RSC, but affect the fractions in the downstream. (G) ΔQ coding persistency at each layer. Persistency progressively decreases in the downstream. Artificial manipulations of ΔQ coding persistency affect the persistency in the downstream. Error bars in (F) and (G) are 95% CI.
- (H) Temporal dynamics of population activity states visualized with dsPCA applied at each layer. Cylindrical dynamics gradually collapses into highly tangled dynamics in downstream layers.

advantages of persistent coding using data-driven machine learning approaches. Coding persistency was both learning and context dependent, and the persistent coding emerged during task learning in both mouse brain and artificial network agents performing the same task. Persistency facilitates the untangled maintenance of information as well as its reliable retrieval by downstream circuits. The observation that persistency is context dependent suggests that certain cortical areas such as the RSC can adjust coding persistency depending on behavioral demands. For example, persistency may be especially preferred when the task context requires extended maintenance of the information, or the maintained information is graded as in the case of value, so that information can be stably maintained and robustly retrieved by downstream areas. Furthermore, we showed that persistent coding in key areas such as the RSC could also contribute to the wide distribution of ΔQ coding across the mouse brain even through nonspecific signal leakage. The same principle may also apply to other task-related signals in various task conditions, providing a possible explanation for the widespread phenomenon of distributed coding across the brain. In other words, a wide distribution of information is expected across the interconnected network of the brain, unless specific connectivity restricts the propagation of particular information. We note that nonspecific leakage is one of potential





mechanisms for signal distribution and it remains to be shown how much such a mechanism contributes to the phenomenon. Furthermore, this mechanism is agnostic as to whether the propagated information has a function in the downstream areas leaked information could contribute to various computations performed in downstream areas.

We trained artificial RNNs to imitate the mouse behavioral strategy using behavioral cloning and investigated the activity dynamics that emerged in the RNNs that were trained without activity constraints. Previous studies trained task-performing artificial neural networks either by using the correct action labels which are defined in each task structure (e.g., action A must be taken after stimulus A) (Masse et al., 2019; Orhan and Ma, 2019) or by RL (Banino et al., 2018; Song et al., 2017; Tsuda et al., 2020; Wang et al., 2018). Both approaches train the networks to learn the optimal strategy in the respective task, independent of the actual behavioral strategy that animals learn in the environment. In our value-based decision task, animals learn to use behavioral history for decisions during training, but the RL strategy that animals develop was suboptimal (Figures 4E and 4F). The origins of the sub-optimality likely include (1) limited memory capacity, (2) low sample efficiency, (3) limited amount of training trials, and (4) inductive bias inherent to each species. Deep RL, an artificial network that learns to solve a task with RL, does not always have these constraints, and thus it learns a nearoptimal strategy unlike animals. These artificial networks may not reflect the mechanisms used by the brain. In another common approach, simpler mathematical models (e.g., regression, classical RL models) directly fit to animal behaviors are useful to understand the behavioral strategies. However, they do not provide insights into potential neural activity dynamics that may mediate the behaviors. To overcome these issues, we trained artificial RNNs, using mice as the teachers, to acquire the sub-optimal RL strategy that mice develop during training. The big data of ~50k decisions collected from expert mice allowed us to successfully train RNNs to imitate mouse behavioral strategy. This data-driven approach to train RNNs to implement animal/human-like behaviors would be a useful approach to obtain the neural network models and analyze what kind of activity dynamics allows the animal strategy in a particular task. Similarly to our approach, convolutional neural networks have been trained in visual object recognition tasks. The training was done to perform the task optimally, as opposed to our approach using behavioral cloning. Nevertheless these networks have been shown to develop some neural activity characteristics that resemble the neural activity in the visual system of animals (Kriegeskorte, 2015; Yamins and DiCarlo, 2016). These deep learning approaches will be a powerful approach to understand what kind of neural activity may mediate given behaviors.

In this study, we developed dsPCA, a novel dimensionality reduction method which combines the strengths of supervised and unsupervised algorithms. The supervised aspect allows us to identify the best demixed linear coding dimensions for targeted task-related variables, and the unsupervised aspect allows us to identify nontargeted correlated signals in the remaining population activity. Therefore, dsPCA is a generally applicable method to understand both the signals of interest and other nontargeted correlational structures in high-dimensional data. Using dsPCA, we found that both mouse brain and artificial RNN agents develop cylindrical dynamics, which consists of within-trial cyclic dynamics and its across-trial transition along ΔQ axis. Similar within-trial dynamics have been well studied in monkey motor cortex during arm movement (Churchland et al., 2012; Russo et al., 2018, 2020). The studies showed that the population activity state draws untangled rotational dynamics during movements. They also showed that the activity state draws a simple cyclic trajectory in the primary motor cortex, while the supplementary motor area draws a helical trajectory that unfolds along a single direction by reflecting the 'context' of the movement (Russo et al., 2020). The activity trajectory that we observed had cylindrical geometry, and the activity state repeatedly transitioned along the ΔQ axis based on the RPE. These spatially confined geometries ensure the untangled representation of ΔQ , which contributes to a robust ΔQ representation in the brain. dsPCA and other RNN-based approaches in this study would facilitate the geometric understanding of population dynamics in both biological and artificial networks.

STAR*METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j. neuron.2021.11.001.

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AUTHOR CONTRIBUTIONS

R.H. and T.K. conceived the project. R.H. performed the experiments, analyses and simulations with suggestions from T.K. R.H. and T.K. wrote the paper.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR*METHODS

KEY RESOURCES TABLE

Reagent or resource	Source	Identifier
Deposited Data		
Behaviors and imaging data in value-based decision task	Hattori et al., 2019,	https://doi.org/10.1016/j.cell.2019.04.027
Experimental Models: Organisms/Strains		
Mouse: CaMKIIa-tTA: B6; CBA-Tg(Camk2a-tTA)1Mmay/J	The Jackson Laboratory	RRID:IMSR_JAX:003010
Mouse: tetO-GCaMP6s: B6;DBA-Tg (tetO-GCaMP6s)2Niell/J	The Jackson Laboratory	RRID:IMSR_JAX:024742
Software and Algorithms		
Python3	Python Software Foundation	RRID:SCR_008394
MATLAB	MathWorks	RRID:SCR_001622
R	R Foundation for Statistical Computing	RRID:SCR_001905
Scanlmage	Vidrio Technologies, LLC	RRID:SCR_014307
Suite2P	Pachitariu et al., 2016	RRID:SCR_016434
BControl	The Brody lab	https://brodylabwiki.princeton.edu/bcontrol/index.php
TensorFlow2	Google	https://github.com/tensorflow/tensorflow
scikit-learn	Pedregosa et al., 2011,	https://github.com/scikit-learn/scikit-learn
SciPy	Virtanen et al., 2020,	https://github.com/scipy/scipy
NumPy	Harris et al., 2020,	https://github.com/numpy/numpy
Statsmodels	Seabold and Perktold, , 2010	https://github.com/statsmodels/ statsmodels
Seaborn	Waskom, 2021	https://github.com/mwaskom/seaborn
Matplotlib	Hunter, 2007	https://github.com/matplotlib/matplotlib
lme4	Bates et al., 2015	https://cran.r-project.org/web/ packages/lme4/
PatchWarp	Hattori and Komiyama, 2021	DOI: 10.5281/zenodo.5590965, https://github.com/ryhattori/PatchWarp
dsPCA	This paper	DOI: 10.5281/zenodo.5620834, https://github.com/ryhattori/dspca

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources should be directed to and will be fulfilled by the lead contact, Takaki Komiyama (tkomiyama@ucsd.edu).

Materials availability

This study did not generate new unique reagents.

Data and code availability

- Data reported in this paper are available from the lead contact upon reasonable request.
- dsPCA code has been deposited at Zenodo and is publicly available. The DOI and the link to the latest code in the GitHub repository are listed in the key resource table.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.





EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animals

The experimental data in the value-based decision task were first reported in ref. (Hattori et al., 2019). The data in the alternate choice task were newly collected for the current study. Both male and female mice were included in both datasets because we did not observe obvious sex-dependent differences in their neural activity patterns. Mice were originally obtained from the Jackson Laboratory (CaMKIIa-tTA: B6;CBA-Tg(Camk2a-tTA)1Mmay/J [JAX 003010]; tetO-GCaMP6s: B6;DBA-Tg(tetO-GCaMP6s)2Niell/J [JAX 003010]; tetO-GCAMP6s] 024742]). All mice (6 weeks or older) were implanted with glass windows above their dorsal cortex for in vivo two-photon calcium imaging. All mice were water-restricted at ~1ml/day during training.

METHOD DETAILS

Surgery

Mice were continuously anesthetized with 1%-2% isoflurane during surgery after subcutaneous injection of dexamethasone (2mg/ kg). After exposing the dorsal skull and removing the connective tissue on the skull surface using a razor blade, we marked on the skull with black ink at the coordinates of [AP from bregma, ML from bregma] = [+3.0 mm, 0 mm], [+2.0 mm, 0 mm], [+1.0 mm, 0 mm], $[0\,\text{mm},0\,\text{mm}],[-1.0\,\text{mm},0\,\text{mm}],[-2.0\,\text{mm},0\,\text{mm}],[-3.0\,\text{mm},0\,\text{mm}],[0\,\text{mm},\pm\,1.0\,\text{mm}],[0\,\text{mm},\pm\,2.0\,\text{mm}],[0\,\text{mm},\pm\,3.0\,\text{mm}],[-2.0\,\text{mm},\pm\,3.0\,\text{mm$ 1.0 mm], $[-2.0 \text{ mm}, \pm 2.0 \text{ mm}]$, $[-2.0 \text{ mm}, \pm 3.0 \text{ mm}]$. We then applied saline on the skull and waited for a few minutes until the skull became transparent enough to visualize vasculature patterns on the brain surface. We took a photo of the vasculature patterns along with marked coordinates and used it to find target cortical areas for two-photon microscopy. A large craniotomy was performed to expose 6 cortical areas, and a hexagonal glass window was implanted on the brain. The glass window was secured on the edges of the remaining skull using 3M Vetbond (WPI), followed by cyanoacrylate glue and dental acrylic cement (Lang Dental). After implanting the glass window, a custom-built metal head-bar was secured on the skull above the cerebellum using cyanoacrylate glue and dental cement. Mice were subcutaneously injected with Buprenorphine (0.1 mg/kg) and Baytril (10 mg/kg) after surgery.

