HOW CONNECTIVITY STRUCTURE SHAPES RICH AND LAZY LEARNING IN NEURAL CIRCUITS

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ABSTRACT

In theoretical neuroscience, recent work leverages deep learning tools to explore how some network attributes critically influence its learning dynamics. Notably, initial weight distributions with small (resp. large) variance may yield a rich (resp. lazy) regime, where significant (resp. minor) changes to network states and representation are observed over the course of learning. However, in biology, neural circuit connectivity generally has a low-rank structure and therefore differs markedly from the random initializations generally used for these studies. As such, here we investigate how the structure of the initial weights — in particular their effective rank — influences the network learning regime. Through both empirical and theoretical analyses, we discover that high-rank initializations typically yield smaller network changes indicative of lazier learning, a finding we also confirm with experimentally-driven initial connectivity in recurrent neural networks. Conversely, low-rank initialization biases learning towards richer learning. Importantly, however, as an exception to this rule, we find lazier learning can still occur with a low-rank initialization that aligns with task and data statistics. Our research highlights the pivotal role of initial weight structures in shaping learning regimes, with implications for metabolic costs of plasticity and risks of catastrophic forgetting.

1 Introduction

In theoretical neuroscience, it has been shown that structural variations can have a pronounced impact on learning dynamics in humans and animals. For instance, studies have revealed that specific neural connectivity patterns can facilitate faster learning of certain tasks (Braun et al., 2022; Raman & O'Leary, 2021; Simard et al., 2005; Xie et al., 2022; Duncker et al., 2020; Goudar et al., 2023; Chang et al., 2023). Similarly, in deep learning, the structure (in the form of architecture or initial connectivity) is a pivotal design component, and it dictates learning speed and quality alongside the data, the objective function, and learning rule (Richards et al., 2019; Zador, 2019; Yang & Molano-Mazón, 2021; Braun et al., 2022).

One particularly notable aspect of the structure is the initial connectivity before training takes place. Specifically, the initial connection weight magnitude can significantly bias learning dynamics, pushing them towards either rich or lazy regimes (Chizat et al., 2019; Flesch et al., 2021). Lazy learning often induces minor changes in the network during the learning process. Such minimal adjustments are advantageous given that plasticity is metabolically costly (Mery & Kawecki, 2005; Plaçais & Preat, 2013), and significant changes in representations might lead to issues like catastrophic forgetting (McCloskey & Cohen, 1989; Kirkpatrick et al., 2017). On the other hand, the rich learning regime can significantly adapt the network's internal representations to task statistics, which can be advantageous

for task feature acquisition and has implications for generalization (Flesch et al., 2021; George et al., 2022). While influential works have highlighted the role of initial weight magnitude in determining learning dynamics (Woodworth et al., 2020; Flesch et al., 2021; Braun et al., 2022), most studies have focused on tabula rasa initializations like isotropic Gaussian/Uniform patterns. These patterns stand in contrast to the connectivity structures observed in biological neural circuits, which frequently exhibit a more pronounced low-rank eigenstructure (Song et al., 2005; Mastrogiuseppe & Ostojic, 2018). This divergence prompts a pivotal question: how does the initial weight structure, given a fixed initial weight magnitude, bias the learning regime?

The goal of this study is to understand how initial weight structure, particularly the effective rank, modulates the *effective* richness or laziness of task learning within the standard training regime. We note that *rich* and *lazy* learning regimes have well established meanings in deep learning theory. The latter being defined as a situation where the Neural Tangent Kernel (NTK) stays stationary during training, while the former refers to the case where the NTK changes. In this work, we slightly extend these definitions and introduce **effective learning richness/laziness**. Unlike the traditional definition, which is based upon initialization, effective learning richness/laziness is defined in terms of post-training adjustment measurements. From this perspective, a learning process is deemed effectively "lazy" if the measured NTK movement is small. For example, consider a network whose initialization puts it in standard rich regime, but for a given task, its NTK moves very little during training. We define learning for this specific situation as effectively lazy. In other words, while the standard regime definition informs us (prior to training) whether the network can adapt significantly to task training or not, our "effective" definition lies in the post-training effects.

1.1 CONTRIBUTIONS

Our main **contributions** and findings can be summarized as follows:

- Through theoretical derivation in two-layer feedforward linear network, we demonstrate that higher-rank initialization results in *effectively* lazier learning **on average** across tasks (Theorem 1). We note that the emphasis of the theorem is on the expectation across tasks.
- We validate our theoretical findings in recurrent neural networks (RNNs) through numerical experiments on well-known neuroscience tasks (Figure 1) and demonstrate the applicability to different initial connectivity structures extracted from neuroscience data (Figures 2 3).
- We identify scenarios where certain low-rank initial weights still result in *effectively* lazier learning for specific tasks (Proposition 1 and Figure 4). We postulate that such patterns emerge when a neural circuit is predisposed perhaps due to evolutionary factors or post-development to certain tasks, ingraining specific inductive biases in neural circuits.

1.2 RELATED WORKS

An extended discussion on related works can also be found in Appendix A.

Theoretical Foundations of Neural Network Regimes and Implications for Neural Circuits: The deep learning community has made tremendous strides in developing theoretical groundings for artificial neural networks (Advani et al., 2020; Jacot et al., 2018; Alemohammad et al., 2020; Agarwala et al., 2022; Atanasov et al., 2021; Azulay et al., 2021; Emami et al., 2021). A focal point is the 'rich' and 'lazy' learning regimes dichotomy, which have distinct impacts on representation and generalization (Chizat et al., 2019; Flesch et al., 2021; Geiger et al., 2020; George et al., 2022; Ghorbani et al., 2020; Woodworth et al., 2020; Paccolat et al., 2021; Nacson et al., 2022; HaoChen et al., 2021; Flesch et al., 2023). The 'lazy' regime results in minimal weight changes, while the 'rich' regime fosters task-specific adaptations. The transition between these is influenced by various factors, including initial weight scale and network width (Chizat et al., 2019; Geiger et al., 2020).

A nexus between deep learning and neuroscience has expanded the applications of deep learning theoretical frameworks to study the learning dynamics of biological neural networks (Bordelon & Pehlevan, 2022; Liu et al., 2022a; Braun et al., 2022; Ghosh et al., 2023; Saxe et al., 2019; Farrell et al., 2022; Papyan et al., 2020; Tishby & Zaslavsky, 2015). For the rich/lazy regime theory, the existence of diverse learning regimes in neural systems is evident through the resource-intensive plasticity-driven transformations prevalent in developmental phases, followed by more subdued

adjustments (Lohmann & Kessels, 2014), and previous investigations characterized neural network behaviors under distinct regimes (Bordelon & Pehlevan, 2022; Schuessler et al., 2023) and discerning which mode yields solutions mimicking neural data (Flesch et al., 2021). Our work builds on these studies by further exploring the precursors of these regimes, focusing on how the brain's initial weight configurations influence task learning.

Neural circuit initialization, connectivity patterns and learning: Extensive research has explored the influence of various random initializations on deep network learning (Saxe et al., 2013; Bahri et al., 2020; Glorot & Bengio, 2010; He et al., 2015; Arora et al., 2019). The literature predominantly focuses on random initialization, but actual neural structures exhibit markedly different connectivity patterns, such as Dale's law and enriched cell-type-specific connectivity motifs (Rajan & Abbott, 2006; Ipsen & Peterson, 2020; Harris et al., 2022; Dahmen et al., 2020; Aljadeff et al., 2015). Given the low-rankedness in neural circuits (Song et al., 2005; Mastrogiuseppe & Ostojic, 2018), our study investigates the impact of effective rank on learning regimes. With emerging connectivity data (Campagnola et al., 2022; Consortium et al., 2021; Dorkenwald et al., 2022; Winnubst et al., 2019; Scheffer et al., 2020), future work is poised to encompass rich additional features of connectivity.

