Theoretical Issues in Deep Networks: Approximation, Optimization and Generalization

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While deep learning is successful in a number of applications, it is not yet well understood theoretically. A satisfactory theoretical characterization of deep learning however, is beginning to emerge. It covers the following questions: 1) representation power of deep networks 2) optimization of the empirical risk 3) generalization properties of gradient descent techniques — why the expected error does not suffer, despite the absence of explicit regularization, when the networks are overparametrized? In this review we discuss recent advances in the three areas. In approximation theory both shallow and deep networks have been shown to approximate any continuous functions on a bounded domain at the expense of an exponential number of parameters (exponential in the dimensionality of the function). However, for a subset of compositional functions, deep networks of the convolutional type (even without weight sharing) can have a linear dependence on dimensionality, unlike shallow networks. In optimization we discuss the loss landscape for the exponential loss function. It turns out that global minima at infinity are completely degenerate. The other critical points of the gradient are less degenerate, with at least one — and typically more — nonzero eigenvalues. This suggests that stochastic gradient descent will find with high probability the global minima. To address the question of generalization for classification tasks, we use classical uniform convergence results to justify minimizing a surrogate exponential-type loss function under a unit norm constraint on the weight matrix at each layer — since the interesting variables for classification are the weight directions rather than the weights. As a side remark, such minimization for (homogeneous) ReLU deep networks implies maximization of the margin. The resulting constrained gradient system turns out to be identical to the well-known weight normalization technique, originally motivated from a rather different way. We also show that standard gradient descent contains an implicit $L_2$ unit norm constraint in the sense that it solves the same constrained minimization problem with the same critical points (but a different dynamics). Our approach, which is supported by several independent new results (1–4), offers a solution to the puzzle about generalization performance of deep overparametrized ReLU networks, uncovering the origin of the underlying hidden complexity control in the case of deep networks.

Machine Learning | Deep learning | Approximation | Optimization | Generalization

1. Introduction

In the last few years, deep learning has been tremendously successful in many important applications of machine learning. However, our theoretical understanding of deep learning, and thus the ability of developing principled improvements, has lagged behind. A satisfactory theoretical characterization of deep learning is emerging. It covers the following areas: 1) approximation properties of deep networks 2) optimization of the empirical risk 3) generalization properties of gradient descent techniques — why the expected error does not suffer, despite the absence of explicit regularization, when the networks are overparametrized?

A. When Can Deep Networks Avoid the Curse of Dimensionality? We start with the first set of questions, summarizing results in (5–7), and (8, 9). The main result is that deep networks have the theoretical guarantee, which shallow networks do not have, that they can avoid the curse of dimensionality for an important class of problems, corresponding to compositional functions, that is functions of functions. An especially interesting subset of such compositional functions are hierarchically local compositional functions where all the constituent functions are local in the sense of bounded small dimensionality. The deep networks that can approximate them without the curse of dimensionality are of the deep convolutional type — though, importantly, weight sharing is not necessary.

Implications of the theorems likely to be relevant in practice are:

a) Deep convolutional architectures have the theoretical guarantee that they can be much better than one layer architectures such as kernel machines for certain classes of problems; b) the problems for which certain deep networks are guaranteed to avoid the curse of dimensionality (see for a nice review (10)) correspond to input-output mappings that are com-

Significance Statement

In the last few years, deep learning has been tremendously successful in many important applications of machine learning. However, our theoretical understanding of deep learning, and thus the ability of developing principled improvements, has lagged behind. A theoretical characterization of deep learning is now beginning to emerge. It covers the following questions: 1) representation power of deep networks 2) optimization of the empirical risk 3) generalization properties of gradient descent techniques — how can deep networks generalize despite being overparametrized — more weights than training data — in the absence of any explicit regularization? We review progress on all three areas showing that 1) for a class of compositional functions deep networks of the convolutional type are exponentially better approximators than shallow networks; 2) only global minima are effectively found by stochastic gradient descent for over-parametrized networks; 3) there is a hidden norm control in the minimization of cross-entropy by gradient descent that allows generalization despite overparametrization.