Behavior task and training

Mice were water-restricted at 1-2 ml/day after a minimum of 5 days of recovery after surgery. We began animal training in pre-training tasks after at least a week of water restriction. We used BControl (C Brody), a real-time system running on Linux communicating with MATLAB, to control behavioral apparatus. We placed 2 lickports in front of head-fixed mice to monitor their licking behaviors and give water rewards. Licking behaviors were monitored by IR beams running in front of each water tube. We used an amber LED (5mm diameter) as the ready cue and a speaker for auditory cues. Each trial begins with a ready period (2 or 2.5 s with the amber LED light), followed by an answer period with an auditory go cue (10 kH tone). The 10 kHz tone was terminated when animals made a choice (the first lick to a lickport) or when the answer period reached the maximum duration of 2 s. Mice received a 50 ms feedback tone (left: 5 kHz, right: 15 kHz) after a choice. ~2.5 μl water was provided to mice on each rewarded trial from a lickport.

Before running in the alternate choice task or value-based decision task, mice were trained in 2 pre-training tasks. In the 1st pretraining task, mice were rewarded for either choice during the answer period. We gradually increased the mean ITI from 1 s to 6 s with ± 1 s jitter. Through training in this task (2-3 days), mice learn that they can obtain water rewards from the 2 lickports if they lick during the answer period. In the 2nd pre-training task, reward location alternated every trial irrespective of their choice directions. Furthermore, licking during ready period was punished by 500 ms white noise alarm sound and trial abort with an extra 2 s ITI in addition to the regular 5-7 s ITI. Through training in this 2nd pre-training task (2-3 days), mice learned to lick from both lickports and withhold licking during the ready period.

Alternate choice task

In the alternate choice task, mice need to change their choice from a previous trial to get a water reward. For example, if a mouse chose left on one trial, regardless of whether the mouse received a reward or not, a water reward is available only from the right choice on the next trial. The mouse will not get any rewards by repeating left choices for many trials because a reward will not be assigned to the left until the mouse collects the assigned reward on the right side. Mice need to rely on which side they chose in the previous trial to make the correct choice. ITI was 5-7 s, and the trials with licking during ready period were classified as alarm trials (500 ms white noise alarm sound and extra 2 s ITI). Mice were trained for at least 2 weeks before starting 2-photon calcium imaging.

Value-based decision task

In the value-based decision task, a reward is probabilistically assigned to each choice. On each trial, a reward may be assigned to each choice according to the reward assignment probabilities that are different between two choices. Once a reward was assigned to a lickport, the reward remained assigned until it was chosen. As a result, the probability that a reward is assigned to a choice gradually increases if the choice has not been selected in the recent past trials. The combinations of reward assignment probabilities were either [60%, 10%] or [52.5%, 17.5%] in a trial, and reward assignment probabilities switched randomly every 60-80 trials in the order of [Left, Right] = ..., [60%, 10%], [10%, 60%], [52.5%, 17.5%], [17.5%, 52.5%], [60%, 10%], The probability switch was postponed if the fraction of choosing the lickport with higher reward assignment probability was below 50% in recent 60 trials until





the fraction reached at least 50%. ITI was 5-7 s, and the trials with licking during ready period were classified as alarm trials (500 ms white noise alarm sound and extra 2 s ITI). Trials in which mice licked during ready period ('alarm trials', 5.15%) and the trials in which mice failed to lick during the answer period ('miss trials', 4.68%) were not rewarded. We did not include alarm and miss trials in neural activity analyses to ensure that the ready periods we analyzed were free of licking behaviors and that mice were engaged in the task in the trials.

Two-photon calcium imaging

We used a two-photon microscope (B-SCOPE, Thorlabs) with a 16 × objective (0.8 NA, Nikon) and 925 nm excitation wavelength (Ti-Sapphire laser, Newport) for *in vivo* calcium imaging. Images were acquired using ScanImage (Vidrio Technologies) running on MATLAB. All calcium imaging was performed using camk2-tTA::tetO-GCaMP6s double transgenic mice that express GCaMP6s in camk2-positive excitatory neurons. Each field-of-view (FOV) (512 × 512 pixels covering 524 × 524 µm) was scanned at ~29 Hz. Areas within the FOV that were not consistently imaged across frames were discarded from analyses (Typically 10 pixels from each edge of the FOV). We imaged and analyzed layer 2/3 neurons of 6 cortical areas in this study: retrosplenial (RSC, 0.4 mm lateral and 2 mm posterior to bregma), posterior parietal (PPC, 1.7 mm lateral and 2 mm posterior to bregma), posterior premotor (pM2, 0.4 mm lateral and 0.5 mm anterior to bregma), anterior lateral motor (ALM, 1.7 mm lateral and 2.25 mm anterior to bregma), primary somatosensory (S1, 1.8 mm lateral and 0.75 mm posterior to bregma), and primary visual (V1, 2.5 mm lateral and 3.25 mm posterior to bregma) cortex. Images from these areas were collected from both hemispheres. We collected only 1 population from each hemisphere for each cortical area of a single mouse. We imaged both hemispheres in two different behavioral sessions if the FOVs on both hemispheres were clear at the time of imaging.

Image processing

Images from 2-photon calcium imaging were processed using a custom-written pipeline (Hattori, 2021). The pipeline corrects motion artifacts using pyramid registration (Mitani and Komiyama, 2018), and slow image distortions were further corrected by affine transformations based on enhanced correlation coefficients between frames (Evangelidis and Psarakis, 2008). We used Suite2P (Pachitariu et al., 2016) to define regions of interests (ROIs) corresponding to individual neurons and extract their GCaMP fluorescence. We selected only cellular ROIs using a user-trained classifier in Suite2P and by manual inspections. At the step of signal extraction from each cellular ROI, we excluded pixels that overlap with the other ROIs.

Neural activity

The neural activity data for the value-based decision task were first reported in ref. (Hattori et al., 2019). We also additionally collected new neural activity data from mice running the alternate choice task. The activity was continuously recorded with *in vivo* two-photon calcium imaging at ~29 Hz from mice during the task performance. GCaMP fluorescence time series were deconvolved to obtain signals that better reflect the kinetics of neural spiking activity using a non-negative deconvolution algorithm (Friedrich et al., 2017; Pachitariu et al., 2018). The deconvolved signal of each neuron was z-score normalized using the activity time series during the entire imaging session before performing all the activity analyses in this study.

For the alternate choice task, we collected and analyzed the activity of 8,524 RSC neurons (14 populations), 3,186 PPC neurons (7 populations), 7,915 pM2 neurons (14 populations) and 4,911 ALM neurons (10 populations) from 9 expert mice while they were running the alternate choice task. For the value-based decision task, we analyzed the activity of 9,254 RSC neurons (15 populations), 6,210 PPC neurons (13 populations), 7,232 pM2 neurons (13 populations) and 5,498 ALM neurons (10 populations) from early sessions (\leq 6th session), and 9,992 RSC neurons (populations), 7,703 PPC neurons (populations), 9,759 pM2 neurons (populations), 6,721 ALM neurons (populations), 7,576 S1 neurons (14 populations) and 2,767 V1 neurons (6 populations) from expert sessions of the data used in ref. (Hattori et al., 2019).

Reinforcement learning model for mouse behaviors

The reinforcement learning model that we used to estimate the action values in each trial was taken from ref. (Hattori et al., 2019). This model was optimized specifically for mouse behaviors and not necessarily ideal for describing the RL action policy of artificial neural network agents (e.g., Optimal RNN agents). Action values of chosen (Q_{ch}) and unchosen (Q_{unch}) options in each trial were updated as follows:

$$Q_{ch}(t+1) = \begin{cases} Q_{ch}(t) + \alpha_{rew} * (R(t) - Q_{ch}(t)) \text{ if rewarded } (R(t) = 1) \\ Q_{ch}(t) + \alpha_{unr} * (R(t) - Q_{ch}(t)) \text{ if unrewarded } (R(t) = 0) \end{cases}$$
 [eq.1]

$$Q_{unch}(t+1) = (1-\delta) * Q_{unch}(t)$$
 [eq.2]

where α_{rew} and α_{unr} are the learning rates for rewarded and unrewarded trials respectively, δ is the forgetting rate for the unchosen option, and R(t) is reward outcome in trial t (1 for rewarded, 0 for unrewarded trials). The learning rates and the forgetting rate were constrained between 0 and 1. In alarm and miss trials, values of both options were discounted by δ . The probability of choosing left (P_L) on trial t is estimated using left (Q_L) and right (Q_R) action values as follows:





$$P_{L}(t) = \frac{1}{1 + e^{-\beta_{\Delta O}(\beta_{0} + Q_{L}(t) - Q_{R}(t))}}$$
 [eq.3]

where β_0 is the value bias which is constant within each session, and $\beta_{\Delta O}$ reflects the behavioral sensitivity to ΔQ . The RL model was fit to the behavioral choice patterns with maximum likelihood estimation.

∆Q-coding neurons

ΔQ-coding neurons in the value-based decision task were identified with the following multiple linear regression model.

$$a_i(t) = \beta_C C(t) + \beta_{\Delta Q} \Delta Q(t) + \beta_{Q_{ch}} Q_{ch}(t) + \beta_{\Sigma Q} \Sigma Q(t) + \beta_0$$
 [eq.4]

where $a_i(t)$ is the mean activity of i^{th} neuron within each 200 ms time bin on trial t (except for some analyses (Figures S1 and S2) where the mean activity within the first 2 s of ready period was used instead), C(t) is the choice on trial t (1 if contralateral choice, -1 if ipsilateral choice), $\Delta Q(t)$ is the value difference between contralateral and ipsilateral options on trial t, $Q_{ch}(t)$ is the value of the chosen option on trial t, and $\sum Q(t)$ is the sum of values of both options on trial t. The regression weights were estimated by the ordinary leastsquares method. ΔQ -coding neurons were identified with two-tailed t test for the $\beta_{\Delta Q}$ regression weight (statistical threshold of either p < 0.05 or p < 0.01 as indicated in the figure legend of each analysis). The t-value for $\beta_{\Delta Q(t)}$ is $T_{\beta_{\Delta Q(t)}} = \frac{\beta_{\Delta Q}}{se(\beta_{\Delta Q})}$ where $se(\beta_{\Delta Q})$ is an estable point of the figure legend of each analysis. timate of the standard error of $\beta_{\Delta Q}$.