2 SETUP AND THEORETICAL FINDINGS

2.1 RNN SETUP

We examine recurrent neural networks (RNNs) because they are commonly adopted for modeling neural circuits (Barak, 2017; Song et al., 2016). We consider a RNN with N_{in} input units, N hidden units and N_{out} readout units (Figure 1A). The update formula for $h_t \in \mathbb{R}^N$ (the hidden state at time t) is governed by (Ehrlich et al., 2021; Molano-Mazon et al., 2022):

$$h_{t+1} = \rho h_t + (1 - \rho)(W_h f(h_t) + W_x x_t), \tag{1}$$

where $\rho=e^{-dt/\tau_m}\in\mathbb{R}$ denotes the leak factor for simulation time step dt and τ_m denotes the membrane time constant; $f(\cdot):\mathbb{R}^N\to\mathbb{R}^N$ is the activation function, for which we use ReLU; $W_h\in\mathbb{R}^{N\times N}$ (resp. $W_x\in\mathbb{R}^{N\times N_{in}}$) is the recurrent (resp. input) weight matrix and $x_t\in\mathbb{R}^{N_{in}}$ is the input at time step t. Readout $\hat{y}_t\in\mathbb{R}^{N_{out}}$, with readout weights $w\in\mathbb{R}^{N_{out}\times N}$, is defined as

$$\hat{y}_t = \langle w, f(h_t) \rangle. \tag{2}$$

The objective is to minimize scalar loss $L \in \mathbb{R}$, for which we use the cross-entropy loss for classification tasks and mean squared error for regression tasks. L is minimized by updating the parameters using variants of gradient descent:

$$\Delta W = -\eta \nabla_W L,\tag{3}$$

for learning rate $\eta \in \mathbb{R}$ and $W = [W_h \quad W_x \quad w^T] \in \mathbb{R}^{N \times (N_{in} + N + N_{out})}$ contains all the trainable parameters. Details of parameter settings can be found in Appendix C.

2.2 EFFECTIVE LAZINESS MEASURES

As mentioned above, we introduce *effective* richness and laziness, with effectively lazier (resp. richer) learning corresponding to less (resp. greater) network change over the course of learning. To quantify network change, we adopt the following three measures that have been used previously (George et al., 2022). We note that these measures can be sensitive to other architectural aspects that bias learning regimes, such as network width, so throughout we hold these variables constant when making the comparisons.

Weight change norm quantifies the vector norm of change in W. Effectively lazier learning should result in a lower weight change norm, and it is quantified as:

$$\|\Delta W\| := \|W^{(f)} - W^{(0)}\|,\tag{4}$$

where $\|\cdot\| = \|\cdot\|_F$; $W^{(0)}$ (resp. $W^{(f)}$) are the weights before (resp. after) training.

Representation alignment (RA) quantifies the directional change in a representational similarity matrix (RSM) before and after training. RSM focuses on the similarity between how two pairs

of input are represented by computing the Gram matrix R of last step hidden activity. Greater representation alignment indicates effectively lazier learning in the network, and it is obtained by

$$RA(R^{(f)}, R^{(0)}) := \frac{Tr(R^{(f)}R^{(0)})}{\|R^{(f)}\|\|R^{(0)}\|}, \text{ where } R := H^TH,$$
 (5)

where $H \in \mathbb{R}^{N \times m}$ is the hidden activity at the last time step; $R^{(0)}$ and $R^{(f)} \in \mathbb{R}^{m \times m}$ are the initial and final RSM, respectively; m is the batch size.

Tangent kernel alignment (KA) quantifies the directional change in the neural tangent kernel (NTK) before and after training; effectively lazier learning should result in higher tangent kernel alignment. The NTK computes the Gram matrix K of the output gradient. Greater tangent kernel alignment points to effectively lazier learning, and it is obtained by

$$KA(K^{(f)}, K^{(0)}) := \frac{Tr(K^{(f)}K^{(0)})}{\|K^{(f)}\|\|K^{(0)}\|}, \quad \text{where } K := \nabla_W \hat{y}^T \nabla_W \hat{y}$$
 (6)

where $K^{(0)}$ and $K^{(f)} \in \mathbb{R}^{m \times m}$ (for the $N_{out} = 1$ case) denote the initial and final NTK, respectively.

2.3 Theoretical findings

This subsection derives the theoretical impact of initial weight effective rank on tangent kernel alignment. First, Theorem 1 focuses on **task-agnostic** settings, treating task definition as random variables and computing the **expected** tangent kernel alignment across tasks. With some assumptions, tangent kernel alignment is maximized when the initial weight singular values are distributed across all dimensions (i.e. high-rank initialization).

In this section, our theoretical results are framed in a simplified feedforward setting, as we use a two-layer network with linear activations. However, we return to RNNs (Eq. 1) for the rest of the paper, and verify the generality of our theoretical findings with numerical experiments for both feedforward and recurrent architectures. Our choice is motivated by the need for theoretical tractability. While research on RNN learning in the NTK regime exists (Yang, 2020; Alemohammad et al., 2020; Emami et al., 2021), we are not aware of any studies featuring the final converged NTK that could serve as a basis for our comparison of the initial and final kernel. Consequently, we have chosen to focus on RNNs for neural circuit modeling and employ linear feedforward networks for theoretical derivations, a strategy also adopted by Farrell et al. (2022); numerous other studies, including Saxe et al. (2019), (Atanasov et al., 2021), (Arora et al., 2019), and (Braun et al., 2022), have similarly concentrated on extracting theoretical insights from linear feedforward networks.

For a two-layer linear network with input data $X \in \mathbb{R}^{d \times m}$, $W_1 \in \mathbb{R}^{N \times d}$ and $W_2 \in \mathbb{R}^{1 \times N}$ as weights for layers 1 and 2, respectively, the NTK throughout training, K, is:

$$K = X^{T}(W_{1}^{T}W_{1} + ||W_{2}||^{2}I)X.$$
(7)

Without the loss of generality, suppose the output target $Y \in \mathbb{R}^{1 \times m}$ is generated from a linear teacher network as $Y = \beta^T X$, for some Gaussian vector $\beta \in \mathbb{R}^d$, with $\beta_i \sim \mathcal{N}(0, 1/d)$.

Theorem 1. (Informal) Consider the network above with its corresponding NTK in Eq. 7, trained under MSE loss with small initialization and whitened data. The expected kernel alignment across tasks is maximized with high-rank initialization, i.e. the singular values of $W_1^{(0)}$ are distributed across all dimensions. (Formal statement and proof are in Appendix B)

We emphasize again that Theorem 1 is **task-agnostic**, i.e. it focuses on the **expected** tangent kernel alignment across input-output definitions. This is in contrast to **task-specific** settings (e.g. Woodworth et al. (2020)) that focus on a given task. In such task-specific settings, certain low-rank initializations can in fact lead to lazier learning. The following proposition predicts that if the task structure is known, low-rank initialization that is already aligned with the task statistics (input/output covariance) can lead to kernel alignment. We revisit this proposition again in Figure 4. We remark that initializing this way can still have high initial error because of randomized $W_2^{(0)}$.

Proposition 1. (Informal) Following the setup and assumptions in Theorem 1, rank-1 initializations of the form $W_1^{(0)} = \sigma[\beta^T/\|\beta\| \ \vec{0} \ ... \ \vec{0}]$ leads to a high tangent kernel alignment. (Formal statement and proof are in Appendix B)

Above, we state technical results in terms of one metric of the effective laziness of learning, based on the NTK; our proof in Appendix B easily extends also to the representation alignment metric. The impact on weight change is also assessed in Appendix Proposition 2. This is in line with our simulations with RNNs, which will show similar trends for all three of the metrics introduced in Section 2.2).

3 SIMULATION RESULTS

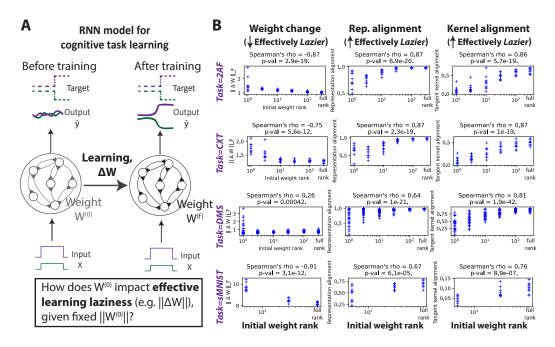


Figure 1: Low-rank initial recurrent weights, generated using SVD, lead to greater changes (or effectively richer learning) in the recurrent neural network. A) Schematic of RNN training setup. B) Measurements of effective richness vs laziness of learning (metrics as defined in Section 2.2), for RNN trained on several cognitive tasks in Neurogym (Molano-Mazon et al., 2022) as well as the sequential MNIST task (sMNIST). For details on SVD weight creation, see Appendix C. Fewer rank points were used for sMNIST due to computational time. Each dot represents a single training run, with each run using a different random initialization (10 runs total for each setting).

In this section we empirically illustrate and verify our main theoretical results, which are: (1) on average, high-rank initialization leads to effectively lazier learning (Theorem 1); (2) it is still possible for certain low-rank initializations that are already aligned to the task statistics to achieve effectively lazier learning (Proposition 1).

