

Signatures of task learning in neural representations

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Highlights

- Learning produces coordinated changes in population activity, both in neural circuits and artificial neural networks (ANNs).
- Learning reshapes neural manifolds through changes in dimensionality and the orthogonalisation of task-relevant dimensions.
- Neural population dynamics support new tasks via changes in initial conditions, dynamic modes, or the attractor landscape.
- Studies in continual learning in ANNs give insight into how neural circuits may organize network activity to support multiple skills.

Abstract

While neural plasticity has long been studied as the basis of learning, the growth of large-scale neural recording techniques provides a unique opportunity to study how learning-induced activity changes are coordinated across neurons within the same circuit. These distributed changes can be understood through an evolution of the geometry of neural manifolds and latent dynamics underlying new computations. In parallel, studies of multi-task and continual learning in artificial neural networks hint at a tradeoff between non-interference and compositionality as guiding principles to understand how neural circuits flexibly support multiple behaviors. In this review, we highlight recent findings from both biological and artificial circuits that together form a new framework for understanding task learning at the population level.

Introduction

Learning to perform new tasks is fundamental to flexible and adaptive behavior in a changing environment. Neural correlates of learning have been observed as modifications of neural firing properties, including increases in single-neuron stimulus discriminability and the recruitment of new neurons encoding task-relevant information [1, 2, 3, 4, 5]. This view of learning mirrors classic principles of neural computation based on single neuron tuning properties. Yet the past decade of systems neuroscience has experienced a pivot towards more abstract population-level dynamics [6, 7], whose structure often reflects stimulus, task, and behavioral information, and which may underlie computations in neural circuits [8]. In this review, we present recent experimental evidence of how population representations and dynamics evolve over learning as well as guiding theoretical principles from task-trained artificial neural networks (ANNs). We focus on three emerging themes: 1) learning as changes to the geometric structure of neural manifolds, 2) learning as changes in the

dynamics governing neural population trajectories, and 3) learning multiple tasks, especially while avoiding catastrophic forgetting. Together these studies form the basis for a new conceptual framework which may provide a unifying picture of how task learning is expressed in population-level representations while admitting substantial degeneracy at the synaptic and neuronal levels.

From pairwise correlations to low-dimensional manifolds

A major theme in the study of neural populations is how interneuronal correlations shape population codes. Decades of theoretical work has shown that the information encoded in neural populations is limited when noise correlations (i.e., trial-to-trial covariability for a fixed stimulus) are aligned with signal correlations (i.e., correlations due to shared tuning properties) [9]. A natural question is therefore whether improvements in task performance over learning are caused by a change in correlation structure that increases the available stimulus information. In animals trained on associative learning tasks, an overall suppression of noise correlations for high signal correlation neuron pairs has indeed been observed for rewarded stimuli [2, 10]. Perceptual learning studies have reported a reduction of average noise correlations but the link to task performance remains unclear - either because the changes were non-specific [11] or because the subjects' choices were shown to be sub-optimal with respect to the population activity structure [12]. Importantly, recent work has suggested that, rather than being stationary properties of population codes, noise correlations can change dynamically due to top-down feedback [13], and that knowledge of this modulation can be useful under certain decoding models [14].

Together, these mixed results point to the need to consider the global structure of variability in neural activity space, rather than average correlations between pairs of neurons [15]. Towards this end, a newer framework considers the low-dimensional structures in which population representations are embedded, often called neural “manifolds” (or when linear, also called “subspaces”). Neural manifold structure has been shown to constrain the speed of learning [16] and the manifold geometry underlying learned behaviors can remain stable over long timescales, even years [17]. While recent work using brain computer interfaces (BCIs) has argued that learning is constrained by pre-existing covariability structure [18, 19], one might expect the manifold structure to be flexible over long-timescale learning of new skills due to synaptic plasticity. Indeed, a follow up BCI study showed that animals were able to produce new neural patterns over several days of training (**Figure 1A**) [20]. In the following two sections, we focus on recent evidence from theoretical and experimental communities of how the geometry of neural manifolds could change to enable new task-specific computations.