Action history coding neurons

Neurons that encode action history from an immediately preceding trial in the alternate choice task and the value-based decision task were identified with the following multiple linear regression model.

$$a_i(t) = \beta_{C_i}C(t) + \beta_{C_{(t-1)}}C(t-1) + \beta_0$$
 [eq.5]

where $a_i(t)$ is the mean activity of i^{th} neuron within each 200 ms time bin on trial t, C(t) is the choice on trial t (1 if contralateral choice, -1 if ipsilateral choice, C(t-1) is the choice on trial (t-1) (1 if contralateral choice, -1 if ipsilateral choice, 0 otherwise). The regression weights were estimated by the ordinary least-squares method. Action history coding neurons were identified with two-tailed t test for the $\beta_{C_{(t-1)}}$ regression weight (statistical threshold of p < 0.05). The t-value for $\beta_{C_{(t-1)}}$ is $T_{\beta_{C_{(t-1)}}} = \frac{\beta_{C_{(t-1)}}}{\sec(\beta_{C_{(t-1)}})}$ where $se(\beta_{C_{(t-1)}})$ is an estimate of the standard error of $\beta_{C_{(t-1)}}$.

Persistency index

Persistency index to quantify the mean persistency of ΔQ coding or action history coding in a population of neurons was defined as follow;

Persistency index =
$$\frac{\frac{1}{m}\sum_{j=1}^{m}\sum_{i=1}^{n}std(T_{shuffled}^{ij} \text{ sequence}) - \sum_{i=1}^{n}std(T_{raw}^{i} \text{ sequence})}{\frac{1}{m}\sum_{i=1}^{m}\sum_{i=1}^{n}std(T_{shuffled}^{ij} \text{ sequence}) - \sum_{i=1}^{n}std(T_{sorted}^{i} \text{ sequence})}$$
 [eq.6]

where T^i_{raw} sequence is the time series of t-values for $\beta_{\Delta Q}$ or $\beta_{C_{(t-1)}}$ that was obtained by fitting the [eq. 4] or [eq. 5] to the activity of each of the non-overlapping 200 ms time bins between 5 s before the ready cue and 2 s after the ready cue. The across-time standard deviation of the T_{raw}^{i} sequence was summed across all n neurons in the population (including neurons with non-significant t-values), and this summed standard deviation was normalized by min-max normalization such that the persistency index ranges between 0 (chance level persistency of a target population) and 1 (maximum persistency of a target population). The maximum persistency of a target population, $\sum_{i=1}^{n} std(T_{sorted}^{i} sequence)$, was obtained by independently sorting the cell identity at each time bin according

to the $\beta_{\Delta Q}$ t-values of each cell in the time bin. The chance level persistency of a target population, $\frac{1}{m}\sum_{j=1}^{m}\sum_{i=1}^{n}std\left(T_{shuffled}^{ij}sequence\right)$

was obtained by independently shuffling the cell identity at each time bin. To minimize the effect of randomness in the shuffling procedure, we iterated the shuffling m times (m = 10) and took the mean of the 10 iterations. This persistency index describes how persistent the target signal coding is above chance and how far the persistency is from the maximum persistency that the target population activity could achieve.

Demixed subspace principal component analysis (dsPCA)

Supervised dimensionality reduction algorithms can identify dimensions that encode targeted signals in high-dimensional data. However, they do not provide any information about signals that are not targeted by the users. As a result, these supervised analyses may miss important signals that exist in the original high-dimensional data. On the other hand, unsupervised dimensionality reduction algorithms can find dimensions for the major signals in the high-dimensional data, but they do not automatically reveal what kind of signals are reflected along each dimension. Furthermore, unsupervised methods may miss the signals of interest if the target signals are much weaker than the other dominant signals in the data.





We developed a novel dimensionality reduction algorithm that combines the strengths of both supervised and unsupervised methods. The demixed subspace principal component analysis (dsPCA) identifies demixed coding axes for targeted variables in a supervised manner, and then identify axes that capture the remaining variance in the data using an unsupervised method. Although previously reported demixed principal component analysis (dPCA) has similar objectives (Kobak et al., 2016), dPCA can only identify targeted coding axes for discrete variables. In contrast, dsPCA can identify demixed axes for both discrete and continuous variables. Furthermore, although dPCA splits each targeted signal into multiple linear axes, dsPCA identifies a single linear coding dimension for each of the target signals, and all the linear information for the target signals are contained within the dimensions identified by these single coding axes. The input to the algorithm is a 3^{rd} -order tensor of population activity with dimensions of Trial $(m) \times \text{Time } (t) \times \text{Neuron } (n)$.

$$X_{trial \times time \times neuron} = X_{m \times t \times n}$$
 [eq.7]

The tensor $X_{m \times t \times n}$ is first averaged over time axis elements within a specified time range, and we get a 2^{nd} -order tensor of $X'_{m \times n}$.

$$\mathbf{X}'_{m \times n} = \begin{pmatrix} x_{1,1} & x_{2,1} & \cdots & x_{n,1} \\ x_{1,2} & x_{2,2} & \cdots & x_{n,2} \\ \vdots & \vdots & \ddots & \vdots \\ x_{1,m} & x_{2,m} & \cdots & x_{n,m} \end{pmatrix}$$
 [eq.8]

To identify the demixed linear coding axes that encode ΔQ , Qch, or ΣQ in the population activity, we fit the following multiple linear regression model to the mean activity of individual neurons during the ready period;

$$\begin{pmatrix} x_{i,1} \\ x_{i,2} \\ \vdots \\ x_{i,m} \end{pmatrix} = \begin{pmatrix} 1 & \Delta Q_{1} & Q_{ch1} & \Sigma Q_{1} \\ 1 & \Delta Q_{2} & Q_{ch2} & \Sigma Q_{2} \\ \vdots & \vdots & \vdots & \vdots \\ 1 & \Delta Q_{m} & Q_{chm} & \Sigma Q_{m} \end{pmatrix} \begin{pmatrix} \beta_{i,0} \\ \beta_{i,\Delta Q} \\ \beta_{i,Q_{ch}} \\ \beta_{i,\Sigma Q} \end{pmatrix} + \begin{pmatrix} \varepsilon_{1} \\ \varepsilon_{2} \\ \vdots \\ \varepsilon_{m} \end{pmatrix}$$
[eq.9]

where $\beta_{i,\Delta O}$, $\beta_{i,Q_{ch}}$, and $\beta_{i,\Delta O}$ are the regression coefficients of the i^{th} neuron. For a population of n neurons, we obtain n regression coefficients for each type of Q-related signal. These regression coefficients are used to define the coding axes as follows;

$$\overrightarrow{\Delta q} = \begin{pmatrix} \Delta q_1 \\ \Delta q_2 \\ \vdots \\ \Delta q_n \end{pmatrix} = \frac{\overrightarrow{\beta_{\Delta Q}}}{\|\overrightarrow{\beta_{\Delta Q}}\|_2} = \frac{\left(\beta_{1, \Delta Q} \quad \beta_{2, \Delta Q} \quad \cdots \quad \beta_{n, \Delta Q}\right)^T}{\sqrt{\sum_{i=1}^n |\beta_{i, \Delta Q}|^2}}$$
 [eq.10]

$$\overrightarrow{\boldsymbol{q}_{ch}} = \begin{pmatrix} q_{ch1} \\ q_{ch2} \\ \vdots \\ q_{chn} \end{pmatrix} = \frac{\overrightarrow{\boldsymbol{\beta}_{\boldsymbol{Q}_{ch}}}}{\|\overrightarrow{\boldsymbol{\beta}_{\boldsymbol{Q}_{ch}}}\|_{2}} = \frac{\left(\beta_{1, Q_{ch}} \quad \beta_{2, Q_{ch}} \quad \cdots \quad \beta_{n, Q_{ch}}\right)^{T}}{\sqrt{\sum_{i=1}^{n} \left|\beta_{i, Q_{ch}}\right|^{2}}} \tag{eq.11}$$

$$\overrightarrow{\Sigma q} = \begin{pmatrix} \Sigma q_1 \\ \Sigma q_2 \\ \vdots \\ \Sigma q_n \end{pmatrix} = \frac{\overrightarrow{\beta_{\Sigma Q}}}{\|\overrightarrow{\beta_{\Sigma Q}}}\|_2 = \frac{\left(\beta_{1, \Sigma Q} \quad \beta_{2, \Sigma Q} \quad \cdots \quad \beta_{n, \Sigma Q}\right)^T}{\sqrt{\sum_{i=1}^n \left|\beta_{i, \Sigma Q}\right|^2}}$$
[eq.12]

Note that these coding axes are 'demixed' coding axes where the activity variance for partially correlated variables are demixed into one of the axes for the partially correlated variables thanks to the linear demixing in the regression model ([eq. 9]). Although some previous studies further orthogonalized these demixed coding axes (Mante et al., 2013), we did not orthogonalize between the coding axes because further orthogonalization would remix these best demixed coding axes.





Next, our goal is to identify a neural subspace that does not encode any of the targeted Q-related signals. To identify the neural subspace that is free of the 3 targeted Q-related signals, we solve the following full QR decomposition of an $n \times 3$ matrix with the 3 coding axis vectors using Householder reflections;

$$\begin{pmatrix} \Delta q_{1} & q_{ch_{1}} & \Sigma q_{1} \\ \Delta q_{2} & q_{ch_{2}} & \Sigma q_{2} \\ \vdots & \vdots & \vdots \\ \Delta q_{n} & q_{ch_{n}} & \Sigma q_{n} \end{pmatrix} = (\mathbf{S}_{\mathbf{Q}}, \mathbf{S}_{free})\mathbf{R}$$

$$= \begin{pmatrix} \overline{\mathbf{q}}_{1}, \overline{\mathbf{q}}_{2}, \overline{\mathbf{q}}_{3}, \overline{\mathbf{f}}_{1}, \overline{\mathbf{f}}_{2}, \cdots \overline{\mathbf{f}}_{(n-3)} \end{pmatrix} \mathbf{R}$$

$$= \begin{pmatrix} q_{1,1} & q_{2,1} & q_{3,1} & f_{1,1} & f_{2,1} & \cdots & f_{(n-3),1} \\ q_{1,2} & q_{2,2} & q_{3,2} & f_{1,2} & f_{2,2} & \cdots & f_{(n-3),2} \\ \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\ q_{1,n} & q_{2,n} & q_{3,n} & f_{1,n} & f_{2,n} & \cdots & f_{(n-3),n} \end{pmatrix} \mathbf{R}$$

where R is an upper triangular matrix, S_Q is a neural subspace that captures all Q-related signals, and S_{free} is the Q-free subspace that is orthogonal to the S_Q . S_Q is formed by 3 orthonormal basis vectors $(\overline{q_1}, \overline{q_2}, \overline{q_3})$, and these basis vectors and the 3 coding axis vectors $(\overrightarrow{\Delta q}, \overrightarrow{q_{ch}}, \overrightarrow{\Sigma q})$ span the identical neural subspace. On the other hand, \mathbf{S}_{free} is formed by (n-3) target-free orthonormal vectors $(\overrightarrow{f_1}, \overrightarrow{f_2}, \cdots \overrightarrow{f_{(n-3)}})$ and capture all the remaining population activity variance that were not captured by the subspace S_Q . The representation of the population activity $X'_{m \times n}$ in S_{free} is given by

$$proj_{\mathbf{S}_{free}}\mathbf{X}' = \mathbf{X}'\mathbf{S}_{free}$$
 [eq.14]

