TRUTH OR BACKPROPAGANDA? AN EMPIRICAL IN-VESTIGATION OF DEEP LEARNING THEORY

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Paper under double-blind review

Abstract

We empirically evaluate common assumptions about neural networks that are widely held by practitioners and theorists alike. We study the prevalence of local minima in loss landscapes, whether small-norm parameter vectors generalize better (and whether this explains the advantages of weight decay), whether widenetwork theories (like the neural tangent kernel) describe the behaviors of classifiers, and whether the rank of weight matrices can be linked to generalization and robustness in real-world networks.

1 INTRODUCTION

Modern deep learning methods are descendent from such long-studied fields as statistical learning, optimization, and signal processing, all of which were built on mathematically rigorous foundations. In statistical learning, principled kernel methods have vastly improved the performance of SVMs and PCA (Suykens & Vandewalle, 1999; Schölkopf et al., 1997), and boosting theory has enabled weak learners to generate strong classifiers (Schapire, 1990). Optimizers in deep learning are borrowed from the field of convex optimization , where momentum optimizers (Nesterov, 1983) and conjugate gradient methods provably solve ill-conditioned problems with high efficiency (Hestenes & Stiefel, 1952). Deep learning harnesses foundational tools from these mature parent fields.

Despite its rigorous roots, deep learning has driven a wedge between theory and practice. Recent theoretical work has certainly made impressive strides towards understanding optimization and generalization in neural networks. But doing so has required researchers to make strong assumptions and study restricted model classes.

In this paper, we seek to understand whether deep learning theories accurately capture the behaviors and network properties that make realistic deep networks work. Following a line of previous work, such as Swirszcz et al. (2016), Zhang et al. (2016), Balduzzi et al. (2017) and Santurkar et al. (2018), we put the assumptions and conclusions of deep learning theory to the test using experiments with both toy networks and realistic ones. We focus on the following important theoretical issues:

- Local minima: Numerous theoretical works argue that all local minima of neural loss functions are globally optimal or that all local minima are nearly optimal. In practice, we find highly suboptimal local minima in realistic neural loss functions, and we discuss reasons why suboptimal local minima exist in the loss surfaces of deep neural networks in general.
- Weight decay and parameter norms: Research inspired by Tikhonov regularization suggests that low-norm minima generalize better, and for many, this is an intuitive justification for simple regularizers like weight decay. Yet for neural networks, it is not at all clear which form of ℓ_2 -regularization is optimal. We show this by constructing a simple alternative: Biasing solutions toward a non-zero norm still works and can even measurably improve performance for modern architectures.
- Rank: Generalization theory has provided guarantees for the performance of low-rank networks. However, we find that regularization which encourages high-rank weight matrices often outperforms that which promotes low-rank matrices. This indicates that low-rank structure may not be a significant force behind generalization in practical networks. We further investigate the adversarial robustness of low-rank networks, which are thought to be more resilient to attack, and we find empirically that their robustness is often lower than the baseline or even a purposefully constructed high-rank network.

 Neural tangent kernels and the wide-network limit: We investigate theoretical results concerning neural tangent kernels of realistic architectures. Astonishingly, stochastic sampling of the tangent kernels suggests that theoretical results on tangent kernels of multi-layer networks may transfer to realistic settings and initializations for multilayer networks and basic convolutional architectures. However, this transfer holds best for extremely wide networks, and does not reliably hold in the regime where practical deep networks live and even reverses when considering ResNet architectures.

2 LOCAL MINIMA IN LOSS LANDSCAPES: DO SUBOPTIMAL MINIMA EXIST?

It is generally accepted that "in practice, poor local minima are rarely a problem with large networks." (LeCun et al., 2015). However, exact theoretical guarantees for this statement are elusive. Various theoretical studies of local minima have investigated spin-glass models (Choromanska et al., 2014), deep linear models (Laurent & Brecht, 2018; Kawaguchi, 2016), parallel subnetworks (Haeffele & Vidal, 2017), and dense fully connected models (Nguyen et al., 2018) and have shown that either all local minima are global or all have a small optimality gap. The apparent scarcity of poor local minima has lead practitioners to develop the intuition that bad local minima ("bad" meaning high loss value and suboptimal training performance) are practically non-existent.

To further muddy the waters, some theoretical works prove the *existence* of local minima. Such results exist for simple fully connected architectures (Swirszcz et al., 2016), single-layer networks (Liang et al., 2018), and two-layer ReLU networks (Safran & Shamir, 2017). Unfortunately, existing analysis of neural loss landscapes requires strong assumptions (e.g. random training data, linear activation functions, fully connected layers, or extremely wide network widths) — so strong, in fact, that it is reasonable to question whether these results have bearing on practical neural networks or describe the underlying cause of good optimization performance in real-world settings.

In this section, we investigate the existence of suboptimal local minima from a theoretical perspective and an empirical one. If suboptimal local minima exist, they are certainly hard to find by standard methods (otherwise training would not work). Thus, we present simple theoretical results that inform us on how to construct non-trivial suboptimal local minima, concretely generalizing previous constructions, such as those by Swirszcz et al. (2016). Using experimental methods inspired by theory, we easily find suboptimal local minima in the loss landscapes of a range of classifiers.