Expansion and compression of dimensionality

A starting point to characterize changing manifold structure is to consider how the dimensionality of neural representations changes over learning. Both dimensionality expansion and compression have been reported at different computational stages, including at different layers of deep networks [21, 22, 23] and in recordings from different regions along the cortical hierarchy [24, 25]. This suggests that the expected relationship between changes in dimensionality and task performance depends on the chosen brain region. Experimental recordings in the motor cortex and associated regions have shown that motor skill learning is often accompanied by a reduction in the dimensionality of the task-encoding activity subspace as well as in the variability of the embedded neural trajectories [26, 27]. While such low-dimensional representations may simply reflect task simplicity and reduced behavioral variability [28], there may also be normative pressures that favor the emergence of more compact representations. Intuitively, low-dimensional task representations allow for robust decoding in the presence of independent neuronal “noise”, and can enable non-interfering representations of different skills to be stored in orthogonal low-dimensional subspaces.

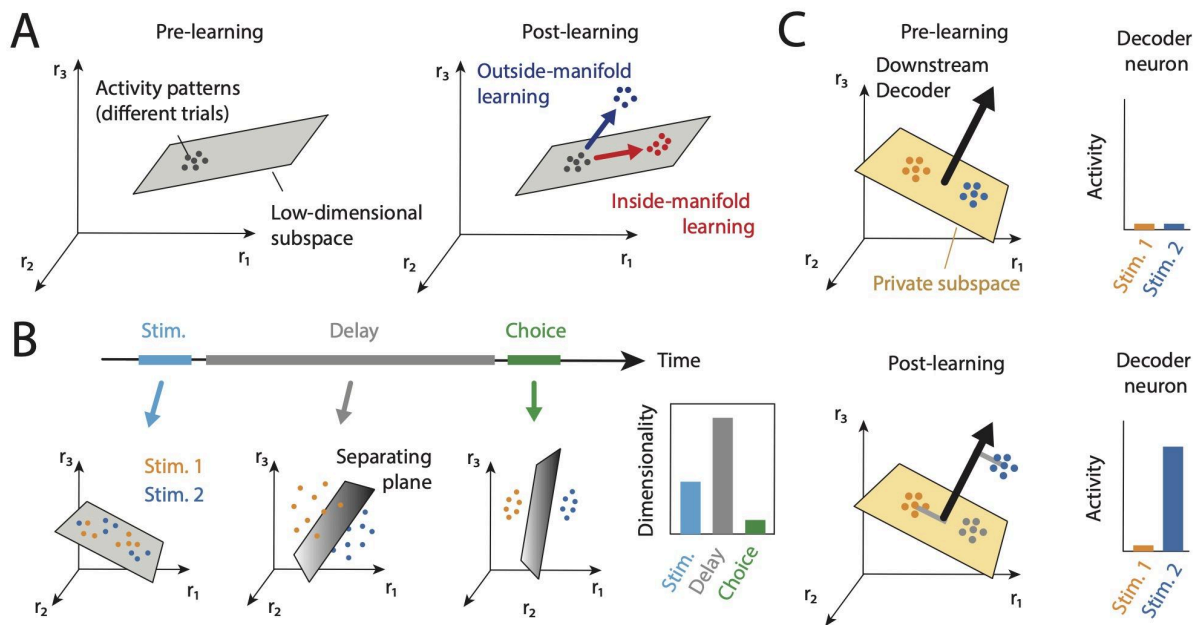


Figure 1: Learning-induced changes in manifold geometry. (A) Left: Neural activity is often observed to lie in a low dimensional subspace (i.e., linear manifold) (gray). Right: The effect of learning can be decomposed into inside-manifold learning (red) or outside-manifold learning (blue) contributions, depending on changes in activity. Both effects have been found to contribute to learning, suggesting that neural manifolds can change over long-timescale learning [20]. (B) The dimensionality of learned neural representations depends on the task period. In an ANN trained on a classification task, the dimensionality of each stimulus representation increases transiently to improve linear discriminability, but becomes more compact during the choice period to enable a robust downstream readout. In this scenario, learning could simultaneously increase delay-period dimensionality while decreasing choice-period dimensionality. Adapted from [29]. (C) Example of re-orientation of task relevant dimensions to a decoder axis. Top: (Pre-learning) Representations of two stimuli are separable but in the same decoder-null subspace that is orthogonal to the dimension representing the decoding axis (left). Both stimuli produce weak responses in the decoder neuron (right). Bottom: After training, the representation of Stim. 2 shifts along the decoding axis, producing a larger projection onto the decoding axis (left) and hence a larger downstream response (right). More generally, network activity can shift between a decoder-null or decoder-potent subspace to suppress or enhance the readout. See also [30].