Lastly, we further realign the dimensions of the Q-free subspace Sfree such that minimum numbers of dimensions are necessary to explain the remained activity variance as much as possible. This realignment is done using the principal component vectors from PCA on $proj_{S_{tree}}X'$. The top p principal component vectors (p $\leq n$ - 3) can be used as the major Q-free subspace dimensions for dimensionality reduction purpose as follows;

$$F_{\rho}' = X'(S_{\text{free}}W_{\rho}^{\ \ pca}) = X'W_{\rho}^{\ \ dspca}$$
 [eq 15]

where the m-by-p matrix ${\bf F_p}'$ is the top p principal components of the activity within the Q-free subspace, the (n-3)-by-p matrix ${\bf W_p}^{pca}$ is the loadings matrix of the dsPCA. The columns of ${\bf W_p}^{dspca}$ are the Q-free axis vectors in the raw n-dimensional population activity space. More generally, the neural subspace that is free of k targeted variables can be obtained by the same [eq 15] with $p \le n - k$.

Through these steps ([eq 7] \sim [eq 15]), dsPCA identified the 3 linearly demixed coding axes for the targeted Q-related signals ($\overline{\Delta q}$). $\overrightarrow{q_{ch}}$, $\overrightarrow{\Sigma q}$), and (n-3) target-free axes (column vectors of W_{n-3}^{dspca}). We confirmed that none of the targeted signals could be linearly decodable from the population activity within the obtained target-free subspace (Figures 3E, 3I, and S3C).

In this manuscript, we decomposed neural population activity into demixed Q subspace and Q-free subspace using dsPCA. The Q subspace consists of demixed linear coding axes for ΔQ , Qch and ΣQ , and all activity variance that linearly relates to these Q-related signals are included in this subspace. On the other hand, all the other activity variance that did not remain in the Q subspace is included in the Q-free subspace. The activity state of the neural population changes across trials within the Q subspace depending on how each of the Q-related signals is updated by choice and its outcome. We also identified the axes that capture the major withintrial temporal activity variance in the Q-free subspace by performing PCA on the 2nd-order tensors that are obtained by averaging proj_{Sfree}X over trial axis elements.

Quantification of Q-related signals in subspaces from dsPCA

dsPCA decomposed population activity into Q subspace and Q-free subspaces. We examined the amount of Q-related signals in each subspace. The strength of Q-related signals in a full population activity with n neurons was quantified using linear decoders given by

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$$\Delta Q(t) = \sum_{i=1}^{n} \beta_i^{\Delta Q} \mathbf{a}_i(t) + \beta_0^{\Delta Q}$$
 [eq.16]

$$Q_{ch}(t) = \sum_{i=1}^{n} \beta_{i}^{Q_{ch}} a_{i}(t) + \beta_{0}^{Q_{ch}}$$
 [eq.17]

$$\Sigma Q(t) = \sum_{i=1}^{n} \beta_i^{\Sigma Q} a_i(t) + \beta_0^{\Sigma Q}$$
 [eq.18]

where $a_i(t)$ is the activity of the i^{th} neuron on trial t, β_i^x is the regression weight for $a_i(t)$, and β_0^x is the constant term. The decoder was trained with an L2 penalty by selecting the regularization parameter by 5-fold cross-validation. The decoding accuracy was obtained with 5-fold cross-validation by separating trials into training and test sets. Similarly, the strength of Q-related signals in the 3-dimensional Q subspace and the (n-3)-dimensional Q-free subspaces were quantified using linear decoders on the projected population activity in each subspace as follows;

$$\Delta Q(t) = \sum_{i=1}^{x} \beta_i^{\Delta Q} s_i(t) + \beta_0^{\Delta Q}$$
 [eq.19]

$$Q_{ch}(t) = \sum_{i=1}^{x} \beta_{i}^{Q_{ch}} \mathbf{s}_{i}(t) + \beta_{0}^{Q_{ch}}$$
 [eq.20]

$$\Sigma Q(t) = \sum_{i=1}^{x} \beta_i^{\Sigma Q} s_i(t) + \beta_0^{\Sigma Q}$$
 [eq.21]

where $s_i(t)$ is the population activity along the i^{th} dimension of the subspace on trial t, β_i^x is the regression weight for $s_i(t)$, and β_i^x is the constant term. x = 3 for Q subspace while x = n - 3 for Q-free subspace. These analyses revealed that all Q-related signals were captured by the Q-subspace, while Q-related signals were completely absent in the Q-free subspace (Figures 3E and 3I and S3C).

RNN agents with optimal or mouse-like RL strategy

The RNN agents trained to perform RL in this study consisted of 2 neurons in the input layer, 100 neurons in the recurrent layer, and 1 neuron in the output layer. The agents were trained to perform RL in the same behavior task environment with 10 time steps per trial. The 2 input neurons receive choice and reward outcome information only at the time step immediately after choice, and the history of the choice outcome information was maintained through the recurrent connectivity in the downstream recurrent layer. The sequence of activity fed into the input neurons was given as vectors with either choice or reward history labels in their elements. The elements that correspond to the time steps immediately after choice took 1 for left choice and -1 for right choice in the choice history vector, and the elements took 1 for reward outcome and -1 for no-reward outcome in the reward history vector. These elements took 0 in miss trials. The other elements of the vectors were all zeros. We sequentially fed 100 time steps of sequences into these input neurons, and the network training was done with unroll length of 100 time steps for backpropagation through time. The choice input neuron and reward input neuron connect with neurons in the recurrent layer. The neurons in the recurrent layer are connected with each other through recurrent connections, which allows each recurrent neuron to receive outputs of the previous time steps. The output of the recurrent layer is given by

$$\mathbf{y}_{(t)} = tanh(\mathbf{W}_{x}\mathbf{x}_{(t)} + \mathbf{W}_{y}\mathbf{y}_{(t-1)} + \mathbf{b})$$
 [eq.22]

where $tanh(\cdot)$ is a hyperbolic tangent activation function of the form $tanh(z) = \frac{e^z - e^{-z}}{e^z + e^{-z}}$, $\mathbf{x}_{(t)}$ is a 2 × 1 vector containing the choice and reward information from a previous time step, $y_{(t-1)}$ is a 100 \times 1 vector containing the layer's outputs at time step t, W_x is a 100 \times 2 matrix containing the connection weights for the inputs of the current time step, W_v is a 100 \times 100 matrix containing the connection weights for the outputs of the previous time step, and \boldsymbol{b} is a 100 \times 1 vector containing each neuron's bias term. The recurrent neurons send their outputs to the output neuron. The output neuron calculates the probability of selecting left action in the trial with a sigmoid activation function of the form $\sigma(z) = \frac{1}{1+e^{-z}}$. The agent then selects an action for the trial probabilistically by following the choice probability from the output neuron. This 3-layer RNN agent was trained to perform either an optimal RL strategy or the RL strategy that mice develop after training using its recurrent activity dynamics.

To train the RNNs to perform optimal RL in the task environment, we directly utilized the reward assignment rule of the task. In the value-based decision task, a reward is assigned to each choice according to the reward assignment probabilities of each choice on each trial. Once a reward was assigned to a lickport, the reward was maintained on the choice until it was chosen by the animal. As a result, the probability that a reward is assigned to a choice gradually increases if the choice has not been selected in the recent trials. The actual cumulative reward probabilities of left and right choices are given by





$$P_L(t) = 1 - \prod_{x=t-N_D(t)}^{t} \{1 - A_L(x)\}$$
 [eq.23]

$$P_R(t) = 1 - \prod_{x=t-N_I(t)}^{t} \{1 - A_R(x)\}$$
 [eq.24]

where $A_c(x)$ is the reward assignment probability of choice c on trial x, $N_c(t)$ is the number of successive c choices before trial t (e.g., $N_B(t) = 3$ when the choice on (t-4) was left and the choices on (t-3), (t-2), (t-1) were right). Therefore, an optimal choice generator would select a choice with higher cumulative reward probability on each trial as follows;

Optimal choice =
$$argmax{P_c(t)}$$
 [eq.25]

We used this optimal choice generator as the teacher to train RNNs to learn a near-optimal RL strategy. Unlike the optimal choice generator that knows the exact reward assignment probabilities $(A_c(x))$ and the reward assignment rule, the RNNs are agnostic to these hidden variables. Therefore, our goal is to train the RNNs to use only the past choice and reward history to make choices that are similar to the choices made by the optimal choice generator. To train the RNNs to imitate the behaviors of the optimal choice generator, we calculated binary cross-entropy as the loss function to be minimized. The cross-entropy is given by

$$H_p = -\frac{1}{M} \sum_{i=1}^{M} \left(a_i^{optimal} log(p_i^{RNN}) + (1 - a_i^{optimal}) log(1 - p_i^{RNN}) \right)$$
 [eq.26]

where M is the total number of training trials, $a_i^{optimal}$ is 1 or 0 when the optimal choice generator selected left or right action on the i^{th} trial respectively, and p_i^{RNN} is the left choice probability of the RNN agent from its output neuron.

To train RNNs to perform mouse-like RL that is suboptimal in the task environment, we used 50,472 decision making trials of expert mice in the task environment. We fed the choice and reward history that expert mice experienced into the RNNs, and trained the RNNs to imitate the choice patterns of expert mice. To do this, we calculated the binary cross-entropy as the loss function to be minimized. The cross-entropy is given by

$$H_p = -\frac{1}{M} \sum_{i=1}^{M} \left(a_i^{mouse} log(p_i^{RNN}) + (1 - a_i^{mouse}) log(1 - p_i^{RNN}) \right)$$
 [eq.27]

where M is the total number of training trials, a_i^{mouse} is 1 or 0 when the expert mouse selected the left or right action in the i^{th} trial respectively, and p_i^{RNN} is the left choice probability of the RNN agent from its output neuron.