Trivial local minima are easy to find in ReLU networks – consider the case where bias values are sufficiently low so that the ReLUs are "dead" (i.e. inputs to ReLUs are strictly negative). Such a point is trivially a local minimum. Below, we make a more subtle observation that multilayer perceptrons (MLPs) must have non-trivial local minima, provided there exists a linear classifier that performs worse than the neural network (an assumption that holds for virtually any standard benchmark problem). Specifically, we show that MLP loss functions contain local minima where they behave identically to a linear classifier on the same data.

We now define a family of low-rank linear functions which represent an MLP. Let "rank-s affine function" denote an operator of the form $G(\mathbf{x}) = A\mathbf{x} + \mathbf{b}$ with rank(A) = s.

Definition 2.1. Consider a family of functions, $\{F_{\phi} : \mathbb{R}^m \to \mathbb{R}^n\}_{\phi \in \mathbb{R}^P}$ parameterized by ϕ . We say this family has *rank-s affine expression* if for all rank-s affine functions $G : \mathbb{R}^m \to \mathbb{R}^n$ and finite subsets $\Omega \subset \mathbb{R}^m$, there exists ϕ with $F_{\phi}(\mathbf{x}) = G(\mathbf{x}), \forall \mathbf{x} \in \Omega$. If $s = \min(n, m)$ we say that this family has *full affine expression*.

We investigate a family of L-layer MLPs with ReLU activation functions, $\{F_{\phi} : \mathbb{R}^m \to \mathbb{R}^n\}_{\phi \in \Phi}$, and parameter vectors ϕ , i.e., $\phi = (A_1, \mathbf{b}_1, A_2, \mathbf{b}_2, \dots, A_L, \mathbf{b}_L)$, $F_{\phi}(\mathbf{x}) = H_L(f(H_{L-1}...f(H_1(\mathbf{x}))))$, where f denotes the ReLU activation function and $H_i(\mathbf{z}) = A_i\mathbf{z} + \mathbf{b}_i$. Let $A_i \in \mathbb{R}^{n_i \times n_{i-1}}$, $\mathbf{b}_i \in \mathbb{R}^{n_i}$ with $n_0 = m$ and $n_L = n$.

Lemma 1. Consider a family of L-layer multilayer perceptrons with ReLU activations $\{F_{\phi} : \mathbb{R}^m \to \mathbb{R}^n\}_{\phi \in \Phi}$, and let $s = \min_i n_i$ be the minimum layer width. Such a family has rank-s affine expression.

Proof. The idea of the proof is to use the singular value decomposition of any rank-s affine function to construct the MLP layers and pick a bias large enough for all activations to remain positive. See Appendix A.1. \Box

The ability of MLPs to represent linear networks allows us to derive a theorem which implies that arbitrarily deep MLPs have local minima at which the performance of the underlying model on the training data is equal to that of a (potentially low-rank) linear model. In other words, neural networks inherit the local minima of elementary linear models.

Theorem 1. Consider a training set, $\{(\mathbf{x}_i, y_i)\}_{i=1}^N$, a family $\{F_{\phi}\}_{\phi}$ of MLPs with $s = \min_i n_i$ being the smallest width. Consider a parameterized affine function $G_{A,\mathbf{b}}$ solving

$$\min_{A,\mathbf{b}} \mathcal{L}(G_{A,\mathbf{b}}; \{(\mathbf{x}_i, y_i)\}_{i=1}^N), \qquad subject \ to \ rank(A) \le s,$$
(1)

for a continuous loss function \mathcal{L} . Then, for each local minimum, (A', \mathbf{b}') , of the above training problem, there exists a local minimum, ϕ' , of the MLP loss $\mathcal{L}(F_{\phi}; \{(\mathbf{x}_i, y_i)\}_{i=1}^N)$ with the property that $F_{\phi'}(\mathbf{x}_i) = G_{A',\mathbf{b}'}(\mathbf{x}_i)$ for i = 1, 2, ..., N.

Proof. See appendix A.2.

The proof of the above theorem constructs a network in which all activations of all training examples are positive. We do, however, expect that the general problem in expressivity occurs every time the support of the activations coincides for all training examples, as the latter reduces the deep network to an affine linear function (on the training set), which relates to the discussion in Balduzzi et al. (2017). We test this hypothesis below by initializing deep networks with biases of high variance. Note that the above constructions of Lemma 1 and Theorem 1 are not limited to MLPs and could be extended to convolutional neural networks with suitably restricted linear mappings G_{ϕ} by using the convolution filters to represent identities and avoiding any negative activations on the training examples. Moreover, shallower MLPs can similarly be embedded into deeper MLPs recursively. Linear classifiers, or even shallow MLPs, often have higher training loss than more expressive networks. Thus, we can use the idea of Theorem 1 to find various suboptimal local minima in the loss landscapes of neural networks. We confirm this with subsequent experiments.

We find that initializing a network at a point that approximately conforms to Theorem 1 is enough to get trapped in a bad local minimum. We verify this by training a linear classifier on CIFAR-10 with weight decay, (which has a test accuracy of 40.53%, loss of 1.57, and gradient norm of 0.00375). We then initialize a multilayer network as described in Lemma 1 to approximate this linear classifier and recompute these statistics on the full network (see Table 1). When training with this initialization, the gradient norm drops futher, moving parameters even closer to the linear minimizer. The final training result still yields positive activations for the entire training dataset.

Moreover, any isolated local minimum of a linear network results in many local minima of an MLP $F_{\phi'}$, as the weights ϕ' constructed in the proof of Theorem 1 can undergo transformations such as scaling, permutation, or even rotation without changing $F_{\phi'}$ as a function during inference, i.e. $F_{\phi'}(\mathbf{x}) = F_{\phi}(\mathbf{x})$ for all \mathbf{x} for an infinite set of parameters ϕ , as soon as F has at least one hidden layer.