Impact on effective laziness by low-rank initialization via SVD in RNNs: As a proof-of-concept, we start in Figure 1 with low-rank initialization in RNNs by truncating an initial Gaussian random matrix via Singular Value Decomposition (SVD), which enables us to precisely control the rank, and rescale it to ensure that the comparison is across the same weight norm (or equivalently, variance, because of zero mean initialization) due to the role of the initial weight magnitude in influencing the learning regime (Schuessler et al., 2020). Additionally, all comparisons were made after training was completed, and all these training sessions achieved comparable losses. For our investigations, we applied this initialization scheme across a variety of cognitive tasks — including two-alternative forced choice (2AF), delayed-match-to-sample (DMS), context-dependent decision-making (CXT) tasks — implemented with Neurogym (Molano-Mazon et al., 2022) and the well-known machine learning benchmark sequential MNIST (sMNIST). Details of our simulation setup can be found in Appendix C. For statistical analysis and significance tests, we used methods in the SciPy Package (Virtanen et al., 2020).

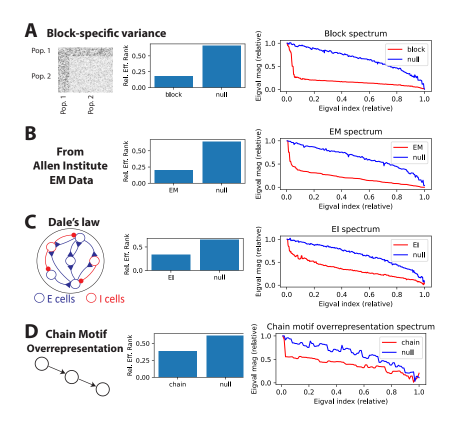


Figure 2: Low-rank weight structure through example biologically inspired structures. We present the eigenspectrum and (relative) effective rank of connectivity A) with cell-type-specific statistics, B) derived from EM data, C) obeying Dale's law, and D) with an over-representation of chain motif. These structures result in a lower effective rank compared to the standard random Gaussian initialization (null) with the same magnitude. We plotted the magnitude of the eigenvalues (Eigval mag) scaled by the dominant eigenvalue magnitude, against their indices divided by the network size N (Eigval index). Please see Appendix C for details on the network initialization.

Figure 1 indicates that low-rank initial weights result in effectively richer learning and greater network changes. These numerical trends are in line with Theorem 1, which is also demonstrated for a two-layer linear network in Appendix Figure 10. In the Appendix, we show the main trends observed in Figure 1 also hold for Uniform initialization (Figure 5), soft initial weight rank (Figure 6), various network sizes (Figure 7), learning rates (Figure 8), and gains (Figure 9).

Low-rank initialization via biologically motivated connectivity in RNNs: To establish a closer connection with biological neural circuits, we have tested our predictions on low-rank initialization using a variety of biologically motivated structures capable of resulting in low-rank connectivity. Here are some of the examples: (A) connectivity with cell-type-specific statistics (Aljadeff et al., 2015), where each block in the weight matrix corresponds to the connections between neurons of two distinct cell types, with the variance of these connections differing from one block to another. In terms of block-specific connectivity statistics, there are infinite possibilities for defining the blocks, each resulting in a unique eigenspectrum. For the example provided here, we adopted the setup from Figure S3 in Aljadeff et al. (2015), with parameters set as $\alpha = 0.02$, $\gamma = 10$, and $1 - \epsilon = 0.8$; these correspond to the fraction of hyperexcitable neurons, gain of hyperexcitable connections and gain of the rest, respectively. We follow this particular setup because it has been demonstrated to create an outlier leading eigenvalue, thereby reducing the effective rank. We also consider (B) connectivity matrix derived from the electron microscopy (EM) data (Institute, 2023), where the synaptic connections between individual neurons are meticulously mapped to create a detailed and comprehensive representation of neural circuits. This high-resolution data enables the analysis of intricate structural patterns and the elucidation of the diverse connectivity properties characteristic

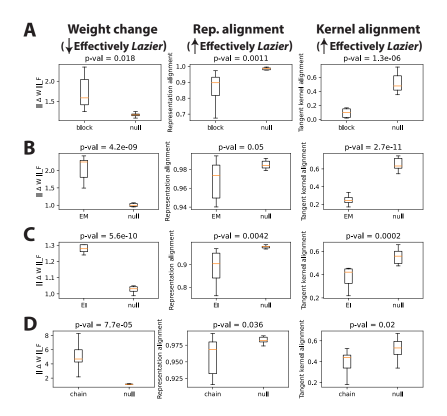


Figure 3: Low-rank initialization, achieved by experimentally-driven connectivity in Figure 2, leads to effectively richer learning. We apply the effective laziness measures in Section 2.2 to compare the effective laziness of the experimentally-driven initial connectivity versus standard random Gaussian initialization (null). Panels A-D corresponds to that of Figure 2. The boxplots are generated from 10 independent runs with different initialization seeds. Again, comparisons are done across equal initial weight magnitude. Details of how the simulations are conducted can be found in Appendix C. Due to the space limit, we include only the 2AF task here and show in Appendix Figures 11 and 12 that similar trends hold for the DMS and CXT tasks.

of biological neural networks. Also, we consider (C) connectivity obeying Dale's law, where each neuron is either excitatory or inhibitory, meaning it can only send out one type of signal – either increasing or decreasing the activity of connected neurons – a principle inspired by the way neurons behave in biological systems (Song et al., 2005). Additionally, (D) the over-representation of certain localized connectivity patterns (or network motifs), e.g. chain motif (Zhao et al., 2011; Hu et al., 2018), in which two cells are connected via a third intermediary cell, creates outliers in the weight eigenspectrum (Dahmen et al., 2020), subsequently lowering the effective rank. Details of these initial connectivity structures are provided in Appendix C. As illustrated in Figure 2, these connectivity structures, motivated by known features of biological neural networks, exhibit a lower effective rank compared to standard random Gaussian initialization, thereby serving as natural testbeds for our theoretical predictions. To quantify (relative) effective rank, we used $\frac{\sum_i |\lambda_i|}{|\lambda_1|N}$, which indicates the number of eigenvalues on the order of the dominant ones scaled by the network size N.

Figure 3, again tested in RNNs, shows that these different low-rank biologically motivated structures can lead to effectively richer learning compared to the standard random Gaussian initialization. This finding supports our overarching prediction, that lower rank leads to greater richness of learning. We remark that low-rank structures only set network initializations, but are not enforced during training.

Low-rank initialization aligned with task statistics: These simulations may be considered to be within our task-agnostic framework. That is, we have chosen a "random" battery of tasks that is not directly matched to the initial network connectivity structures. Thus, our findings that lower rank

initializations lead to richer learning are expected from our theoretical prediction on the task-averaged alignment (Theorem 1), rather than something task-specific. However, Proposition 1 also predicts that low-rank initialization can lead to lazy learning if the initialization is already aligned to the task structure. To test this (see Figure caption for details), we observe in Figure 4 that a considerably higher alignment can be achieved when the initialization aligns solely with the dominant task features, especially when the relative strength of these dominant features is high. We postulate that such alignment may occur in biological settings if the circuit has evolved to preferentially learn specific tasks.

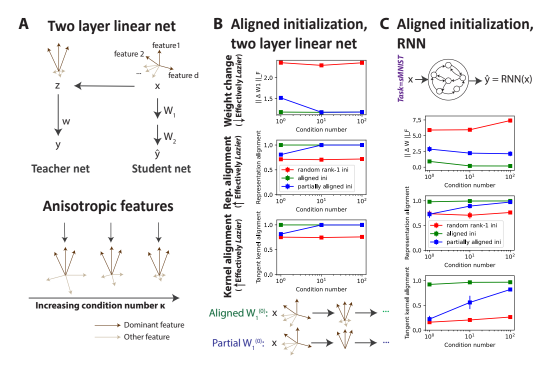


Figure 4: Low-rank initializations can still achieve high alignment for specific tasks (see **Proposition 1).** A) The student-teacher two-layer linear network setup as described in Section 2.3, but with feature anisotropy controlled by a feature modulation matrix F. The condition number of F dictates the relative feature strength. We set the top half of the singular values of F are set to κ , while the bottom half are set to 1, where κ represents the condition number of F. B) The aligned initialization (green) is achieved by setting W_1 as described in Proposition 1 (with $\beta = w^T F$, w is as illustrated), so that the initialization aligns with the task statistics. The partial alignment (blue) mirrors the aligned case, but F is substituted with its rank-(d/2) truncation, causing the network to align only with the dominant features. We observe that a considerably higher alignment can be achieved when the initialization aligns solely with the dominant features, especially when the relative strength of these dominant features is high. (C) The analysis from B) is replicated for RNNs learning the sMNIST task. As the ground truth network function is elusive, we use a teacher network with pre-trained weights. Once again, we replace F with its rank-(d/2) truncation for partial alignment. Details on the input/output definitions and initializations, as well as other simulation specifics, are available in Appendix C. We note that in all scenarios presented here, the initial error remains high since the readout weights are initialized randomly, rendering it a valid learning problem.