While low-dimensional activity can be a signature of systems consolidation at the end of learning, high *initial* dimensionality could prove useful for probing new neural patterns in a search for optimal control [31]. In line with this idea, one study of BCI learning found an initially high contribution of “private” variability of individual motor cortical neurons to task performance, which reduced over trials as the population transitioned to using lower-dimensional “shared” variability for control [32]. By running either private or shared signals through the BCI decoder separately, Athalye and colleagues demonstrated that movements driven by the shared signal were faster and more direct, suggesting that the observed high-dimensional activity followed by compression reflected an implicit change in strategy for efficient goal-directed motor control. A compression of the dimensionality of stimulus representations was also observed in the primate prefrontal cortex in a recent classical conditioning study [33]. The authors argued that this compression could arise from the dual pressures of increased generalization and metabolic constraints [33], further showcasing the different tradeoffs inherent in dimensionality expansion and compression.

Additional intuition can be obtained from theoretical studies of learning in recurrent neural networks (RNNs), which are trained to minimize the error of a readout unit. For example, one recent study found that learning in RNNs trained on simple tasks induces weight changes that are low-rank [34], a form of connectivity structure which has previously been linked to low-dimensional neural activity [35]. More recent work has additionally found low-rank structure in the evolution of the weights over learning [36]. But similar to [32], RNNs often show benefits of being initialized with high-dimensional activity in early epochs [34, 37]. Another key study showed that RNNs trained with stochastic gradient descent tend to compress representations specifically during the “decision” period, even in tasks where transient high-dimensional activity is computationally useful (**Figure 1B**) [29]. The authors pinpointed the cause of this dimensionality compression to the stochasticity inherent to the weight updates, which acts as an effective regularizer to enforce robust representations. While these studies show that low-dimensional neural representations could emerge as a result of task structure and gradient-based optimization, it remains an open question whether similar effects on dimensionality are found with biologically plausible plasticity rules which are local and may have task-irrelevant components [38]. Towards this end, one study has shown that dimensionality can be systematically affected by circuit motifs between pairs and triplets of neurons [39], suggesting a possible pathway for the regulation of global dimensionality through local synaptic plasticity.

Reorienting task-relevant dimensions

Beyond global changes in dimensionality, learning may drive specific changes in the geometry of neural representations. Examining how task-relevant dimensions reorient themselves over trials can provide insight regarding computational strategies that develop for solving the task. In particular, recent work has shown that neural population activity is often structured into orthogonal subspaces, enabling the circuit to separate sensory-driven from spontaneous activity patterns [40], to independently encode distinct task variables [41, 42], or to arrange dynamics in separate subspaces for context-specific computations [43, 44, 45]. Orthogonalisation of neural dimensions can also occur at different periods within a task in order to store non-interfering representations of past stimuli in short-term memory. This effect was recently observed in the auditory cortex during an implicit learning paradigm with auditory sequences: the optimal decoding axis for early stimuli evolved throughout sequence progression, and was eventually orthogonal to the original stimulus encoding dimensions [46]. This allowed previous stimuli in the sequence to be encoded by the same circuit within a “memory” subspace orthogonal to the “sensory” subspace. While Libby and Buschman reported an increase in this orthogonalisation with passive exposure, the absence of behavioral readout precludes analysis of the timescale of statistical learning. Thus a more direct comparison of pre- and post-learning activity is needed to verify that such orthogonalisation is shaped by task requirements.