While our first experiment initializes a deep MLP at a local minimum it inherited from a linear one to empirically illustrate our findings of Theorem 1, Table 1 also illustrates that similarly bad local minima are obtained when choosing large biases (third row) and choosing biases with large variance (fourth row) as conjectured above. To significantly reduce the bias, however, and still obtain a subpar optimum, we need to rerun the experiment with SGD without momentum, as shown in the last row, reflecting common intuition that momentum is helpful to move away from bad local optima.

In light of our finding that neural networks trained with unconventional initialization reach suboptimal local minima, we conclude that poor local minima can readily be found with a poor choice of hyperparameters. Suboptimal minima are less scarce than previously believed, and neural networks avoid these because good initializations and stochastic optimizers have been fine-tuned over time. Fortunately, promising theoretical directions may explain good optimization performance while remaining compatible with empirical observations. The approach followed by Du et al. (2019) analyzes the loss trajectory of SGD, showing that it avoids bad minima. While this work assumes (unrealistically) large network widths, this theoretical direction is compatible with empirical studies, such as Goodfellow et al. (2014), showing that the training trajectory of realistic deep networks does not encounter significant local minima.

| | At Initialization | | | After training | | |
|------------------------------------|-------------------|--------|---------|----------------|--------|---------|
| Init. Type | Loss | Grad. | Min. EV | Loss | Grad. | Min. EV |
| Default | 4.5963 | 0.5752 | -1.5549 | 0.0061 | 0.0074 | 0.0007 |
| Lemma 1 | 1.5702 | 0.0992 | 0.03125 | 1.5699 | 0.0414 | 0.0156 |
| Bias+20 | 31.204 | 343.99 | -1.7421 | 2.3301 | 0.0090 | 0.0005 |
| Bias $\in \mathcal{U}(-50, 50)$ | 51.445 | 378.36 | -430.49 | 2.3153 | 0.0048 | 0.0000 |
| $Bias \in \mathcal{U}(-10, 10)$ NM | 12.209 | 42.454 | -47.733 | 0.2198 | 0.0564 | 0.0013 |

Table 1: Local minima for MLPs generated via various initializations. We show loss, euclidean norm of the gradient vector, and minimum eigenvalue of the Hessian before and after training. We use 500 iterations of the power method on a shifted Hessian matrix computed on the full dataset to find the minimum eigenvalue. The experiment in the last row is trained with no momentum (NM).

3 WEIGHT DECAY: ARE SMALL ℓ_2 -NORM SOLUTIONS BETTER?

Classical learning theory advocates regularization for linear models, such as SVM and linear regression. For SVM, ℓ_2 regularization endows linear classifiers with a wide-margin property (Cortes & Vapnik, 1995), and recent work on neural networks has shown that minimum norm neural network interpolators benefit from over-parametrization (Hastie et al., 2019). Following the long history of explicit parameter norm regularization for linear models, weight decay is used for training nearly all high performance neural networks (He et al., 2015a; Chollet, 2016; Huang et al., 2017; Sandler et al., 2018).

In combination with weight decay, all of these cutting-edge architectures also employ batch normalization after convolutional layers (Ioffe & Szegedy, 2015). With that in mind, van Laarhoven (2017) shows that the regularizing effect of weight decay is counteracted by batch normalization, which removes the effect of shrinking weight matrices. Zhang et al. (2018) argue that the synergistic interaction between weight decay and batch norm arises because weight decay plays a large role in regulating the effective learning rate of networks, since scaling down the weights of convolutional layers amplifies the effect of each optimization step, effectively increasing the learning rate. Thus, weight decay increases the effective learning rate as the regularizer drags the parameters closer and closer towards the origin. The authors also suggest that data augmentation and carefully chosen learning rate schedules are more powerful than explicit regularizers like weight decay.

Other work echos this sentiment and claims that weight decay and dropout have little effect on performance, especially when using data augmentation (Hernández-García & König, 2018). Hoffer et al. (2018) further study the relationship between weight decay and batch normalization, and they develop normalization with respect to other norms. Shah et al. (2018) instead suggest that minimum norm solutions may not generalize well in the over-parametrized setting.

We find that the difference between performance of standard network architectures with and without weight decay is often statistically significant, even with a high level of data augmentation, for example, horizontal flips and random crops on CIFAR-10 (see Tables 2 and 3). But is weight decay the most effective form of ℓ_2 regularization? Furthermore, is the positive effect of weight decay because the regularizer promotes small norm solutions? We generalize weight decay by biasing the ℓ_2 norm of the weight vector towards other values using the following regularizer, which we call *norm-bias*:

$$R_{\mu}(\phi) = \left| \left(\sum_{i=1}^{P} \phi_i^2 \right) - \mu^2 \right|.$$
⁽²⁾

 R_0 is equivalent to weight decay, but we find that we can further improve performance by biasing the weights towards higher norms (see Tables 2 and 3). In our experiments on CIFAR-10, networks are trained using weight decay coefficients from their respective original papers. ResNet-18 and DenseNet are trained with $\mu^2 = 2500$ and norm-bias coefficient 0.005, and MobileNetV2 is trained with $\mu^2 = 5000$ and norm-bias coefficient 0.001. While we find that weight decay improves results over a non-regularized baseline for all three models, we also find that models trained with large norm bias (i.e., large μ) outperform models trained with weight decay. These results lend weight to the argument that explicit parameter norm regularization is in fact useful for training networks, even deep CNNs with batch normalization and data augmentation. However, the fact that norm-biased networks can outperform networks trained with weight decay suggests that any benefits of weight decay are unlikely to originate from the superiority of small-norm solutions.