4 Discussion

Our investigation casts light on the nuanced influence of initial weight structure, particularly its effective rank, on learning dynamics. Anchored by Theorem 1, our theoretical findings underscore that high-rank random initialization generally facilitates *effectively* lazier learning on average across various task definitions, with an emphasis on the expectation across tasks rather than on a specific task. This is rather intuitive, as in high-dimensional spaces, the probability of two low-rank matrices

(one corresponds to the initial network and the other corresponds to the task) being nearly orthogonal is high. However, certain low-rank initial weights, when naturally predisposed to specific tasks, may lead to effectively lazier learning, suggesting an interesting interplay between evolutionary or developmental biases and learning dynamics (Proposition 1). Our numerical experiments on RNNs further validate these theoretical findings, illustrating the nuanced behavior of low and high-rank initializations in diverse settings (Section 3).

4.1 POTENTIAL IMPLICATIONS TO BOTH NEUROSCIENCE AND DEEP LEARNING

Understanding the shift between rich and lazy learning regimes is paramount not just for its theoretical value, but also for its practical implications in neural systems. Synaptic plasticity, while potent, comes with costs, including energy expenditure and the potential for catastrophic forgetting (McCloskey & Cohen, 1989; Placais & Preat, 2013; Mery & Kawecki, 2005). Our research underscores the nuanced influence of initial weight structure even when the initial weight magnitude remains constant. When an organism learns de novo, as might be the case in early developmental stages or in unpredictable environments, our findings suggest that high-rank initialization generally promotes effectively lazier learning — indicative of minimal shifts in weights and representation. However, when an organism possesses neural circuits refined for specific tasks post-development (or via evolution), the presence of low-rank structures that are already somewhat aligned with the statistics of the world, and the behavior the animal must produce within it, encourages more effective lazy learning compared to the random initialization. Discussion of multiple learning regimes in the brain is in line with certain observed neural phenomena — more subdued alterations post-development preceded by significant synaptic modifications during development (Lohmann & Kessels, 2014) — thereby enriching the discourse on the benefits of dual-regime learning in neural circuits, and emphasizing how it can judiciously curb synaptic changes to conserve resources (Plaçais & Preat, 2013). Such understanding of the link between the initial weight rank and learning dynamics could also guide design choices, potentially improving training efficiency in neuromorphic hardware. Furthermore, the application of regime-specific theoretical deep learning tools to biological neural networks has just started to be explored (Pogodin et al., 2023). Our study contributes to this discourse by suggesting which regime a network operates under based on its initial structure, consequently serving as a bridge, merging tools from deep learning theory with computational neuroscience.

4.2 LIMITATIONS AND FUTURE DIRECTIONS:

Our study predominantly focused on the weight (effective) rank, leaving the exploration of other facets of weight shape on the effective learning regime as an open avenue. Also, the ramifications of effective learning regimes on learning speed — given the known results on kernel alignment and ease of learning (Bartlett et al., 2021) and present mixed findings in the existing literature (Flesch et al., 2021; George et al., 2022) — warrant further exploration. Also, enhancing the breadth of our study necessitates examinations across a broader range of tasks and neural network architectures, and while it is grounded on the backpropagation learning rule, the implications when applied to biologically plausible learning rules are yet to be explored. Additionally, we ensured the consistency of outcomes against factors like width, learning rate, and initial gain (see Appendix D), but the strict NTK regime limit and other factors such as noise remain unexplored (HaoChen et al., 2021). On that note, the study's focus on RNNs with finite task duration prompts further investigation into the implications for tasks with extended time steps and how conclusions for feedforward network depth (Xiao et al., 2020; Seleznova & Kutyniok, 2022) translate to RNN sequence length. Moreover, postulating that effectively lazier learning might preserve representations and rich learning could reduce interference, we identify exploring the impact of effective learning regimes in continual learning (Libby & Buschman, 2021; Lee et al., 2022; Flesch et al., 2023) as a future direction. In fact, tangent kernel alignment, one of the laziness measures examined here, has been exploited for regularization to facilitate continual learning (Doan et al., 2021). Furthermore, a more profound exploration into the neuroscientific implications of effective learning regimes is crucial, particularly how they affect phenomena like "neural collapse" and "information bottleneck", as well as their varied capacities for generalization and unique susceptibilities to spurious correlations (Papyan et al., 2020; Farrell et al., 2022; Tishby & Zaslavsky, 2015; Saxe et al., 2019; Flesch et al., 2021; George et al., 2022; Schuessler et al., 2023). Our current study did not delve into how initial weight rank affects these facets of learning, representing an essential future direction in connecting weight rank to these theoretical implications in both biological and artificial neural networks. Lastly, while our

theoretical derivations primarily target regression problems, sidestepping classification tasks, there remains unexplored terrain regarding the interplay between the number of task classes and weight rank, which is pivotal to uncovering a more precise relationship between the effective learning regime and the initial weight rank (Dubreuil et al., 2022; Gao et al., 2017).

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A EXTENDED DISCUSSIONS ON RELATED WORKS

Theoretical Foundations of Neural Network Regimes and Implications for Neural Circuits: The journey of understanding deep learning systems has borne witness to unprecedented progress

in the mathematical dissection of neural network functionalities (Advani et al., 2020; Jacot et al., 2018; Pezeshki et al., 2021; Baratin et al., 2021; Alemohammad et al., 2020; Yang, 2020; Agarwala et al., 2022; Atanasov et al., 2021; Azulay et al., 2021; Emami et al., 2021). These theoretical findings, until recently confined predominantly to artificial domains, have embarked upon explorations into biological neural networks, elucidating the intricate dynamics of learning and computational properties (Bordelon & Pehlevan, 2022; Liu et al., 2022a; Braun et al., 2022; Ghosh et al., 2023). Among the vanguard of these theoretical endeavors stands the dichotomy of 'rich' and 'lazy' learning regimes. Both lead to task learning, yet they carry distinct ramifications for representation and generalization (Chizat et al., 2019; Flesch et al., 2021; Geiger et al., 2020; George et al., 2022; Ghorbani et al., 2020; Woodworth et al., 2020; Paccolat et al., 2021; Nacson et al., 2022; HaoChen et al., 2021; Flesch et al., 2023). In the 'lazy' regime, which is typically associated with large initial weights, learning predominantly centers on adjusting the readout weights. This leads to minimal alterations in the network weights and representation, while capitalizing on the expansive dimensionality provided by the hidden layer's random projections (Flesch et al., 2021). In contrast, the 'rich' regime, defined by smaller initial weights, fosters the development of highly tailored hidden unit representations specifically aligned with task demands, resulting in considerable adaptations in weights and representation. It's essential to highlight that the transition and dominance between these regimes are influenced by more than just the initial weight scale. Other factors, ranging from network width to the output gain (often referred to as the α parameter), play a pivotal role (Chizat et al., 2019; Geiger et al., 2020).

A nexus between deep learning theoretical frameworks and neuroscience has unveiled applications of the rich/lazy regimes. Previous investigations characterized neural network behaviors under distinct regimes (Bordelon & Pehlevan, 2022; Schuessler et al., 2023) and discerning which mode yields solutions mimicking empirical data (Flesch et al., 2021). It is compelling to observe that the existence of multiple learning regimes isn't an isolated phenomenon in artificial systems; analogous learning patterns echo in neural circuits as well. For instance, while plasticity-driven transformations might be resource-intensive, they manifest robustly during such developmental phases, followed by minor changes afterwards (Lohmann & Kessels, 2014). Building upon these findings, our research delves deeper into the precursors of these regimes. We examine how inherent factors in the brain, especially initial weight configurations, influence the inclination towards either rich or lazy learning. This understanding is crucial for assessing the applicability of regime-specific tools in neural contexts and for shedding light on the potential benefits of having both learning regimes coexist in the brain.