Rearranging activity into orthogonal subspaces can also be useful for selective information routing between brain regions. A single neuron can be viewed as a linear decoder, which projects upstream neural patterns onto a single dimension; any changes in upstream activity orthogonal to this dimension have no effect on its firing (**Figure 1C**). Similarly, a downstream population can be viewed as a linear subspace determined by the decoder axes of all its neurons. Neural circuits could thus regulate downstream responses by re-orienting activity from a “decoder-null” to a “decoder-potent” subspace to communicate the outcome of an internal computation to other brain regions [30, 43, 44, 47]. For example, learning of a visuomotor association task specifically orthogonalised responses for motor-associated stimuli in V1 [48]. Another study found that in animals trained on auditory Go/No-Go tasks, auditory cortical representations were reoriented during task engagement so that an inferred downstream stimulus decoder would observe enhanced activation during a target stimulus (which required a behavioral change), whereas reference stimuli would be indistinguishable from baseline activity [49]. In both studies, the orthogonalisation of task-relevant dimensions was not necessarily for an increase in stimulus discriminability. Rather, this selective and contextual re-orientation may

correspond to an alignment with a decoder-potent subspace that could be used to trigger stronger downstream responses for behaviorally relevant stimuli (**Figure 1C**).

So far, we have focused on learning-induced changes to the geometry of neural manifolds or of decoder axes. An alternative perspective instead focuses on the low-dimensional subspaces that may control how neural trajectories unfold as a dynamical system [50], rather than low-dimensional subspaces in which neural activity is embedded. For example, rather than orthogonalization of dimensions encoding stimuli, a re-alignment of the dimensions dictating the flow of neural dynamics can be used to shape how upstream inputs are integrated by neural populations. Towards this end, [51] analyzed mouse V1 responses during a visual discrimination task, and inferred recurrent interactions and feedforward input by fitting autoregressive models to the data. They observed that task learning led to an increase in stimulus information in V1 without a concurrent increase in the inferred external input. Rather, improvements in V1 were better explained by enhanced temporal integration of task-relevant input after learning, and was achieved in the network by realigning the slowly decaying modes of recurrent dynamics with the most informative direction of network input (**Figure 2A**). This finding underscores the importance of not only considering learning-induced changes in manifold geometry, but also in the underlying dynamical processes that sculpt neural representations.

Reshaping the dynamics underlying task performance

Besides changes to the geometry of neural representations (e.g., realignment of stimulus-decoding dimensions; **Figure 1C**), learning can require changes to the dynamical structure that govern how population activity evolves under different conditions (e.g., realignment of slow and fast modes; **Figure 2A**). Activity-based methods focus on the structure of population activity in neural activity space, often with the implicit assumption that neural data points are drawn from some stationary distribution with no notion of time (e.g., PCA). On the other hand, dynamics-based methods consider the forces that determine how population activity changes from one time point to the next. This latter view takes insight from dynamical systems theory to ask how recurrent connectivity and neuron-intrinsic nonlinearities combine to shape the time-varying neural trajectories which implement population-level computations. These two views are not exclusive and can even be combined for a fuller view of task learning, as is demonstrated in recent work on motor learning: in the motor cortex, preparatory activity is often constrained to an “output-null” subspace orthogonal to movement-related activity [47], which is hypothesized to separate preparation from the execution of motor plans. However, orthogonal representations do not necessarily mean that the dynamics do not interact, as activity within the preparatory subspace can be causally important for the quality of succeeding movement kinematics [52, 53]. Refinement of motor skills may thus involve targeting preparatory activity to specific subspaces and setting appropriate “initial conditions” to trigger the relevant network dynamics during the movement phase (**Figure 2B**). This hypothesis was tested by Perich and colleagues [54] who simultaneously recorded activity in primate dorsal premotor (PMd) and primary motor (M1) cortices during motor adaptation. By fitting predictive models of M1 neuron spiking using PMd and M1 population activity, the authors inferred that learning-induced changes in M1 activity were driven by upstream changes in planning-related activity in PMd. A more recent study observed structured rotations in preparatory activity states during curl force-field adaptation, which were specific to the direction of the curl field and the subset of targets showing behavioral signatures of adaptation [55]. These findings are consistent with a view that adapted neural dynamics in M1 can be generated by altered initial conditions without a change to the intrinsic M1 dynamical repertoire (although further experiments are needed to examine the potential contribution of small, correlated synaptic changes within M1 [56]).