To further investigate the effect of weight decay and parameter norm on generalization, we also consider models without batch norm. In this case, weight decay directly penalizes the norm of the linear operators inside a network, since there are no batch norm coefficients to compensate for the effect of shrinking weights. Or goal is to determine whether small-norm solutions are superior in this setting where the norm of the parameter vector is more meaningful.

In our first experiment without batch norm, we experience improved performance training an MLP with *norm-bias* (see Table 3). In a state-of-the-art setting, we consider ResNet-20 with Fixup initialization, a ResNet variant that removes batch norm and instead uses a sophisticated initialization that solves the exploding gradient problem (Zhang et al., 2019). We observe that weight decay substantially improves training over SGD with no explicit regularization — in fact, ResNets with this initialization scheme train quite poorly without explicit regularization and data normalization. Still, we find that *norm-bias* with $\mu^2 = 1000$ and norm-bias coefficient 0.0005 achieves better results than weight decay (see Table 3). This once again refutes the theory that small-norm parameters generalize better and brings into doubt any relationship between classical Tikhonov regularization and weight decay in neural networks. See Appendix A.5 for a discussion concerning the final parameter norms of Fixup networks.

Table 2: ResNet-18, DenseNet-40, and MobileNetV2 models trained on non-normalized CIFAR-10 data with various regularizers. Numerical entries represent the average accuracy over 10 runs.

| Model | No weight decay (%) | Weight decay (%) | Norm-bias (%) |
|-------------|---------------------|--------------------|---------------------------|
| ResNet | $93.46 (\pm 0.05)$ | $94.06(\pm 0.07)$ | 94.86 (± 0.05) |
| DenseNet | $89.26 (\pm 0.08)$ | $92.27 (\pm 0.06)$ | $92.49 (\pm 0.06)$ |
| MobileNetV2 | $92.88 (\pm 0.06)$ | $92.88 (\pm 0.09)$ | 93.50 (± 0.09) |

Table 3: ResNet-18, DenseNet-40, MobileNetV2, and ResNet-20 with Fixup initialization trained on normalized CIFAR-10 data with various regularizers. Numerical entries represent the average accuracy over 10 runs.

| Model | No weight decay (%) | Weight decay (%) | Norm-bias (%) |
|--------------|---------------------|--------------------|---------------------------|
| ResNet | $93.40(\pm 0.04)$ | $94.76(\pm 0.03)$ | $94.99(\pm 0.05)$ |
| DenseNet | $90.78 (\pm 0.08)$ | $92.26 (\pm 0.06)$ | 92.46 (±0.04) |
| MobileNetV2 | $92.84 (\pm 0.05)$ | $93.64 (\pm 0.05)$ | $93.64 (\pm 0.03)$ |
| ResNet Fixup | $10.00 (\pm 0.00)$ | $91.42 (\pm 0.04)$ | $91.55 (\pm 0.07)$ |
| MLP | $58.88 (\pm 0.10)$ | $58.95 (\pm 0.07)$ | 59.13 (± 0.09) |

4 KERNEL THEORY AND THE INFINITE-WIDTH LIMIT

In light of the recent surge of works discussing the properties of neural networks in the infinitewidth limit, in particular, connections between infinite-width deep neural networks and Gaussian processes, see Lee et al. (2017), several interesting theoretical works have appeared. The wide network limit and Gaussian process interpretations have inspired work on the neural tangent kernel (Jacot et al., 2018), while Lee et al. (2019) and Bietti et al. (2018) have used wide network assumptions to analyze the training dynamics of deep networks. The connection of deep neural networks to kernel-based learning theory seems promising, but how closely do current architectures match the predictions made for simple networks in the large-width limit?

We focus on the Neural Tangent Kernel (NTK), developed in Jacot et al. (2018). Theory dictates that, in the wide-network limit, the neural tangent kernel remains nearly constant as a network trains. Furthermore, neural network training dynamics can be described as gradient descent on a convex functional, provided the NTK remains nearly constant during training (Lee et al., 2019). In this section, we experimentally test the validity of these theoretical assumptions.



Figure 1: The correlation coefficient of the neural tangent kernel after training with its initialization. We expect this coefficient to converge toward 1 in the infinite-width limit for multi-layer networks as in Jacot et al. (2018), the effect on different architectures and especially residual neural networks is much less clear. We find that the trend toward infinite-width is essentially confirmed for the convolutional neural network, while it is falsified for the ResNet (which features batch normalization).