Interplay of Neural Learning and structure: Understanding how the brain learns using its myriad elements is a perennial quest in neuroscience. Addressing this, certain studies have unveiled biologically plausible learning rules (Lillicrap et al., 2020; Scellier & Bengio, 2017; Diederich & Opper, 1987; Hinton, 2022; Laborieux & Zenke, 2022; Greedy et al., 2022; Sacramento et al., 2018; Payeur et al., 2021; Roelfsema & Holtmaat, 2018; Meulemans et al., 2022; Murray, 2019; Bellec et al., 2020; Liu et al., 2021; 2022b; Marschall et al., 2020), suggesting potential neural algorithms involving known neural ingredients. Concurrently, given the three primary components of a neural network's design — task, learning rule, and architecture — another avenue of research delves deep into the architectural facet, specifically focusing on how it interacts with the learning rule to enhance learning (Richards et al., 2019; Zador, 2019; Yang & Molano-Mazón, 2021). Under the structural umbrella, the neural unit's complexity and initial connectivity patterns are two crucial aspects. Complex neuron models, for instance, have shown the potential in boosting learning performance by allowing implicit forms of memory and computations at the single neuron level (Salaj et al., 2021; Winston et al., 2023). Moreover, A large body of work has investigated the effect of different random initializations on learning in deep networks (Saxe et al., 2013; Bahri et al., 2020; Glorot & Bengio, 2010; He et al., 2015; Arora et al., 2019). For instance, the variance in random initial weights can induce pronounced shifts in network behavior, ranging from the "lazy" to the "rich" regimes (Chizat et al., 2019; Flesch et al., 2021). This introduces unique inductive biases during the learning process, with distinct preferences for learning certain features (George et al., 2022). Our discourse primarily orbits around connectivity and its implications on learning dynamics in networks with simple rectified units. Our results sit within the purview of these regimes, with a widely adopted assumption of gradient

descent via backpropagation as the learning rule, while remaining open to encompassing a wider spectrum of rules in future explorations.

Neural circuit connectivity pattern and eigenspectrum: While the importance of initial weights on function and learning is clear, the impact of specific weight shapes, apart from weight scale, on rich or lazy dynamics remains less explored. The predominant focus in the literature has been on random initialization. Yet, neural circuit structures significantly diverge from this paradigm. Illustratively, one finds connectivity principles or patterns markedly different from what one observes with a mere random initialization (Pogodin et al., 2023), resulting in distinct neural dynamics; these connectivity principles or patterns include Dale's law (Rajan & Abbott, 2006; Ipsen & Peterson, 2020; Harris et al., 2022; Liu et al., 2020), an over-representation of higher-order motifs (Dahmen et al., 2020) and cell-type-specific connectivity statistics (Aljadeff et al., 2015), to name a few. Given the prominence of low-rankedness observed in neural circuits (Song et al., 2005), our study centers on the influence of effective rank on the effective learning regime. As the next generation of connectivity data becomes available (Campagnola et al., 2022; Consortium et al., 2021; Dorkenwald et al., 2022; Winnubst et al., 2019; Scheffer et al., 2020), future explorations will broaden the scope to other weight characteristics.

B PROOFS

B.1 Proofs for main text Theorem and Proposition

Notation Let $f(x) = W_2W_1x$ denote a two-layer linear network with N hidden units on d-dimensional inputs $x \in \mathbb{R}^d$, with weight matrices $W_1 \in \mathbb{R}^{N \times d}$ and $W_2 \in \mathbb{R}^{1 \times N}$. We consider m training inputs $x_1, \cdots x_m$ and the corresponding data matrix $X = [x_1^T \cdots x_m^T] \in \mathbb{R}^{d \times m}$; the output target is generated from a linear teacher network as $Y = \beta^T X$, where $\beta_i \sim \mathcal{N}(0, 1/d)$.

Since our goal is to investigate how the *shape* of the initial weights impacts network change, we will consider a fixed small (Froebenius) norm for these; i.e.,

$$||W_1^{(0)}||_F = ||W_2^{(0)}||_F := \sigma \ll 1$$

We denote by s_1, \dots, s_d denote the singular values of $W_1^{(0)}$; they satisfy $\sum_{j=1}^d s_j^2 = \sigma^2$.

In what follows we focus on the whitened setting, where X has all its non zero singular values equal to 1. We also assume $m \ge d$ for simplicity (this assumption can easily be relaxed in our analysis), so that the whitened data assumption translates as $XX^T = I_d$.

Prior results Our analysis builds on prior results Atanasov et al. (2021) on the evolution of the NTK for two-layer linear networks trained by gradient flow of the mean square error. In the above setting, Atanasov et al. (2021) show that the final NTK $K^{(f)}$ (i.e. the asymptotic NTK as the number of iterations goes to infinity) is given by

$$K^{(f)} = ||\beta||X^{T}(\hat{\beta}\hat{\beta}^{T} + I_{d})X + O(\sigma^{2}).$$
(8)

where $\hat{\beta} := \beta/\|\beta\|$. We are interested in the expected kernel alignment over the tasks, in the small initialization regime:

$$\mathbb{E}_{\beta}[KA(K^{(f)}, K^{(0)})] := \mathbb{E}_{\beta}\left[\frac{\text{Tr}(K^{(f)}K^{(0)})}{\|K^{(f)}\|_F \|K^{(0)}\|_F}\right]. \tag{9}$$

Theorem 1. In the above setting, when considering all possible initializations $W_1^{(0)}$ with small fixed norm σ , the expected kernel alignment $\mathbb{E}_{\beta}[KA]$ (defined in Eq. 9) is maximized with high-rank isotropic initialization, i.e with $W_1^{(0)}$ that has all its non-zero singular values equal in absolute value.

Proof. Let us write $K^{(0)} = X^T M_0 X$ with $M_0 := W_1^{(0)T} W_1^{(0)} + \sigma^2 I_d$. Up to $O(\sigma^4)$ terms, the numerator in Eq. 9 takes the form

$$\operatorname{Tr}(K^{(f)}K^{(0)}) = \|\beta\| \operatorname{Tr}(X^{T}(\hat{\beta}\hat{\beta}^{T} + I_{d})XX^{T}M_{0}X)$$

$$\stackrel{(a)}{=} \|\beta\| \operatorname{Tr}(X^{T}(\hat{\beta}\hat{\beta}^{T} + I_{d})M_{0}X)$$

$$\stackrel{(b)}{=} \|\beta\| \operatorname{Tr}((\hat{\beta}\hat{\beta}^{T} + I_{d})M_{0}XX^{T})$$

$$\stackrel{(a)}{=} \|\beta\| \operatorname{Tr}((\hat{\beta}\hat{\beta}^{T} + I_{d})M_{0})$$

$$\stackrel{(c)}{=} \|\beta\|(\hat{\beta}^{T}M_{0}\hat{\beta} + \operatorname{Tr}M_{0})$$

$$(10)$$

where (a) uses $XX^T = I_d$, (b) the cyclicity of the trace, and (c) the fact that $\hat{\beta}^T M_0 \hat{\beta}$ is a scalar. As for the denominator in Eq. 9), we have,

$$||K^{(0)}||_F^2 = \text{Tr}(K^{(0)}K^{(0)})$$

$$= \text{Tr}(X^T M_0 X X^T M_0 X)$$

$$\stackrel{(a)}{=} \text{Tr}(M_0^2)$$
(11)

and, up to $O(\sigma^4)$ terms,

$$||K^{(f)}||_F^2 = \operatorname{Tr}(K^{(f)}K^{(f)})$$

$$= ||\beta||^2 \operatorname{Tr}(X^T(\hat{\beta}\hat{\beta}^T + I_d)XX^T(\hat{\beta}\hat{\beta}^T + I_d))$$

$$= ||\beta||^2 \operatorname{Tr}(X^T(\hat{\beta}\hat{\beta}^T + I_d)X)$$

$$\stackrel{(a)}{=} ||\beta||^2 \operatorname{Tr}(\hat{\beta}\hat{\beta}^T + I_d)^2$$

$$\stackrel{(b)}{=} ||\beta||^2 (d+3)$$
(12)

where (a) in these two calculations uses $XX^T=I_d$ and the cyclicity of the trace; and (b) notes that the $d\times d$ matrix $\hat{\beta}\hat{\beta}^T+I_d$ has d-1 eigenvalues equal to 1 and one equal to 2. Eq. 11 and 12 yield

$$||K^{(f)}||_F ||K^{(0)}||_F = ||\beta|| \sqrt{(d+3)\operatorname{Tr} M_0^2}$$
(13)

Putting together Eq. 10, 13, we obtain, up to additive $O(\sigma^2)$ terms,

$$KA(K^{(f)}, K^{(0)}) = \frac{\hat{\beta}^T M_0 \hat{\beta} + Tr M_0}{\sqrt{(d+3) Tr M_0^2}}$$
(14)

Next, averaging over the tasks requires computing the Gaussian average

$$A[M_0] := \mathbb{E}_{\beta} \left[\hat{\beta}^T M_0 \hat{\beta} \right] = \mathbb{E}_{\beta} \left[\frac{\beta^T M_0 \beta}{\|\beta\|^2} \right].$$

Lemma 1. The map A is invariant under the action of the orthogonal group, i.e $A[UMU^T] = A[M]$ for all M and all orthogonal matrix U.