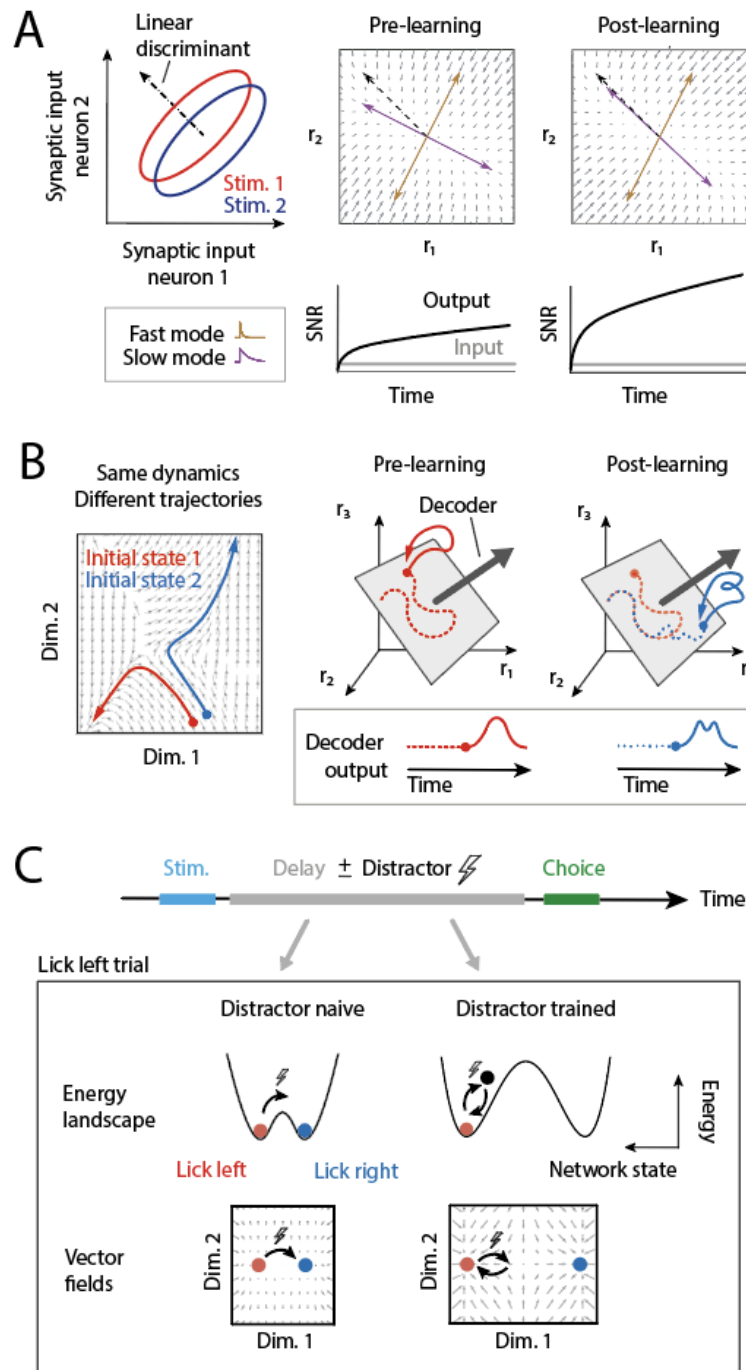


Figure 2: Learning-induced changes in dynamics. **(A)** Alignment of slow network dynamics with informative input modes. The network learns to discriminate between two stimuli by performing leaky integration of noisy feedforward input. Left: Distributions of synaptic input onto two neurons for two stimuli at any given time. The linear discriminant axis separates the two representations (dashed black) and is the most informative input direction. Middle: Recurrent dynamics illustrated by a vector field diagram. The gray arrows indicate the (instantaneous) velocity of the system state (here, neural population activity). In this example, recurrent dynamics are decomposed into a slowly-decaying (purple) and a fast-decaying (orange) mode (top). Stimulus information (or SNR; signal-to-noise) in network output grows as compared to input. Right: Over training, realignment of the slow mode to the input linear discriminant axis means stimulus information decays less quickly during integration, leading to more rapid evidence accumulation. Adapted from [51]. **(B)** Changes in initial conditions can trigger different neural trajectories. Left: Example vector