Fixing a network architecture, we use \mathcal{F} to denote the function space parametrized by $\phi \in \mathbb{R}^p$. For the mapping $F : \mathbb{R}^P \to \mathcal{F}$, the NTK is defined by

$$\Phi(\phi) = \sum_{p=1}^{P} \partial_{\phi_p} F(\phi) \otimes \partial_{\phi_p} F(\phi), \tag{3}$$

where the derivatives $\partial_{\phi_p} F(\phi)$ are evaluated at a particular choice of ϕ describing a neural network. The NTK can be thought of as a similarity measure between images; given any two images as input, the NTK returns an $n \times n$ matrix, where n is the dimensionality of the feature embedding of the neural network. We sample entries from the NTK by drawing a set of N images $\{x_i\}$ from a dataset, and computing the entries in the NTK corresponding to all pairs of images in our image set. We do this for a random neural network $f : \mathbb{R}^m \to \mathbb{R}^n$ and computing the tensor $\Phi(\phi) \in \mathbb{R}^{N \times N \times n \times n}$ of all pairwise realizations, restricted to the given data:

$$\Phi(\phi)_{ijkl} = \sum_{p=1}^{P} \partial_{\phi_p} f(\mathbf{x}_i, \phi)_k \cdot \partial_{\phi_p} f(\mathbf{x}_j, \phi)_l$$
(4)

By evaluating Equation 4 using automatic differentiation, we compute slices from the NTK before and after training for a large range of architectures and network widths. We consider image classification on CIFAR-10 and compare a two-layer MLP, a four-layer MLP, a simple 5-layer ConvNet, and a ResNet. We draw 25 random images from CIFAR-10 to sample the NTK before and after training. We measure the change in the NTK by computing the correlation coefficient of the (vectorized) NTK before and after training. We do this for many network widths, and see what happens in the wide network limit. For MLPs we increase the width of the hidden layers, for the ConvNet (6-Layer, Convolutions, ReLU, MaxPooling), we increase the number of convolutional filters, for the ResNet we consider the WideResnet (Zagoruyko & Komodakis, 2016) architecture, where we increase its width parameter. We initialize all models with uniform He initialization as discussed in He et al. (2015b), departing from specific Gaussian initializations in theoretical works to analyze the effects for modern architectures and methodologies.

The results are visualized in Figure 1, where we plot parameters of the NTK for these different architectures, showing how the number of parameters impacts the relative change in the NTK $(||\Phi_1 - \Phi_0||/||\Phi_0||)$, where Φ_0/Φ_1 denotes the sub-sampled NTK before/after training) and correlation coefficient ($Cov(\Phi_1, \Phi_0)/\sigma(\Phi_1)/\sigma(\Phi_0)$). Jacot et al. (2018) predicts that the NTK should change very little during training in the infinite-width limit.

At first glance, it might seem that these expectations are hardly met for our (non-infinite) experiments. Figure 1a shows that the relative change in the NTK during training (and also the magnitude of the NTK) is rapidly increasing with width and remains large in magnitude for a whole range of widths of convolutional architectures. The MLP architectures do show a trend toward small changes in the NTK, yet convergence to zero is slower in the 4-Layer case than in the 2-Layer case. However, a closer look shows that almost all of the relative change in the NTK seen in Figure 1a is explained by a simple linear re-scaling of the NTK. It should be noted that the scaling of the NTK is strongly effected by the magnitude of parameters at initialization. Within the NTK theory of Lee et al. (2017), a linear rescaling of the NTK during training corresponds simply to a change in learning rate, and so it makes more sense to measure similarity using a scale-invariant metric.

Measuring similarity between sub-sampled NTKs using the scale-invariant correlation coefficient, as in Figure 1b, is more promising. Surprisingly, we find that, as predicted in Jacot et al. (2018), the NTK changes very little (beyond a linear rescaling) for the wide ConvNet architectures. For the dense networks, the predicted trend toward small changes in the NTK also holds for most of the evaluated widths, although there is a dropoff at the end which may be an artifact of the difficulty of training these wide networks on CIFAR-10. For the Wide Residual Neural Networks, however, the general trend toward higher correlation in the wide network limit is completely reversed. The correlation coefficient decreases as network width increases, suggesting that the neural tangent kernel at initialization and after training becomes qualitatively more different as network width increases.

In summary, while we find that trends towards stability of the NTK hold suprisingly well for some realistic architectures in the wide network limit, the NTK is frequently unstable in standard parameter regimes and for network architectures that work extremely well in practice. This gives us hope that kernel-based theories of neural networks may yield guarantees for realistic (albeit wide) models. However, the good behavior of models with unstable NTKs is an indicator that good optimization and generalization behavior does not fundamentally hinge on the stability of the NTK.

5 RANK: DO NETWORKS WITH LOW-RANK LAYERS GENERALIZE BETTER?

State-of-the-art neural networks are highly over-parameterized, and their large number of parameters is a problem both for learning theory and for practical use. In the theoretical setting, rank has been used to tighten bounds on the generalization gap of neural networks. Generalization bounds from Harvey et al. (2017) are improved under conditions of low rank and high sparsity (Neyshabur et al., 2017) of parameter matrices, and the compressibility of low-rank matrices (and other lowdimensional structure) can be directly exploited to provide bounds (Arora et al., 2018). Further studies show a tendency of stochastic gradient methods to find low-rank solutions (Ji & Telgarsky, 2018). The tendency of SGD to find low-rank operators, in conjunction with results showing generalization bounds for low-rank operators, might suggest that the low-rank nature of these operators is important for generalization.

Langenberg et al. (2019) claim that low-rank networks, in addition to generalizing well to test data, are more robust to adversarial attacks. Theoretical and empirical results from the aforementioned paper lead the authors to make two major claims. First, the authors claim that networks which undergo adversarial training have low-rank and sparse matrices. Second, they claim that networks with low-rank and sparse parameter matrices are more robust to adversarial attacks. We find in our experiments that neither claim holds up in the standard robustness setting of CIFAR-10 image classification with ResNet-18.