Proof. This is a consequence of the invariance of the Gaussian measure under the action of the orthogonal group. Explicitly, given an orthogonal matrix U,

$$A[UMU^{T}] = \frac{1}{(2\pi d)^{d/2}} \int d^{d}\beta e^{-\|\beta\|^{2}/d} \left[\frac{\beta^{T}UMU^{T}\beta}{\|\beta\|^{2}} \right]$$

$$\beta' := U^{T}\beta \frac{1}{(2\pi d)^{d/2}} \int d^{d}\beta' |\det U| e^{-\|U\beta'\|^{2}/d} \left[\frac{\beta'^{T}M\beta'}{\|U\beta'\|^{2}} \right]$$

$$= \frac{1}{(2\pi d)^{d/2}} \int d^{d}\beta' e^{-\|\beta'\|^{2}/d} \left[\frac{\beta'^{T}M\beta'}{\|\beta'\|^{2}} \right]$$

$$= A[M]$$
(15)

where the third equality follows from $|\det U| = 1$ and $||U\beta|| = ||\beta||$.

Lemma 2. There is a constant c such that $A[M] = c\operatorname{Tr}(M)$ for any symmetric matrix M.

Proof. Given a symmetric matrix M, it can be diagonalized as $M = UDU^T$ where $D = \text{Diag}(\mu_1, \dots \mu_d)$ is diagonal and U is orthogonal. By rotation invariance, we have A[M] = A[D], and

$$A[D] = \mathbb{E}_{\beta} \left[\hat{\beta}^T D \hat{\beta} \right] = \mathbb{E}_{\beta} \left[\sum_{j=1}^d \hat{\beta}_j^2 \mu_j \right] = \sum_{j=1}^d \mathbb{E}_{\beta} \left[\frac{\beta_j^2}{\|\beta\|^2} \right] \mu_j := \sum_{j=1}^d c_j \mu_j$$
 (16)

We conclude by noting that, by invariance of the (isotropic) Gaussian measure under permutation of the vector components, the coefficients c_j are independent of j, i.e $c_j \equiv c$ for all j. In sum,

$$A[M] = A[D] = c \operatorname{Tr} D = c \operatorname{Tr} M. \tag{17}$$

The expected kernel alignment thus takes the form,

$$\mathbb{E}_{\beta}[KA(K^{(f)}, K^{(0)})] = \frac{(1+c)\operatorname{Tr} M_0}{\sqrt{(d+3)\operatorname{Tr} M_0^2}}$$
(18)

up to additive $O(\sigma^2)$ terms. Finally, we note that

$$\operatorname{Tr} M_0 = \operatorname{Tr}(W_1^{(0)T} W_1^{(0)} + \sigma^2 I_d)$$

$$= \|W_1^{(0)}\|_F^2 + d\sigma^2$$

$$= (d+1)\sigma^2$$
(19)

and

$$\operatorname{Tr} M_0^2 = \operatorname{Tr}(W_1^{(0)T} W_1^{(0)} + \sigma I_d)^2$$

$$= \sum_{j=1}^d (s_j^2 + \sigma^2)^2$$

$$= \sum_{j=1}^d s_j^4 + 2\sigma^2 \sum_{j=1}^d s_j^2 + d\sigma^4$$

$$= \sum_{j=1}^d s_j^4 + (d+2)\sigma^4$$
(20)

Substituting into Eq. 18, we have, up to additive $O(\sigma^2)$ terms,

$$\mathbb{E}_{\beta}[KA(K^{(f)}, K^{(0)})] = \frac{(1+c)(d+1)}{\sqrt{(d+3)(d+2+\sum_{j=1}^{d} (s_j/\sigma)^4)}}$$
(21)

Finally, we see in Eq 21 that the maximization of $\mathbb{E}_{\beta}[KA]$ reduces to the following convex constrained optimization problem:

$$\min_{s} \sum_{j} s_{j}^{4}, \quad \text{subject to } \sum_{j} s_{j}^{2} = \sigma^{2}. \tag{22}$$

The KKT solutions satisfy $s_i^2 = \sigma^2/d$ for all $j = 1 \cdots d$. This implies that the expected tangent kernel alignment is maximized when the initial weight singular values $|s_i|$ are distributed evenly across dimensions, which corresponds to a high-rank initialization.

Proposition 1. Following the setup and assumptions in Theorem 1, rank-1 initialization with $W_1^{(0)} = \sigma[\hat{\beta}^T \ \vec{0} \ ... \ \vec{0}]$ leads to maximal alignment, i.e, $KA(K^{(f)}, K^{(0)}) = 1$ up to additive $O(\sigma^2)$ terms.

Proof. We indeed have,

$$K^{(0)} = X^{T} (W_{1}^{(0)^{T}} W_{1}^{(0)} + \|W_{2}^{(0)}\|^{2} I) X$$
$$= \sigma^{2} X^{T} (\hat{\beta} \hat{\beta}^{T} + I) X$$
(23)

Thus, writing $K := X^T (\hat{\beta} \hat{\beta}^T + I) X$ and using Eq. 8, the alignment takes the form

$$KA(K^{(f)}, K^{(0)}) := \frac{Tr(K^{(f)}K^{(0)})}{\|K^{(f)}\|_F \|K^{(0)}\|_F}$$

$$= \frac{Tr(K(K + O(\sigma^2))}{\|K\|_F \|K + O(\sigma^2)\|_F}$$

$$= \frac{Tr(K^2)}{\|K\|_F^2} + O(\sigma^2)$$

$$= 1 + O(\sigma^2)$$
(24)

B.2 Learning requirement based on $W_h^{(0)}$ rank

The focus of this idea is to show that no changes to hidden weights W_h is not possible (e.g. reservoir settings) for zero-error when the initial weight rank falls below a certain threshold. Freezing the hidden weights W_h would be a special case of lazy learning.

Proposition 2. Consider a linear RNN with input at time t as $X_t \in \mathbb{R}^{N \times d}$ (for t = 1, ..., T - 1), target output $Y \in \mathbb{R}^{N_{out} \times d}$ only at the last step, recurrent weight matrix $W_h \in \mathbb{R}^{N \times N}$ and readout weight matrix $w \in \mathbb{R}^{N_{out} \times N}$. Here, N, N_{out}, d and T are the number of hidden units, number of classes, number of data points and number of time steps, respectively, and we assume $N, d > N_{out}$. Define initial recurrent weight $W_h^{(0)}$ and final recurrent weight $W_h^{(f)}$ that achieves zero error. Then, for arbitrary input X and target output $Y, W_h^{(f)} = W_h^{(0)}$ is not possible when $rank(W_h^{(0)}) < N_{out}$.

Proof. We have the following based on the assumption of the RNN structure, if zero-error learning is achieved:

$$Y = w^{(f)} W_h^{(f)} \left(\sum_{t=1}^{T-1} W_h^{(f)^{T-t-1}} X_t \right).$$
 (25)

We can prove by contradiction. Suppose $W_h^{(f)} = W_h^{(0)}$, then

$$Y = w^{(f)} W_h^{(0)} \left(\sum_{t=1}^{T-1} W_h^{(0)T-t-1} X_t \right).$$
 (26)

Since Y is arbitrary, we can have $rank(Y) = N_{out}$ (by the assumption of $N, d > N_{out}$). Applying $rank(W_h^{(0)}) < N_{out}$ we have

$$rank(Y) = rank(w^{(f)}W_h^{(0)} \left(\sum_{t=1}^{T-1} W_h^{(0)^{T-t-1}} X_t\right))$$

$$\stackrel{(a)}{\leq} min(rank(w^{(f)}), rank(W_h^{(0)}), \left(\sum_{t=1}^{T-1} W_h^{(0)^{T-t-1}} X_t\right))$$

$$< N_{out}, \tag{27}$$

where (a) is because $rank(W_h^{(0)}) < N_{out}$ so the minimum has to be less than N_{out} . This would contradict an arbitrary Y with $rank(Y) = N_{out}$. Thus, $W_h^{(f)} = W_h^{(0)}$ cannot happen and recurrent weights have to be adjusted.