For the training of both the optimal RNN agents and mouse-like RNN agents, the cross-entropy loss was calculated at variable time steps for each trial to reflect the temporal variability of the timing of decision making in this task (variable ITI, variable ready-period, variable reaction time), and all the synaptic weights of the RNN agent were trained with backpropagation through time. The training was optimized using mini-batch gradient descent with Nesterov momentum optimization (learning rate of 0.001 and momentum of 0.9. batch size of 128), and the training was terminated when the loss for a validation set (1/5 of trials) stopped decreasing for the consecutive 50 epochs as a form of regularization (Early stopping). The trained RNN agents ran the task in a simulated environment with the length of 500 trials/session, and the RL behavioral strategy in the simulated environment was quantified by a RL model optimized to describe expert mouse behaviors [eq. 1 –3] and a logistic regression model [eq. 28].

Quantification of history-dependent behavioral strategy

The quantification of behavioral strategy for mice and RNN agents was performed with either a RL model [eq. 1 – 3] or a logistic regression model [eq. 28]. The logistic regression model predicts an action in each trial based on 3 types of history from the past 10 trials. The model is given by

$$logit(P_{L}(t)) = \sum_{i=1}^{10} \beta_{RewC(t-i)} * RewC(t-i) + \sum_{i=1}^{10} \beta_{UnrC(t-i)} * UnrC(t-i)$$

$$+ \sum_{i=1}^{10} \beta_{C(t-i)} * C(t-i) + \beta_{0}$$
[eq.28]

where $P_L(t)$ is the probability of choosing left on trial t, RewC(t-i) is the rewarded choice history on trial t-i (1 if rewarded left choice, -1 if rewarded right choice, 0 otherwise), U of the unrewarded choice history on trial t-i (1 if unrewarded left choice, -1 if unrewarded right choice, 0 otherwise), C(t-i) is the outcome-independent choice history on trial t-i (1 if left choice, -1 if right choice, 0 otherwise). $\beta_{RewC(t-i)}$, $\beta_{UnrC(t-i)}$, and $\beta_{C(t-i)}$ are the raw regression weights of each history predictor, and β_0 is the history tory-independent constant bias term. The sizes of these raw weights reflect the relative contribution of each history variable to decision making in a behavior session. However, the weight size does not reflect the absolute strength of the contribution to decision





making because the strength of each history effect on decision making is determined by not only the regression weight but also the choice prediction accuracy of the regression model. Therefore, we normalized the regression weights by the choice predictability of the regression model as follows;

Normalized
$$\beta_{x} = \left(\frac{N_{choice}^{correct}}{N_{choice}^{all}} - 0.5\right) * \frac{\beta_{x}}{\sum_{i=1}^{10} (\left|\beta_{RewC(t-i)}\right| + \left|\beta_{UnrC(t-i)}\right| + \left|\beta_{C(t-i)}\right|) + \left|\beta_{0}\right|}$$
 [eq.29]

where N_{choice}^{all} is the number of choice trials in the session, and $N_{choice}^{correct}$ is the number of choice trials that were correctly predicted by the [eq. 28]. Each regression weight is divided by the sum of absolute values of all the regression weights before being multiplied by the choice prediction accuracy. This normalization turns raw regression weights to reflect the fraction of choice predictability by each of the history variable. These normalized weights are comparable across different behavior sessions or mice because they reflect the absolute strength of each history event on decision making. We used these normalized weights to compare the history dependence of expert mice and trained RNN agents for their decision making.

Artificial population activity sequence

Artificial population activity sequences with either persistent or non-persistent rate coding of ΔQ were created based on the distributions of the tuning curves of ΔQ coding among RSC neurons. Each population consisted of 200 neurons with 5 time bins, and we assigned 20% of neurons at each time bin to encode ΔQ . The tuning curve slope of ΔQ coding of each RSC neuron $(\beta_{\Delta Q})$ was measured by fitting [eq. 4] to the activity during ready period. We defined across-trial standard deviation of $\beta_{\Delta O} \Delta Q(t)$ from [eq. 4] as the signal standard deviation of ΔQ coding. To derive the noise standard deviation, we first subtracted $\beta_{\Lambda Q}\Delta Q(t)$ from the ready period activity sequence of each trial. The residual ready period activity sequences were then concatenated across trials. The standard deviation of the concatenated activity sequence was defined as the noise standard deviation. The SNR of ΔQ coding was defined as the ratio of the signal standard deviation to the noise standard deviation. The tuning curve slope for each activity time bin was randomly sampled without replacement from the distributions of ΔQ-coding neurons. The ΔQ signal was linearly encoded at each time bin according to the sampled tuning curve slope, and additional Gaussian noise was added to the neural activity. The other non-ΔQ coding activity time bins simply exhibited Gaussian noise. We created populations with 3 different types of rate coding modes (Persistent, Non-persistent 1, Non-persistent 2). In the populations with Persistent mode, the identical 20% of neurons encoded ΔQ at all 5 time bins. In the populations with Non-persistent 1 mode, we randomly selected 20% of neurons at each time bin as the ΔQ -coding neurons and allowed each neuron to encode ΔQ with different tuning curve slopes at different time bins. Non-persistent 2 mode is similar to Non-persistent 1, except that each neuron in the population encoded ΔQ at only one of the time bins.

In addition to the 3 rate coding schemes, we also considered a coding mode that encodes ΔQ as specific sequential activity patterns across cells in a population. In this 3^{rd} non-persistent coding mode (*Non-persistent 3*), neural activity at each time bin can take only binary states (0: inactive, 1: active). Therefore, this population encodes ΔQ using only the identity of active cells. We encoded 10 different sequences in a population such that each sequence uniquely corresponds to one of the 10 binned ΔQ (-1 to 1 with binning of 0.2 width). For each sequence, we randomly assigned 20% of neurons at each time bin as active neurons with a constraint that each neuron can be active only at a single time step in a sequence. After encoding the 10 different sequences in a population, we added Gaussian noise to the activity of each neuron. We defined the SNR of this coding scheme as the ratio of the across-time standard deviation of the activity of a neuron to the standard deviation of its added Gaussian noise.

△Q retrieval by RNN

RNNs were trained to retrieve ΔQ information from the input population activity sequence. The RNN had 40 recurrent neurons with tanh activation functions and an output neuron with linear activation function. The network weights were updated by backpropagation through time with RMSprop to minimize mean-squared-error (MSE) between the network outputs and ΔQ values of the trials in a training set. The network training was terminated when the MSE of a validation set stopped decreasing for the consecutive 20 epochs as a form of regularization (Early stopping). For each training iteration, we used 20% of available trials as a test set to calculate the ΔQ retrieval accuracy by the trained network, and the remaining 80% of the trials were further split into validation set (10%) and training set (70%). We repeated the network training 5 times by using different sets of trials as the test set such that we can obtain ΔQ predictions by the trained networks for all available trials in a cross-validated way. The ΔQ retrieval accuracy was calculated by comparing the ΔQ predictions to the true ΔQ from the RL model. For the ΔQ retrieval from cortical activity, we used only 240 cells as the inputs to match the number of cells across different cortical areas. For each neural population, we subsampled 240 cells in each iteration allowing repetitions with the smallest number of iterations to include every cell at least once for decoding, and the ΔQ retrieval accuracy from the iterations were averaged.

Denoising recurrent autoencoder

Autoencoder is an artificial neural network that learns to extract efficient coding of its input without supervision. It consists of an encoder network and a decoder network, and they are sequentially connected through a coding layer with small number of neurons. In a trained autoencoder, the encoder extracts essential signals in the input into the coding layer, while the decoder tries to reconstruct





the original input from activity in the coding layer. When the number of neurons in the coding layer is smaller than the dimensions of the input, only signals that are dominant in the input remains in the coding layer of a trained autoencoder network. Among various types of autoencoders, we used denoising recurrent autoencoders (Maas et al., 2012; Vincent et al., 2010) to extract dominant signals embedded in each population activity sequence. Although autoencoders with only feedforward connections or convolutional neural networks can also extract latent signals in a population activity sequence, we used recurrent neural networks that sequentially process the input activity because the neural networks in a brain also process input activity sequentially. Our goal is to understand whether such biologically relevant recurrent networks can extract signals from input activity without explicit teaching labels (i.e., unsupervised learning). The latent signals extracted by a recurrent autoencoder represent the latent signals from the perspective of a recurrent network that processes input activity sequentially through its recurrent connectivity.

The autoencoders that we used to visualize extracted dynamics from example populations (Figure 7A) consisted of 3 hidden layers with recurrent connectivity (1st: 50 neurons, 2nd: 10 neurons, 3rd: 50 neurons), and the activity of all neurons in a population was used as the input to the autoencoder. On the other hand, the autoencoders that we used for quantitative across-area comparisons (Figure 7B-E) consisted of 3 hidden layers with recurrent connectivity (1st: 20 neurons, 2nd: N neurons, 3rd: 20 neurons) and processed input activity of subsampled 240 cells. Note that 3 layers are the minimum number of layers that are required for an autoencoder network. All recurrent neurons in the hidden layers had tanh activation functions. All neurons except for the neurons in the middle hidden layer (coding layer) sent activity sequentially to the neurons in the next layer. However, the neurons in the coding layer sent only the activity at the last time step to the next hidden layer. The last-time-step activity is the result of the temporal integration of the original population activity sequence through recurrent connectivity, and the activity reflects the latent representations in the original population activity sequence. The hidden layers after the coding layer reconstructed the original population activity sequence from the latent representations in the coding layer. The network weights were updated by backpropagation through time with RMSprop to minimize mean-squared-error (MSE) between the original population activity sequence and the reconstructed population activity sequence. To ensure stable training of network weights, we clipped the gradients of network weights if their L2 norms were greater than 1 (Gradient clipping (Pascanu et al., 2012)). To add noise robustness to the autoencoders, we applied dropout (Hinton et al., 2012; Srivastava et al., 2014) to the connections between the input neurons and the neurons in the 1st hidden layer such that 50% of randomly selected connections are ablated at each training step. The network training was terminated when the MSE of a validation set (20% of trials for Figure 7A, 10% of trials for Figure 7B-E) stopped decreasing for the consecutive 20 epochs as another form of regularization (Early stopping). The activity of the 10 coding neurons for Figure 7A were further reduced to 2 dimensions with multidimensional scaling to visualize the dominant population activity states. To quantify the strength of ΔQ signal in the activity of N coding neurons for Figure 7B-E, we performed decoding of ΔQ from the activity of N coding neurons using a simple feedforward neural network where all the N coding neurons are connected to an output neuron with tanh activation function. For each training iteration, we used 20% of available trials as a test set to calculate the ΔQ decoding accuracy by the trained network, and the remaining 80% of the trials were further split into validation set (10%) and training set (70%). We repeated the network training 5 times by using different sets of trials as the test set such that we can obtain ΔQ predictions by the trained networks for all available trials in a cross-validated way. For these ΔQ decoding analyses, we also matched the number of cells included in the inputs to the autoencoders across different decoding by subsampling 240 cells from the original population. For each neural population, we subsampled 240 cells in each iteration allowing repetitions with the smallest number of iterations to include every cell at least once for decoding, and the ΔQ decoding accuracy from the iterations were averaged.