We test the generalization and robustness properties of neural networks with low-rank and highrank operators by promoting low-rank or high-rank parameter matrices in late epochs. We employ the regularizer introduced in Sedghi et al. (2018) to create the protocols RankMin, to find low-rank parameters, and RankMax, to find high-rank parameters. RankMin involves fine-tuning a pre-trained model by replacing linear operators with their low-rank approximations, retraining, and repeating this process. Similarly, RankMax involves fine-tuning a pre-trained model by clipping singular values from the SVD of parameter matrices in order to find high-rank approximations. We are able to manipulate the rank of matrices without strongly affecting the performance of the network. We use both natural training and 7-step projected gradient descent (PGD) adversarial training routines (Madry et al., 2017). The goal of the experiment is to observe how the rank of weight matrices impacts generalization and robustness. We start by attacking naturally trained models with the standard PGD adversarial attack with $\epsilon = 8/255$. Then, we move to the adversarial training setting and test the effect of manipulating rank on generalization and on robustness.

In order to compare our results with Langenberg et al. (2019), we borrow the notion of effective rank, denoted by r(W) for some matrix W. This continuous relaxation of rank is defined as follows.

| Training method | Training | Clean | Robust (%) | Robust (%) |
|-----------------------|--------------|--------------|--------------------|--------------------|
| | Accuracy (%) | Accuracy (%) | $\epsilon = 8/255$ | $\epsilon = 1/255$ |
| Naturally Trained | 99.99 | 94.66 | 0.00 | 31.98 |
| RankMax | 99.76 | 93.66 | 0.00 | 22.01 |
| RankMin | 99.90 | 94.44 | 0.00 | 31.53 |
| Adversarially Trained | 89.66 | 85.06 | 33.10 | 80.49 |
| RankMaxAdv | 89.21 | 84.72 | 32.40 | 80.11 |
| RankMinAdv | 88.33 | 84.68 | 30.57 | 79.98 |

Table 4: All models tested are ResNet-18s. Robust accuracy is measured with 20-step PGD attacks with the ϵ values specified at the top of the column.



(a) Effective rank of naturally trained models.

(b) Effective rank of adversarially trained models.

Figure 2: This plot shows the effective rank of each filter. The filters are indexed on the x-axis, so moving to the right is like moving through the layers of the network. Our routines designed to manipulate the rank have exactly the desired effect as shown here.

 $r(W) = \frac{\|W\|_*}{\|W\|_F}$ where $\|\cdot\|_*, \|\cdot\|_1$, and $\|\cdot\|_F$ are the nuclear norm, the 1-norm, and the Frobenius norm, respectively. Note that the singular values of convolution operators can be found quickly with a method from Sedghi et al. (2018), and this method is used here.

In Table 4, RankMin and RankMax achieve similar generalization. Figure 2 confirms that these two training routines do, in fact, control effective rank. Thus, increasing rank using an analogue of rank minimizing algorithms does not harm performance. Moreover, we observe that adversarial robustness does not imply low-rank operators, nor do low-rank operators imply robustness. The findings in Ji & Telgarsky (2018) are corroborated here as the black dots in Figures 2 show that initializations are higher in rank than trained models. Our investigation into what useful intuition in practical cases can be gained from the theoretical work on the rank of CNNs and from the claims about adversarial robustness revealed that rank plays little to no role in the performance of residual networks in the setting of image classification.

6 CONCLUSION

Theoretical work on neural networks has made significant progress in recent years, and there are great reasons to be optimistic about its future. While there is a divide between theory and practice in deep learning, we have seen major benefits when theory and practice come together. For example, certifiably robust classifiers may allow users to ensure security against adversarial attacks (Raghunathan et al., 2018; Cohen et al., 2019), and Wasserstein GANs avoid many of the notorious training difficulties of generative adversarial networks (Arjovsky et al., 2017). This work is meant to underscore the need to carefully examine the assumptions of theory and also to highlight ways to steer theory towards more realistic and less understood models, like residual networks, deep networks, non-linear nets, and models that live far away from the wide-network/kernel limit.

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A APPENDIX

A.1 PROOF OF LEMMA 1

Lemma 1. Consider a family of L-layer multilayer perceptrons with ReLU activations $\{F_{\phi} : \mathbb{R}^m \to \mathbb{R}^n\}$ and let $s = \min_i n_i$ be the minimum layer width. Then this family has rank-s affine expression.

Proof. Let G be a rank-s affine function, and $\Omega \subset \mathbb{R}^m$ be a finite set. Let $G(\mathbf{x}) = A\mathbf{x} + \mathbf{b}$ with $A = U\Sigma V$ being the singular value decomposition of A with $U \in \mathbb{R}^{n \times s}$ and $V \in \mathbb{R}^{s \times m}$.

We define

$$A_1 = \begin{bmatrix} \Sigma V \\ \mathbf{0} \end{bmatrix}$$

where **0** is a (possibly void) $(n_1 - s) \times m$ matrix of all zeros, and $b_1 = c\mathbf{1}$ for $c = \max_{\mathbf{x}_i \in \Omega, 1 \le j \le n_1} |(A_1\mathbf{x}_i)_j| + 1$ and $\mathbf{1} \in \mathbb{R}^{n_1}$ being a vector of all ones. We further choose $A_l \in \mathbb{R}^{n_l \times n_{l-1}}$ to have an $s \times s$ identity matrix in the upper left, and fill all other entries with

zeros. This choice is possible since $n_l \ge s$ for all l. We define $\mathbf{b}_l = \begin{bmatrix} \mathbf{0} & c \mathbf{1} \end{bmatrix}^T \in \mathbb{R}^{n_l}$ where $\mathbf{0} \in \mathbb{R}^{1 \times s}$ is a vector of all zeros and $\mathbf{1} \in \mathbb{R}^{1 \times (n_l - s)}$ is a (possibly void) vector of all ones.