C SETUP AND SIMULATION DETAILS

C.1 INITIAL LOW-RANK WEIGHTS CREATION

For the null case, we initialized with random Gaussian distributions where each weight element $W_{ij} \sim \mathcal{N}(0, g^2/N)$, with an initial weight variance of g. Unless otherwise mentioned, we set g=1.5 and network size N=300, though we also validated across other parameter choices (see Appendix D).

To create low-rank weights using SVD, we generated temporary weights $\hat{W}_{ij} \sim \mathcal{N}(0, g^2/N)$. Subsequently, we applied SVD to \hat{W} and retained the top components based on the desired rank. To ensure comparisons are made across constant initial weight magnitudes, the resultant weight matrix was rescaled to match the Frobenius norm of \hat{W} .

Furthermore, we present details for experimentally-driven low-rank weights. For block-specific statistics, we followed the setup in Figure S3 of Aljadeff et al. (2015), setting parameters as $\alpha = 0.02$, $\gamma = 10$, and $1 - \epsilon = 0.8$. These parameters substantially influence the weight eigenspectrum, as depicted in Figure S3 of Aljadeff et al. (2015); we selected these values specifically to emphasize the outliers and achieve a lower effective rank. These parameters represent the fraction of hyperexcitable neurons (population 1), gain of hyperexcitable connections, and the gain of remaining connections, respectively. For the creation of a chain motif, we employed the procedure described in Section S3.10 of Dahmen et al. (2020), setting $\tau_{chn} = 0.03$ (and $\tau_{chn} = -0.1$ for over-representation or under-representation of the chain motif, respectively). Here, we set N=100. These parameters were chosen to provide enough distinctions from the null case, while still ensuring stability and effective task learning. The electron microscopy (EM) connectivity (of the V1 cortical column model) is obtained from Institute (2023), which includes dendritic tree reconstructions and local axonal projections for hundreds of thousands of neurons, detailing their 0.5 billion synaptic connections. From this, we selected 198 cells, focusing on fully proofread neurons closest to the midpoint between layers 2/3 and 4. Connectivity strength for each neuron is determined by summing the volume of each post-synaptic density to target cells, distinguishing between excitatory and inhibitory cell types. For instance, if cell 'a' forms 10 synapses with cell 'b', the connection strength of connection[a,b] represents the combined volume of synaptic densities at cell 'b'. Inhibitory connections are assigned a sign of -1, while excitatory ones receive +1. For the Dale's law obeying initial connectivity, initialization was done following the process in Yang & Wang (2020) with 80% excitatory and 20% inhibitory neurons (see the notebook EIRNN.ipynb).

It's crucial to highlight that, in testing our Theorem, which examines the effect of the **initial** weight rank, all low-rank modifications are not enforced during training (although the impact of enforcing these structures could be an interesting avenue for future exploration). Weights are adjusted freely based on gradient descent learning.

C.2 TASK AND TRAINING DETAILS

Our code is accessible at https://anonymous.4open.science/r/BioRNN_RichLazy-59F0/README.md. We used PyTorch Version 1.10.2 (Paszke et al., 2019). Simulations were executed on a computer server with x2 20-core Intel(R) Xeon(R) CPU E5-2698 v4 at 2.20GHz, with the average task training duration being around 10 minutes. Following the procedure in George et al. (2022), which delved deeply into effective laziness metrics, we employed gradient-descent learning with the SGD optimizer. Unless mentioned otherwise, the learning rate was 3e-3, but we validated that our findings remain consistent across various learning rates (see Appendix D). All weights — input, recurrent and readout — were trained.

For the neuroscience tasks, we adopted the Neurogym framework (Molano-Mazon et al., 2022). Within this paper, these tasks are denoted as "2AF", "DMS", and "CXT", mirroring Neurogym settings: task =' PerceptualDecisionMaking - v0', task =' DelayMatchSample - v0', and task =' ContextDecisionMaking - v0', respectively. To expedite simulations and facilitate numerous runs, we operated with $dt = \tau_m = 100ms$ and abbreviated task durations: for 2AF, settings were stimulus = 700ms and decision = 100ms; for DMS, they were sample = 100ms, delay = 500ms, test = 100ms, and decision = 100ms; for CXT, they comprised stimulus = 100ms

200ms, delay = 500ms, and decision = 100ms. For these three tasks, we used a batch size of 32 and trained for 10000 iterations.

Regarding the sequential MNIST task LeCun (1998), we employed a row-by-row format to hasten simulations. Inputs were delivered via $N_{in}=28$ units, each presenting a row's grey-scaled value, culminating in 28 steps with network predictions rendered at the final step. Training hinged on the cross-entropy loss function; targets were provided throughout training for the neuroscience tasks, as per Neurogym implementation, and targets were provided at only a trial's conclusion for the sequential MNIST task. For this task, we used a batch size of 200 and trained for 10000 iterations.

For the student-teacher two-layer linear network simulations in Figure 10 and Figure 4, we set N=1000, d=2 (also found similar trends for d=20 and d=100), z=Fx, all entries of w (or β when F=I) to 1 and entries of X are sampled from a uniform distribution over the interval [-2,2]. We used standard Normal initialization for both W_1 and W_2 with $\sigma=0.001$. For Figure 4, F is constructed from SVD, i.e. $F=USV^T$, with U and V generated from arbitrary orthogonal matrices, and S is a diagonal matrix consisting of the singular values with the top half of the singular values set to κ and bottom half set to 1, where κ is the condition number of F. For the aligned initialization, W_1 is initialized as given in Proposition 1 with $\beta=w^TF$ (w here is illustrated in Figure 4), and the F is replaced by its rank-(d/2) truncation for the partially aligned initialization case.

D ADDITIONAL SIMULATIONS

We perform additional simulations to show the robustness of our main trends, Low-rank initial recurrent weights lead to greater changes (or effectively richer learning) in RNNs. We show the main trends observed in Figure 1 holds also for Uniform initialization (Figure 5), soft initial weight rank (Figure 6), various network sizes (Figure 7), learning rates (Figure 8), gains (Figure 9). The trends in Figure 3 also applies to the DMS task (Figure 11) and the CXT task (Figure 12). Also, without the low-rankedness in the shuffled EM connectivity, the impact on effective laziness also goes away (Figure 13). In Figure 10 we confirm that the results, shown in Figure 1 and predicted by Theorem 1, are also observed in a two-layer linear network setup. Again, we find that in situations where initializations are random, higher rank initialization leads to greater tangent kernel alignment than lower rank cases.

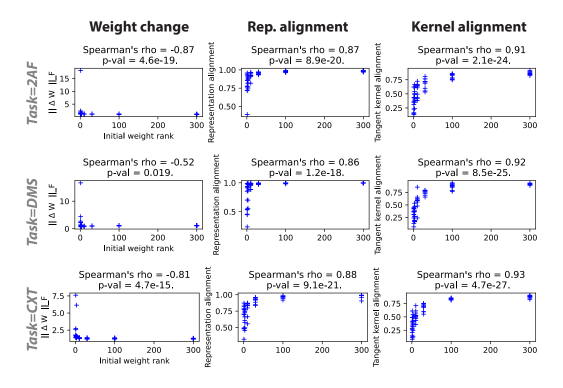


Figure 5: Consistent trends observed in Figure 1 also for Uniform initialization. We replicated the results of Figure 1 — where the initial weights follow a zero-mean Gaussian distribution $W_{ij} \sim \mathcal{N}(0, g^2/N)$ — but now for Uniform initialization $W_{ij} \sim \mathcal{U}\left(-\frac{g}{\sqrt{N}}, \frac{g}{\sqrt{N}}\right)$. Plotting conventions follow that of Figure 1.