Deep RNN with non-specific connectivity

Neural networks with 5 recurrent layers were used to simulate how the input population activity transforms in the downstream recurrent layers when the synaptic weights are non-specific throughout the networks. Each recurrent layer had 1,000 neurons with *tanh* activation functions, and the 5 recurrent layers were sequentially connected through feedforward connections. All neurons of a recorded cortical population were directly connected to the 1st recurrent layer. Each neuron in a recurrent layer was connected with all the other neurons in the same layer, but we made the connections between successive layers sparse by setting the connection probability of a neuron to the neurons in the next layer to 1%, 5%, 10%, 20%, or 50%. The non-specific synaptic weights were randomly drawn from a uniform distribution on [-1, 1).

Anatomical connectivity analyses

We analyzed neural projections from the areas with high ΔQ coding persistency (RSC, PPC, pM2) using the neural tracing data available in the Allen Mouse Brain Connectivity Atlas (Oh et al., 2014). These projection data were originally acquired by injecting adenoassociated virus (AAV) encoding EGFP into various target brain areas and scanning EGFP-labeled axons throughout the brain with high-throughput serial 2-photon tomography. We used their software development kit (SDK), *allensdk*, to access and process their data in Python.

Dorsal view of the Allen Reference Atlas

Allen Reference Atlas is a high-resolution anatomical 3D reference atlas for the adult mouse brain. Different brain structures are colored differently in this atlas. All projection data in the Connectivity Atlas are registered to this reference atlas. We created a dorsal view of the Allen Reference Atlas to indicate the virus injection coordinates and cortical projection density in the dorsal cortex. First,





we downloaded the 3D RGB-colored atlas at the resolution of 25 μm/pix. At each anterior-posterior (AP) and medial-lateral (ML) coordinate of the 3D atlas, we picked up the RGB value of the most dorsal brain surface. We obtained a dorsal view of the atlas by projecting these dorsal RGB values onto a single 2D plane.

Selection of injection data

In the Allen Mouse Brain Connectivity Atlas, each injection experiment is labeled with the name of the injected structure. First, we narrowed injection experiments using these annotations. We selected experiments with virus injections into retrosplenial area (RSP), anterior area (VISa) of posterior parietal association area (PTLp), and secondary motor area (MOs). Then, we further narrowed down injection experiments based on the exact injection coordinates. As we indicated in Figure S9, we isolated medial RSP injections, anterior VISa injections, and posterior MOs injections for RSC, PPC, and pM2, respectively. The database contains experiments that were performed on wild-type mice and Cre transgenic mice for cell-type specific tracing. We used experiments from only WT mice or combined data (WT + Cre). The projection patterns were similar in both cases (Figure S9).

Axon projection density in dorsal cortex

We analyzed the axon projection density from RSC, PPC, and pM2 in the dorsal cortex. For each injection experiment, we calculated the projection density at each AP-ML coordinate as [# of positive pixels] / [# of all pixels] in the volume of 25 μm (AP axis) × 25 μm (ML axis) × 1000 μm (DV axis, from dorsal surface at each AP-ML coordinate). To create a mean projection density map, experiments with left hemisphere injections were mirrored relative to midline before averaging. We also quantified mean projection density within each imaging FOV that we used for in vivo 2-photon calcium imaging. Although our imaging FOVs were based on stereotactic coordinates from the bregma in the Paxinos' atlas (Paxinos and Franklin, 2004), the Allen Reference Atlas does not include coordinates from the bregma. To register our imaging FOVs to the Allen Reference Atlas Coordinate, we calculated the scaling factors for the AP and ML dimensions of the mouse brain to match the brain in the Paxino's atlas to the brain in the Allen Reference Atlas. Using the scaling factors, we estimated the coordinates of each imaging FOV on the Allen Reference Atlas. We calculated the mean signal density within each imaging FOV of the size 500 μm × 500 μm. The mean signal density of each FOV was used to construct the connectivity matrix in Figure 8C.

QUANTIFICATION AND STATISTICAL ANALYSIS

All data analyses and network simulations were performed in Python3.7 with libraries of TensorFlow (Abadi et al., 2016), scikit-learn (Pedregosa et al., 2011), NumPy (Harris et al., 2020), SciPy (Virtanen et al., 2020), and Statsmodels (Seabold and Perktold, 2010). Statistical tests were performed either in Python with SciPy and Statsmodels or in R with its statistics libraries. All accuracy measures reported in this study were obtained with cross-validation. Unless otherwise noted, we split trials into training set (70%), validation set (10%), and test set (20%) for each iteration of decoding, and repeated the network training 5 times by using different sets of trials as the test set. When we compared ΔQ retrieval/decoding accuracy across different cortical populations, we matched the number of cells in the input population activity by subsampling 240 cells in each iteration allowing repetitions with the smallest number of iterations to include every cell at least once for decoding, and the accuracies from the iterations were averaged. For all the simulations with artificial population activity, we created 10 distinct populations for each of the 3 types of coding modes by independently sampling tuning curve slopes from RSC neurons. These repetitions allowed us to tell the variability that originates from the randomness of ΔQ signal assignments and randomness of network trainings. All figure plots were created using Matplotlib (Hunter, 2007) and seaborn (Waskom, 2021) in Python.

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Supplemental information

Context-dependent persistency as a coding mechanism for robust and widely distributed value coding

Ryoma Hattori and Takaki Komiyama

Figure S1. Distributed action value coding is independent of motor plan, Related to Figure 1.

- (A-F) The tuning curves of ΔQ coding averaged across neurons with significant ΔQ coding (Regression on mean ready-period activity (first 2 sec window), P < 0.05, two-sided t-test) for RSC (A), PPC (B), pM2 (C), ALM (D), S1 (E) and V1 (F). Neurons with positive and negative slopes were separately averaged. Error bars are 95% CI. The proportions of neurons with positive and negative slopes were balanced in each cortical area (RSC: 15.92 ± 1.69 % positive, 15.56 ± 1.63 % negative; PPC: 14.82 ± 1.69 % positive, 14.86 ± 1.59 % negative; pM2: 9.84 ± 1.22 % positive, 9.77 ± 1.15 % negative; ALM: 9.45 ± 0.99 % positive, 9.71 ± 0.95 % negative; S1: 9.70 ± 1.58 % positive, 10.41 ± 1.63 % negative; V1: 12.62 ± 3.05 % positive, 11.53 ± 2.80 % negative; mean \pm s.e.m). Note that these fractions are larger than the fractions reported in Fig. 1c because 2-sec window was used instead of 200-ms window to obtain mean activity.
- (G) Population activity states of an example RSC population during ready period in the subspace with ΔQ axis and choice axis from dsPCA. Mean \pm SD of the activity states of trials within each ΔQ bin. dsPCA identified this subspace where ΔQ and the motor plan for the upcoming decision making are encoded along different axes. (H) Activity of ΔQ coding neurons that were identified at different time windows (yellow shadings) during ready period for example PPC, pM2, ALM, and V1 populations. The activity of ΔQ coding neurons that were identified
- (I) t-values for ΔQ coding at each time bin of ready period for example populations of PPC, pM2, ALM, and V1. Neurons were sorted based on the t-values at the last time bin. t-values are from multiple linear regression model (Methods).

from each time bin was averaged according to the binned ΔQ of each trial. Error bars are s.e.m.

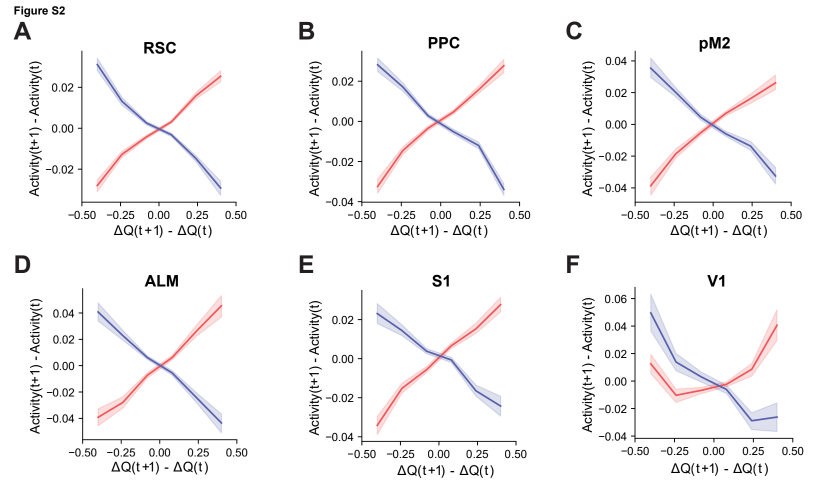


Figure S2. Distributed action value coding is updated at single-trial resolution, Related to Figure 1. (A-F) Change across adjacent trials in mean ready-period activity of neurons with significant ΔQ coding, plotted against the updates of ΔQ across adjacent trials for RSC (A), PPC (B), pM2 (C), ALM (D), S1 (E) and V1 (F). ΔQ coding neurons track updates of ΔQ on a trial-by-trial basis.