field visualized along two dimensions of neural activity space. By changing the initial network activity state (red and blue dots), the same underlying dynamical structure can be used to produce different neural trajectories, which may have different consequences for downstream decoding. Right: Motor planning-related activity (dashed red) is orthogonal to the movement-decoding axis and can therefore set the initial condition (red circle) without producing any movement. Upon receiving a 'Go-cue', the network activity evolves as the trajectory in red and produces motor output. After training, the network follows a different preparatory trajectory (dashed blue) leading to a new initial condition (blue circle) at the end of motor planning. This triggers a new trajectory (blue) that changes the motor output. (C) Changes in attractor dynamics during a stimulus detection task. Top: Mice were trained to report the presence (lick right) or absence (lick left) of a stimulus; during the cue, a distractor stimulus (lightning bolt) was sometimes added as a perturbation. Bottom: Schematic of the attractor landscape and corresponding vector field in distractor-naïve (left) and distractor-trained (right) mice. During the delay period, network activity is in one of the attractor basins, maintaining a motor plan. In distractor-naïve mice, distractors are able to switch the network activity from one attractor to another. In distractor-trained mice, the network state is more robust due to greater separation of attractor basins. Adapted from [57].

In addition to setting new initial conditions for dynamics [54] or realignment of slow dynamical modes [51], learning could shape the internally generated attractor landscape. This was recently demonstrated in the anterolateral motor cortex (ALM) during a stimulus detection task [57]. Mice were trained to report the presence or absence of optogenetic stimulation in vibrissal somatosensory cortex (vS1) by licking left or right after a delay period, and to ignore further vS1 “distractor” stimulation, which occurred during the delay period. In distractor-trained mice, ALM network activity quickly recovered back to the correct choice-related activity. By contrast, in distractor-naïve animals, even weak distractors led to more persistent perturbations of ALM activity, with frequent switches to the incorrect motor plan. Using task-trained ANNs, the authors inferred that robustness against distractors developed due to increased separation and stability of the two choice-related attractors, particularly in the late-delay period (**Figure 2C**). Together, these studies highlight how learning can shape task-relevant dynamics via rearrangement of input and recurrent interactions.

Learning multiple tasks via non-interference and compositionality

Beyond learning of individual tasks, a growing body of work asks how neural circuits can flexibly support multiple simultaneously learned tasks. Expanding from the ideas above, one effective strategy to minimize interference between tasks (or “contexts”) is to use orthogonal task representations. This can take the form of separating task features along different dimensions for selective integration [41]; compression of stimulus representations along task-irrelevant dimensions [58]; or at its extreme, the use of non-overlapping neural populations in different contexts [59, 60]. Such a strategy is most useful when those tasks don’t share common computations and/or require knowledge of independent stimulus features. But as organisms acquire a vast, flexible repertoire of overlapping skills over their lifetimes, it becomes more efficient to instead decompose tasks into modular operations that can be recombined in many ways [61]. Such compositionality has been observed in ANNs trained simultaneously on 15–20 tasks, both in the algebraic relationships of different task representations [62] as well as regarding the reuse of dynamical structure for similar tasks [63, 64]. While these results were observed without architectural constraints, a bias for compositional representations could be imposed through structural bottlenecks and gating inputs to force shared representations [65].

A related perspective emerges in recent work that focuses on sequential (as opposed to simultaneous) learning: how does new task learning interfere with previously acquired skills? In particular, ANNs are notoriously susceptible to “catastrophic forgetting” of prior tasks when new tasks are learned. Multiple solutions to