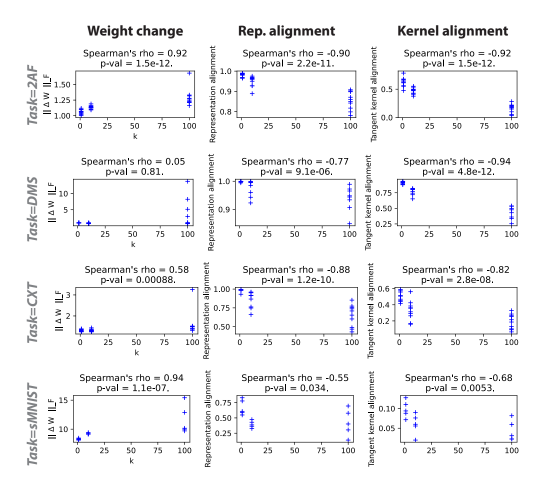


Figure 6: Consistent trends observed in Figure 1 also for "softer" low-rank weights. Here, instead of the "hard" low-rank weights in Figure 1 — where the i^{th} weight singular value s_i is set to 0 if i>r for rank r — we introduce a smoother decay in singular value, where we replace the singular values with $s_i=s_1(1-i/N)^k$ after performing SVD; this means that greater k leads to lower effective rank. Plotting conventions follow that of Figure 1

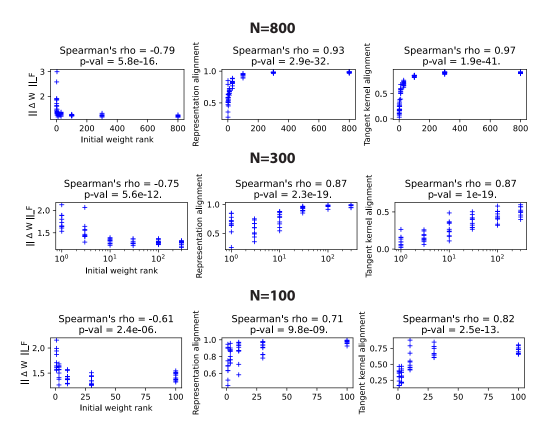


Figure 7: Consistent trends observed in Figure 1 across various network sizes (N). We replicated the results of Figure 1 for different values of N, using the CXT task as an illustrative example. However, the observed trend remains consistent for both the 2AF and DMS tasks. Plotting conventions follow that of Figure 1.

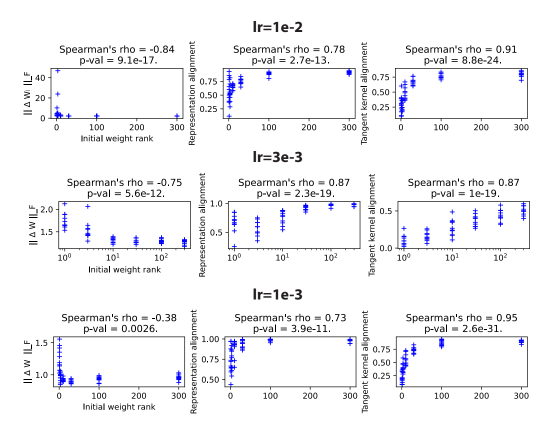


Figure 8: Consistent trends observed in Figure 1 across various learning rates (lr). We replicated the results of Figure 1 for different learning rates, using the CXT task as an illustrative example. However, the observed trend remains consistent for both the 2AF and DMS tasks. Plotting conventions follow that of Figure 1.

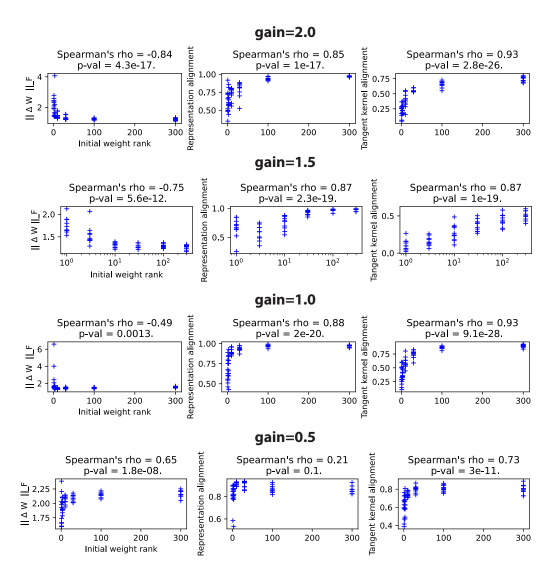


Figure 9: Consistent trends observed in Figure 1 across various initial gain. Here, the gain refers to g, as weights are initialized as $W_{ij} \sim \mathcal{N}(0, g^2/N)$. The trends hold for most typical range of g from 1.0 to 2.0, but gets weakened for smaller values, g < 1.0 (a closer examination of the regime bias in such setting in RNNs is left for future work). We replicated the results of Figure 1 for different learning rates, using the CXT task as an illustrative example. However, the observed trend remains consistent for both the 2AF and DMS tasks. Plotting conventions follow that of Figure 1.

Student-teacher setup for two layer linear net

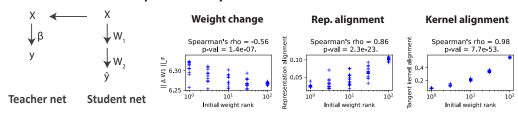


Figure 10: As predicted by the theoretical results, higher rank random initialization leads to effectively lazier in two-layer linear network. We use the student-teacher two-layer linear network setup described in Section 2.3. Plotting convention follows that of Figure 1.

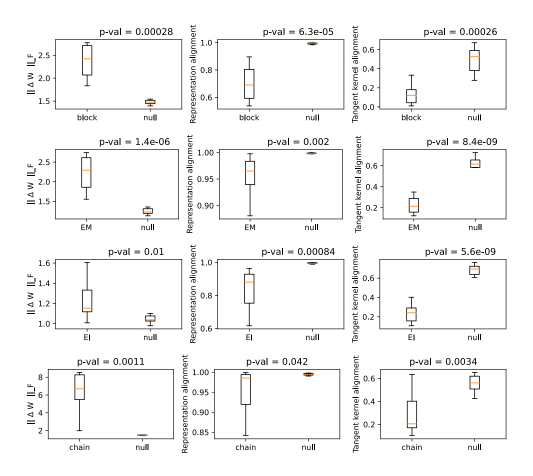


Figure 11: We repeated Figure 3 for the DMS task and observed similar trends: low-rank initialization, achieved by experimentally-driven initial connectivity in Figure 2, leads to effectively richer learning.

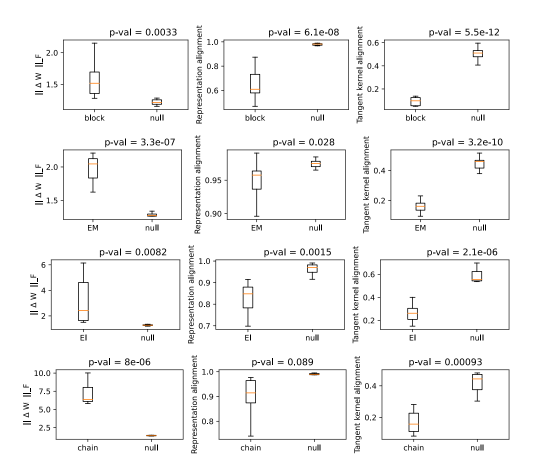


Figure 12: We repeated Figure 3 for the CXT task and observed similar trends: low-rank initialization, achieved by experimentally-driven initial connectivity in Figure 2, leads to effectively richer learning.

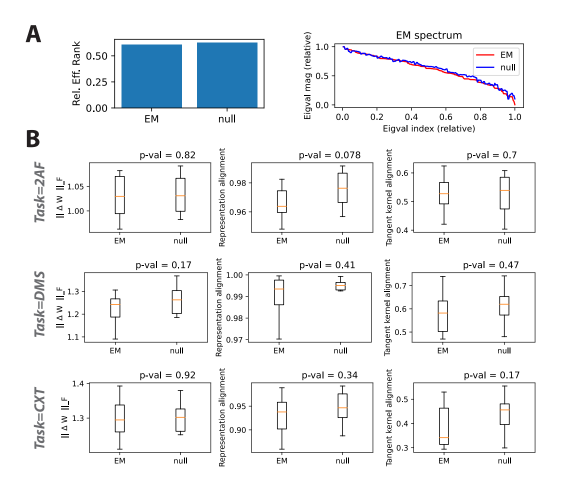


Figure 13: Shuffling the EM connectivity destroys the low-rankedness and the impact on effective laziness. We repeated the analyses with the EM initial connectivity in Figures 2 and 3 but performed random shuffling on the EM connectivity, to see if the low-rankedness and the impact on effective laziness is due to the sparsity in the dataset. Performing such shuffling destroys these trends. Plotting conventions follow that of Figures 2 and 3.