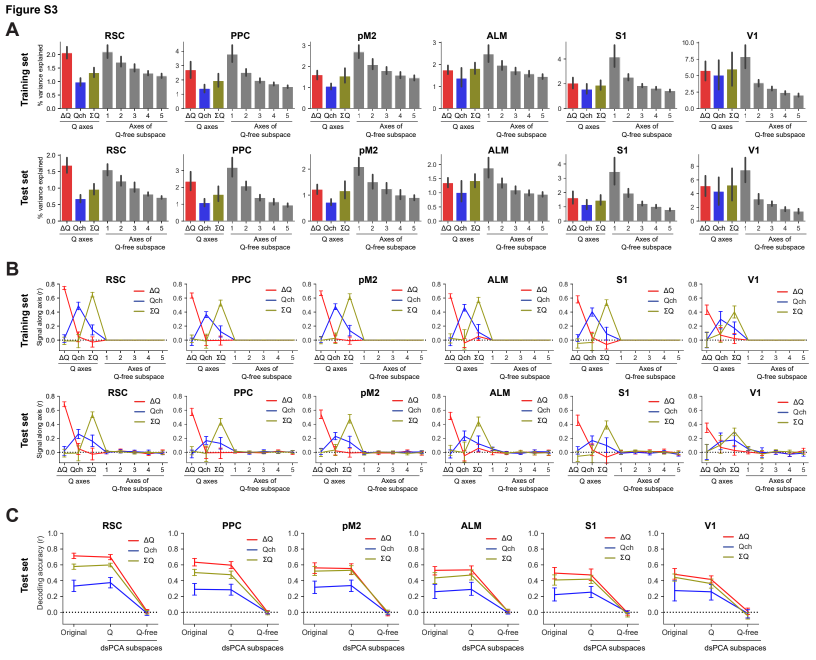


Figure S3. Demixing performance of dsPCA on the population activity from 6 cortical areas, Related to Figure 3.

(A) Fraction of activity variance out of total activity variance, along each Q-related axis and the top 5 PC axes from the Q-free subspace. dsPCA was applied on a subset of trials for RSC populations to identify axes (training set), and the performance on both the training set and the held-out trials (test set) are shown. Error bars are 95% CI.

(B) Q-related signals along each dimension identified with dsPCA for RSC populations. dsPCA was applied on a subset of trials to identify axes (training set), and the performance on both the training set and the held-out trials (test set) are shown. Pearson correlation coefficient was calculated between the projected activity and each signal. Error bars are 95% CI. Q-related signals are undetectable in Q-free subspace.

(C) Decoding accuracy of Q signals from the original population activity, activity in the Q subspace (3 dimensions), and activity in the target-free subspace. Linear decoders with L2 regularization were used. Error bars are 95% CI. Q-related signals are restricted to the Q subspace after dsPCA.

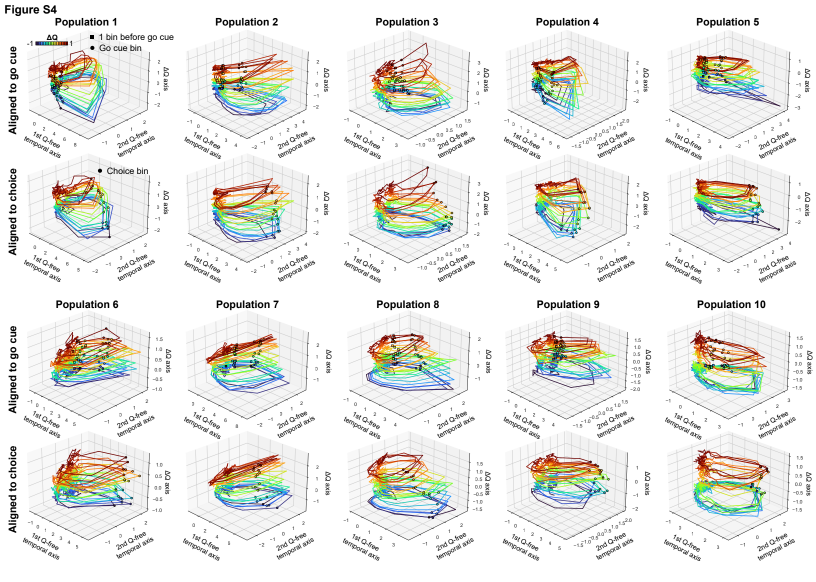


Figure S4. Population dynamics of 10 distinct RSC populations visualized by dsPCA, Related to Figure 3. The population activity dynamics of 10 distinct RSC neural populations in the neuronal manifolds where the ΔQ axis is paired with the axes that reflect major within-trial temporal activity variance in Q-free subspace. The neural trajectories were aligned to either go cue (top row) or choice (bottom row). The activity was binned with 200 ms non-overlapping sliding windows. In the trajectories aligned to go cue, the time bins that coincide with go cue and the time bins immediately preceding the go cue bin were marked by circles and squares, respectively. In the trajectories aligned to choice, the time bins that coincide with choice timing were marked by circles. The time from go cue to choice was 335.36 ± 9.12 ms (mean \pm s.e.m.).

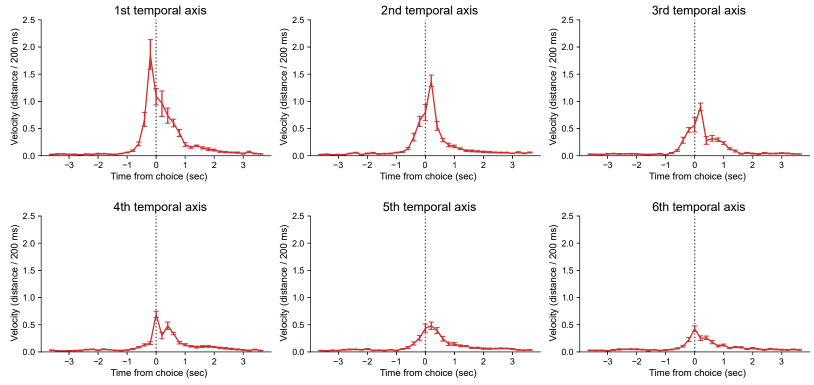


Figure S5. Velocity of RSC population dynamics along temporal axes, Related to Figure 3. Velocity of the RSC population activity dynamics along temporal axes identified within Q-free subspace. The velocity was calculated as the distance traveled between adjacent 200 ms time bins along each temporal axis. The RSC population trajectories accelerate ~1 sec before choice and slowly decelerate after choice. Error bars are s.e.m..

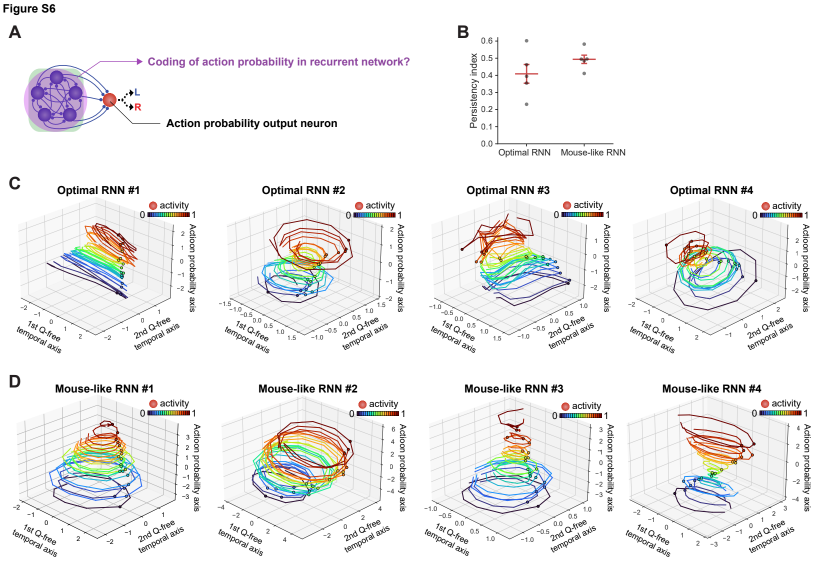


Figure S6. Population coding of action probability in the recurrent layers of trained optimal and mouse-like RNN agents, Related to Figure 4.

- (A) Activity of the action probability output neuron is the ground truth policy values (ΔQ) that directly drives decisions of artificial networks. We examined the representation of the action probability in the population of the recurrent layer presynaptic to the output neuron.
- (B) Persistency index for action probability coding in the recurrent layers of trained optimal and mouse-like RNN agents (mean ± CI).
- (C-D) Population activity dynamics of the recurrent layers of optimal RNN agents (C) or mouse-like RNN agents (D) in neuronal manifolds where the action probability axis is paired with axes that reflect major within-trial temporal activity variance. We analyzed the dynamics of fully trained networks while they ran the task for 10,000 trials. dsPCA was applied on the activity between -5 and -1 time steps from choice, and the population activity between -5 and +5 time steps from choice was projected onto the identified axes to visualize the temporal dynamics of activity within these manifolds. 4 independently trained mouse-like RNN agents are shown. Circles indicate the choice time. Untangled cylindrical dynamics emerged in the recurrent networks.

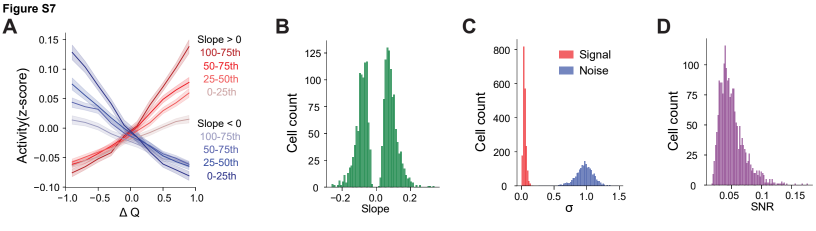


Figure S7. Distributions of the tuning curve slopes and signal-to-noise ratios of ΔQ coding among RSC neurons, Related to Figure 6.

- (A) Tuning curves of ΔQ coding neurons in RSC were separately averaged according to the binned percentile of the tuning curve slope, illustrating the heterogeneity of the tuning curve slopes.
- (B) Distribution of tuning curve slopes of ΔQ coding among RSC neurons with significant ΔQ coding in the mean 2-sec ready period activity (P < 0.01, two-sided t-test).
- (C) Distribution of the standard deviation of ΔQ signal (red) and noise (blue) of the ΔQ -coding RSC neurons (Methods).
- (D) Distribution of the signal-to-noise ratio (SNR) of ΔQ coding among RSC neurons with significant ΔQ coding.