Finally, we choose $A_L = \begin{bmatrix} U & \mathbf{0} \end{bmatrix}$, where now **0** is a (possibly void) $n \times (n_{L-1} - s)$ matrix of all zeros, and $\mathbf{b}_L = -cA_L \mathbf{1} + \mathbf{b}$ for $\mathbf{1} \in \mathbb{R}^{n_{L-1}}$ being a vector of all ones.

Then one readily checks that $F_{\phi}(\mathbf{x}) = G(\mathbf{x})$ holds for all $x \in \Omega$. Note that all entries of all activations are greater or equal to c > 0, such that no ReLU ever maps an entry to zero.

A.2 PROOF OF THEOREM 1

Theorem 1. Consider a training set, $\{(\mathbf{x}_i, y_i)\}_{i=1}^N$, a family $\{F_{\phi}\}$ of MLPs with $s = \min_i n_i$ being the smallest width. Consider the training of a rank-s linear classifier $G_{A,\mathbf{b}}$, i.e.,

$$\min_{A,\mathbf{b}} \mathcal{L}(G_{A,\mathbf{b}}; \{(\mathbf{x}_i, y_i)\}_{i=1}^N), \qquad subject \ to \ rank(A) \le s, \tag{5}$$

for any continuous loss function \mathcal{L} . Then for each local minimum, (A', \mathbf{b}') , of the above training problem, there exists a local minimum, ϕ' , of $\mathcal{L}(F_{\phi}; \{(\mathbf{x}_i, y_i)\}_{i=1}^N)$ with the property that $F_{\phi'}(\mathbf{x}_i) = G_{A',\mathbf{b}'}(\mathbf{x}_i)$ for i = 1, 2, ..., N.

Proof. Based on the definition of a local minimium, there exists an open ball D around (A', \mathbf{b}') such that

$$\mathcal{L}(G_{A',\mathbf{b}'};\{(\mathbf{x}_i,y_i)\}_{i=1}^N) \le \mathcal{L}(G_{A,\mathbf{b}};\{(\mathbf{x}_i,y_i)\}_{i=1}^N) \quad \forall (A,\mathbf{b}) \in D \text{ with } \operatorname{rank}(A) \le s.$$
(6)

First, we use the same construction as in the proof of Lemma 1 to find a function $F_{\phi'}$ with $F_{\phi'}(\mathbf{x}_i) = G_{A',\mathbf{b}'}(\mathbf{x}_i)$ for all training example \mathbf{x}_i . Because the mapping $\phi \mapsto F_{\phi}(\mathbf{x}_i)$ is continuous (not only for the entire network F but also for all subnetworks), and because all activations of $F_{\phi'}$ are greater or equal to c > 0, there exists an open ball $B(\phi', \delta_1)$ around ϕ' such that the activations of F_{ϕ} remain positive for all \mathbf{x}_i and all $\phi \in B(\phi', \delta_1)$.

Consequently, the restriction of F_{ϕ} to the training set remains affine linear for $\phi \in B(\phi', \delta_1)$. In other words, for any $\phi \in B(\phi', \delta_1)$ we can write

$$F_{\phi}(\mathbf{x}_i) = A(\phi)\mathbf{x}_i + \mathbf{b}(\phi) \qquad \forall \mathbf{x}_i$$

by defining $A(\phi) = A_L A_{L-1} \dots A_1$ and $\mathbf{b}(\phi) = \sum_{l=1}^{L} A_L A_{L-1} \dots A_{l+1} \mathbf{b}_l$. Note that due to $s = \min_i n_i$, the resulting $A(\phi)$ satisfies rank $(A(\phi)) \leq s$.

After restricting ϕ to an open ball $B(\phi', \delta_2)$, for $\delta_2 \leq \delta_1$ sufficiently small, the above $(A(\phi), \mathbf{b}(\phi))$ satisfy $(A(\phi), \mathbf{b}(\phi)) \in D$ for all $\phi \in B(\phi', \delta_2)$. On this set, we, however, already know that the loss can only be greater or equal to $\mathcal{L}(F_{\phi'}; \{(\mathbf{x}_i, y_i)\}_{i=1}^N)$ due to equation 6. Thus, ϕ' is a local minimum of the underlying loss function.

A.3 ADDITIONAL COMMENTS REGARDING THEOREM 1

Note that our theoretical and experimental results do not contradict theoretical guarantees for deep linear networks (Kawaguchi, 2016; Laurent & Brecht, 2018) which show that all local minima are global. A deep linear network with $s = \min(n, m)$ is equivalent to a linear classifier, and in this case, the local minima constructed by Theorem 1 are global. However, this observation shows that Theorem 1 characterizes the gap between deep linear and deep nonlinear networks; the global minima predicted by linear network theories are inherited as (usually suboptimal) local minima when ReLU's are added. Thus, linear networks do not accurately describe the distribution of minima in non-linear networks.