catastrophic forgetting have recently been proposed under the umbrella term of continual learning [66, 67], including penalizing weight changes that would incur a loss in performance on previous tasks [68], orthogonalizing weight updates to encourage non-interfering representations (**Figure 3A**) [69, 70, 71, 72] and contextual gating induced by Hebbian learning rules [73]. While these theory-driven hypotheses remain challenging to validate in neural data, one recent study observed an interesting parallel when monkeys were sequentially trained on two BCI mappings A and B [74]. After learning the second map B, neural activity during re-exposure to map A had moved along the “null space” of task A (i.e., orthogonal to map A) and didn’t impede their performance. However these shifted neural activity patterns could also produce improved control signals via map B and could serve as a “memory trace” for B (**Figure 3B**). More broadly, sequential learning studies in primates have observed that information about previous training is reflected in the geometry of task-specific representations [55, 75], echoing findings that the emergent solutions found by ANNs generally depend on pre-training (and can even reorganize representations corresponding to previously learnt tasks without a drop in performance) [63, 76]. However, a key difference between ANNs and biological brains is the high degree of (hierarchical) modularity found in the latter, with disparate learning rules identified across regions, suggesting that a distinct strategy may be at play to enable learning throughout an organism’s lifetime. More work needs to be done, both in artificial and biological neural networks, to fully understand the tradeoffs between non-interference and compositionality as guiding principles for learning multiple tasks.

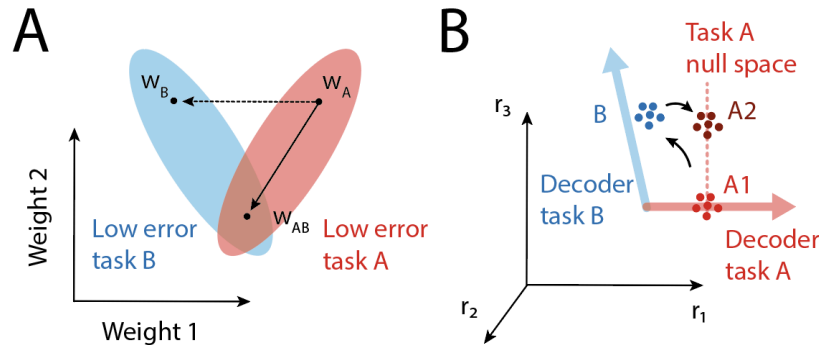


Figure 3: Sequential skill acquisition depends on previously acquired skills. (A) Continual learning algorithms provide a solution to catastrophic forgetting by constraining weight changes so that they do not lead to poor performance in previously-learned tasks. In this example, an ANN that has been pre-trained on task A (initial weights w_A) lies in the red oval corresponding to solutions with low task A error. When trained on task B, standard gradient descent leads to weights w_B , which have low error on task B (blue oval) but poor performance on task A (“forgetting”; dotted line). Instead, continual learning updates the weights so that the resulting network can perform both tasks (w_{AB}). Figure adapted from [68] and [72]. **(B)** Schematic of learning in subjects trained sequentially on two BCI tasks A and B (determined by red and blue decoder axes). Over learning, neural representations are updated from having high projection on decoder A to having high projection on decoder B. Interestingly, when subjects are then re-exposed to task A (A2; maroon), the representations lie in a region corresponding to high performance in both tasks, revealing a ‘memory trace’ of map B. Adapted from [74].

Outlook and challenges ahead

Viewing task learning as shaping of population dynamics to support new computations provides a powerful framework to interpret the diverse learning-induced changes that have been observed in neural data. While we mainly focused on neocortical circuits in this perspective article, these principles extend to learning-related reorganization in other brain regions such as hippocampal [77], cerebellar [26, 78] and limbic [79] circuits, and need to be bridged with decades-long insights on synaptic and intrinsic plasticity mechanisms that enable learning [80, 81]. To forge ahead, several experimental and theoretical challenges need to be overcome. Longitudinal monitoring of the same neural population throughout learning is critical for teasing apart refinement of existing motifs from emergence of new features of population activity [82-84]. Data-analysis tools such as alignment [17, 85, 86] or identification of latent structure across sessions [36, 87, 88], and disentangling input from recurrent contributions [89, 90] will be crucial to track distributed changes in neural representations over learning. Reporting training history and inferring the implicit strategies of animals will help link these neural changes to the behaviors they support, and account for inter-individual variability [75, 91, 92]. Lastly, leveraging further insights from learning in ANNs requires a better understanding of how different (or similar) gradient-based optimization is to biological learning [19, 93] as well as an examination of the dependence of learnt task representations on specific learning rules [94, 95]. Together, these advances will provide a better understanding of the mechanisms underlying behavioral flexibility, guiding future work on lifelong learning in biological and artificial agents.

Declaration of Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Data availability

No data was used for the research described in the article.

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