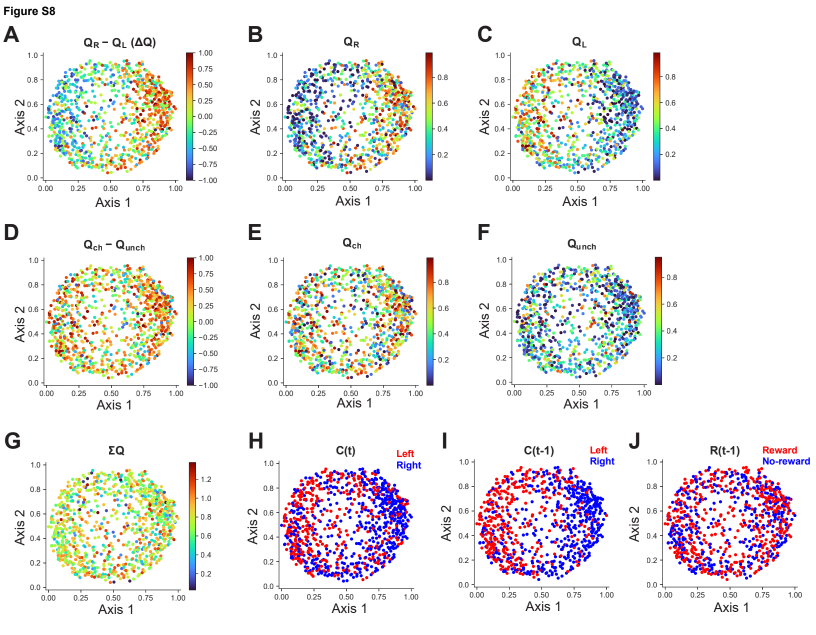


Figure S8. Task-related variables that are not directly related to ΔQ are not represented as the dominant signals in the activity of neurons in the coding layer of trained denoising recurrent autoencoder networks, Related to Figure 7.

Representation of input population activity in the coding layer of a denoising recurrent autoencoder network which was trained to extract major signals from an example RSC population activity. Population activity sequence during ready-period was used as the input. The autoencoders had 10 neurons in the coding layer. The dominant signals extracted in the activity of coding layer neurons were visualized by reducing the dimension to 2 using multidimensional scaling.

 Q_R-Q_L (ΔQ): Action value difference between right and left actions (A). Q_R : Right action value (B). Q_L : Left action value (C). These variable (ΔQ and its components) are clearly represented in these dominant dimensions. $Q_{ch}-Q_{unch}$: Action value difference between chosen and unchosen actions in the upcoming answer period (D). Q_{ch} : Chosen action value (E). Q_{unch} : Unchosen action value (F). Q_{unch} : Unchosen action value (F). Q_{unch} : Unchosen action value (F). Q_{unch} : Reward outcome in the previous trial (J). These variables that are not directly related to Q_{unch} are not clearly represented in these dimensions.

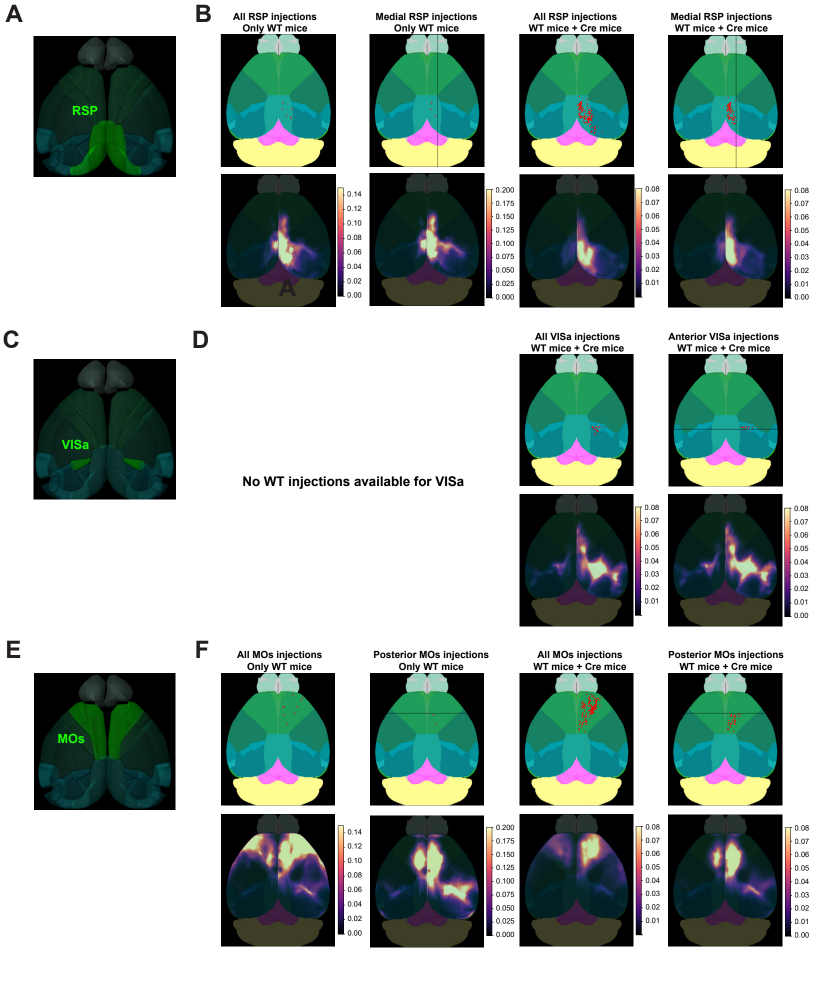


Figure S9. Neural projections from RSC, PPC and pM2 from the Allen Connectivity Atlas dataset, Related to Figure 8.

- (A) 3D model highlighting the area that is annotated as retrosplenial area (RSP) in the Allen Connectivity Atlas.
- (B) Injection coordinates (*top*) and the resulting mean neural projection density (*bottom*) for the injections that are annotated as 'RSP injections' in the database. Experiments with left hemisphere injections were mirrored horizontally. All injections from WT mice (6 experiments), medial injections from WT mice that are within the medial RSC from which RSC recording data were obtained (3 experiments), all injections from both WT and various Cre lines (111 experiments), medial injections from both WT and various Cre lines (60 experiments, this condition was used for Figure 8). Projection density at each AP-ML coordinate was calculated as [# of positive pixels] / [# of all pixels] in the volume of 25μm (AP axis) × 25μm (ML axis) × 1000μm (DV axis, from dorsal surface at each AP-ML coordinate).
- (C) 3D model highlighting the area that is annotated as VISa, which is an anterior subregion of posterior parietal association area (PTLp), in the Allen Connectivity Atlas.
- (D) Injection coordinates (*top*) and the resulting mean neural projection density (*bottom*) for the injections that are annotated as 'VISa injections' in the database. Experiments with left hemisphere injections were mirrored horizontally. All injections from various Cre lines (17 experiments), anterior injections from various Cre lines that are within anterior VISa from which PPC recording data were obtained (9 experiments, this condition was used for Figure 8). No experiments from WT mice was available for VISa injections in the database.
- (E) 3D model highlighting the area that is annotated as secondary motor area (MOs) in the Allen Connectivity Atlas.
- (F) Injection coordinates (*top*) and the resulting mean neural projection density (*bottom*) for the injections that are annotated as 'MOs injections' in the database. Experiments with left hemisphere injections were mirrored horizontally. All injections from WT mice (9 experiments), posterior injections from WT mice that are within posterior MOs, from which pM2 recording data were obtained (2 experiments), all injections from both WT and various Cre lines (145 experiments), posterior injections from both WT and various Cre lines (33 experiments, this condition was used for Figure 8).

Figure S10 В C D Α 15 15.0 1.0 Sorted Sorted Persistency index Raw Raw 12.5 Fraction (%) D2 Shuffled Shuffled D3 Fraction (%) 10.0 D4 7.5 5.0 2.5 0.0 0 0.0 0.1 0.5 0.9 1.3 1.7 D4 PPC D1 D2 D3 D4 D5 PPC D1 D2 D3 Time (sec) Non-specific connectivity F G Н Ε 1.0 12 Sorted Sorted рМ2 10.0 Raw Raw Persistency index 8.0 8.0 8.0 8.0 10 D2 Shuffled Shuffled Fraction (%) 7.5 Fraction (%) D3 6 5.0 2.5 2 0.0 0.0 0 0.1 0.5 0.9 1.3 1.7 pM2 D1 D2 D3 D4 D5 pM2 D1 D2 D3 Ď4 D5 Time (sec) Non-specific connectivity Connection probability from layer to layer 50% 10% 1% RSC D1 15 Fraction (%) D3 D4 D5 0.1 0.5 0.9 1.3 1.7 0.1 0.5 0.9 1.3 1.7 0.1 0.5 0.9 1.3 1.7 0.1 0.5 0.9 1.3 1.7 Time (sec) Time (sec) Time (sec) Time (sec) Connection probability from layer to layer 50% 10% 5% 1% 20 Sorted Sorted Sorted Sorted Raw Raw Raw Fraction (%) Raw Shuffled Shuffled Shuffled **Shuffled** RSC RSC RSC D1 D5 D1 D2 D1 D3 D2 D3 D4 RSC D1 D2 D3 Κ Connection probability from layer to layer 1.0 50% 10% 5% 1%

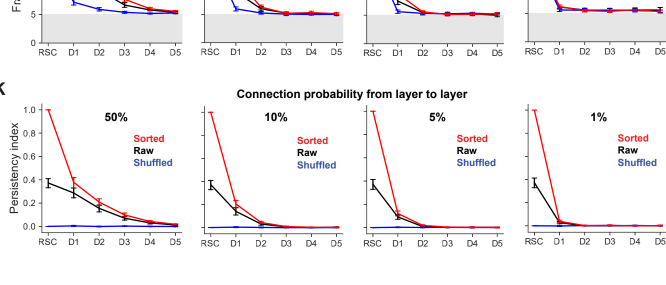


Figure S10. Signal leakage simulations with different areas and different connection probabilities, Related to Figure 8.

- (A) PPC population activity sequences were processed through 5 recurrent layers with non-specific connectivity. Connection probability from layer to layer was set to 20%.
- (B) Fractions of ΔQ coding neurons at each of the 200 ms time bins during ready period (Regression, P < 0.05, two-sided t-test). Error bars are s.e.m.
- (C) Mean fractions of ΔQ coding neurons at each layer during ready period. Fractions across time bins within the ready period were averaged for each population. Artificial manipulations of ΔQ coding persistency does not affect the fractions of ΔQ coding neurons in the area, but affect the fractions in the downstream areas. Error bars are 95% CI.
- (D) ΔQ coding persistency at each layer. Artificial manipulations of ΔQ coding persistency affect the persistency in the downstream areas. Error bars are 95% CI.
- (E-H) Same as (A-D) for pM2 populations.
- (I) Fractions of ΔQ coding neurons at each of the 200 ms time bins during ready period when the connection probability of feed-forward connections between layers was varied. RSC population activity was used. Error bars are s.e.m.
- (J) Mean fractions of ΔQ coding neurons at each layer when the connection probability of feed-forward connections between layers was varied. Error bars are 95% CI.
- (K) ΔQ coding persistency at each layer when the connection probability of feed-forward connections between layers was varied. Error bars are 95% CI.