A.4 ADDITIONAL RESULTS FOR SUBOPTIMAL LOCAL OPTIMA

Table 5 adds more experiments, repeating the previous experiment (training on CIFAR-10 until convergence from different initializations) for a full ResNet-18 architecture trained with gradient descent. We find that essentially the same results appear for the deeper architecture, initializing with very high bias leads to highly non-optimal solutions. In this case even solutions that are equally bad as a zero-norm initialization.

| | At Initi | alization | After t | raining |
|---------------------------------|----------|-----------|----------|----------|
| Init. Type | Loss | Grad. | Loss | Grad. |
| Default | 2.30312 | 0.0500 | 0.00014 | 0.0141 |
| Zero | 2.30258 | 0.000245 | 2.302585 | 0.000129 |
| Bias $\in \mathcal{U}(-10, 10)$ | 12.9679 | 214.686 | 2.3026 | 0.00123 |
| $Bias \in \mathcal{U}(-50, 50)$ | 84.678 | 1190.235 | 2.3026 | 0.007021 |

Table 5: Local minima for ResNets generated via initialization and trained by vanilla gradient descent, showing loss, euclidean norm of the gradient vector, and minimum eigenvalue of the Hessian before and after training.

A.5 DETAILS CONCERNING LOW-NORM REGULARIZATION EXPERIMENTS

Our experiments comparing regularizers all run for 300 epochs with an initial learning rate of 0.1 and decreases by a factor of 10 at epochs 100, 175, 225, and 275. We use the SGD optimizer with momentum 0.9. In error bounds of form $(\pm \sigma \%)$, σ represents the standard error, i.e. $\frac{\sigma_{sample}}{\sqrt{n}}$.

We also tried negative weight decay coefficients, which leads to ResNet-18 CIFAR-10 performance above 90% while blowing up parameter norm, but this performance is still suboptimal and is not informative concerning the optimality of minimum norm solutions. One might wonder if high norm-bias coefficients lead to even lower parameter norm than low weight decay coefficients. This question may not be meaningful in the case of networks with batch normalization. In the case of ResNet-20 with Fixup, which does not contain running mean and standard deviation, the average parameter ℓ_2 norm after training with weight decay is 24.51 while that of models trained with norm – bias is 31.62.

A.6 DETAILS ON THE NEURAL TANGENT KERNEL EXPERIMENT

For further reference, we include details on the NTK sampling during training epochs in Figure 3. We see that the parameter norm (Right) behaves normally (all of these experiments are trained with a standard weight decay parameter of 0.0005), yet the NTK norm (Left) rapidly increases. Most of this increase, however is scaling of the kernel, as the correlation plot (Middle) is much less drastic. We do see that most change happens in the very first epochs of training, whereas the kernel only changes slowly later on.



Figure 3: Plotting the evolution of NTK parameters during training epochs. Left: Norm of the NTK Tensor, Middle: Correlation of current NTK iterate versus initial NTK. Right: Reference plot of the network parameter norms.

A.7 DETAILS ON RANKMIN AND RANKMAX

We developed routines to promote both low-rank and high-rank parameter matrices. We did this by computing approximations to the linear operators at each layer. Since convolutional layers are linear operations, we know that there is a matrix whose dimensions are the number of parameters in the input to the convolution and the number of parameters in the output of the convolution. In order to compute low-rank approximations of these operators, one could write down the matrix corresponding to the convolution, and then compute a low-rank approximation using a singular value decomposition (SVD). In order to make this problem computationally tractable we used the method for computing singular values of convolution operators derived in Sedghi et al. (2018). We



Figure 4: The similarity coefficient of the neural tangent kernel after training with its initialization. We expect this coefficient to converge toward 1 in the infinite-width limit for multi-layer networks. Also shown is the direct relative difference of the NTK norms, which behaves similarly to the normalized direct difference from figure 1.



Figure 5: For reference we record the test accuracy of all models from 1 in the left plot and the relative change in parameters in the right plot.

were then able to do low-rank approximation in the classical sense, by setting each singular value below some threshold to zero. In order to compute high-rank operators, we clipped the singular values so that when mulitplying the SVD factors, we set each singular value to be equal to the minimum of some chosen constant and the true singular value. It is important to note here that these approximations to the convolutional layers, when done naively, can return convolutions with larger filters. To be precise, an $n \times n$ filter will map to a $k \times k$ filter through our rank modifications, where $k \ge n$. We follow the method in Sedghi et al. (2018), where these filters are pruned back down by only using $n \times n$ entries in the output.

The training routine was then modified in the last 20 epochs to periodically approximate the parameters of the network with one of our two approximations. In each case we use pre-trained ResNet-18 models trained for 200 epochs on CIFAR-10 data, with the learning rate initiated to 0.1 and decreasing by a factor of 10 at epochs 100 and 150. The data augmentation during training includes random crops and horizontal flips. For adversarial training we use ℓ_{∞} 7-step PGD with step size equal to 2/255, and total $\epsilon = 8/255$. Starting with one naturally and one adversarially pre-trained model, we trained for an additional 20 epochs with the learning rate decreasing by a factor of 10 again at epochs 203 and 208. For any run in which we project to low-rank or high-rank operators, these projections occur after epochs 201, 205, and 210. As specified in Table 4, we test the accuracy of each model on clean CIFAR-10 test data, as well as 20-step PGD attack with $\epsilon = 8/255$ (with step size equal to 2/255) and $\epsilon = 1/255$ (with step size equal to .25/255).