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of this paper is the \( \sup \) norm in keeping with the standard choice in approximation theory. As it turns out, the results of this section require the \( \sup \) norm in order to be independent from the unknown distribution of the input data.

Let \( V_N \) be the \( \ell \) be set of all networks of a given kind with \( N \) units (which we take to be or measure of the complexity of the approximant network). The \textit{degree of approximation} is defined by \( \text{dist}(f; V_N) = \inf_{P \in V_N} |f - P| \). For example, if \( \text{dist}(f; V_N) = \mathcal{O}(N^{-\gamma}) \) for some \( \gamma > 0 \), then a network with complexity \( N = \mathcal{O}(\epsilon^{-\frac{1}{2}}) \) will be sufficient to guarantee an approximation with accuracy at least \( \epsilon \). The only a priori information on the class of target functions \( f \), is codified by the statement that \( f \in W \) for some subspace \( W \subseteq X \). This subspace is a smoothness and compositional class, characterized by the parameters \( m \) and \( d \) (\( d = 2 \) in the example of Figure 1 ; it is the size of the kernel in a convolutional network).

D. Shallow and deep networks. This section characterizes conditions under which deep networks are “better” than shallow network in approximating functions. Thus we compare shallow (one-hidden layer) networks with deep networks as shown in Figure 1. Both types of networks use the same small set of operations – dot products, linear combinations, a fixed nonlinear function of one variable, possibly convolution and pooling. Each node in the networks corresponds to a node in the graph of the function to be approximated, as shown in the Figure. A unit is a neuron which computes \((x, w) + b_+ \), where \( w \) is the vector of weights on the vector input \( x \). Both \( w \) and the real number \( b \) are parameters tuned by learning. We assume here that each node in the networks computes the linear combination of \( r \) such units \( \sum_{i=1}^r c_i(x, w_i) + b_+ \). Notice that in our main example of a network corresponding to a function with a binary tree graph, the resulting architecture is an idealized version of deep convolutional neural networks described in the literature. In particular, it has only one output at the top unlike most of the deep architectures with many channels and many top-level outputs. Correspondingly, each node computes a single value instead of multiple channels, using the combination of several units. However our results hold also for these more complex networks (see (25)).

The sequence of results is as follows.

- **Both shallow (a) and deep (b) networks are universal,** that is they can approximate arbitrarily well any continuous function of \( n \) variables on a compact domain. The result for shallow networks is classical.

- We consider a special class of functions of \( n \) variables on a compact domain that are hierarchical compositions of local functions, such as \( f(x_1, \cdots, x_8) = h_3(h_21(h_11(x_1,x_2), h_12(x_3, x_4)), h_22(h_13(x_5, x_6), h_14(x_7,x_8))) \)

The structure of the function in Figure 1 b) is represented by a graph of the binary tree reflecting dimensionality \( d = 2 \) for the constituent functions \( h \). In general, \( d \) is arbitrary but fixed and independent of the dimensionality \( n \) of the compositional function \( f \). (25) formalizes the more general compositional case using directed acyclic graphs.

- The approximation of functions with a compositional structure – can be achieved with the same degree of accuracy by deep and shallow networks but the number of
parameters are much smaller for the deep networks than for the shallow network with equivalent approximation accuracy.

We approximate functions with networks in which the activation nonlinearity is a smoothed version of the so called ReLU, originally called ramp by Breiman and given by \( \sigma(x) = x_+ = \max(0, x) \). The architecture of the deep networks reflects the function graph with each node \( h_i \) being a ridge function, comprising one or more neurons.

Let \( I^n = [-1, 1]^n \), \( X = C(I^n) \) be the space of all continuous functions on \( I^n \), with \( ||f|| = \max_{x \in I^n} |f(x)| \). Let \( S_{N,n} \) denote the class of all shallow networks with \( N \) units of the form

\[
x \mapsto \sum_{k=1}^{N} a_k \sigma(\langle w_k, x \rangle + b_k),
\]

where \( w_k \in \mathbb{R}^n, b_k, a_k \in \mathbb{R} \). The number of trainable parameters here is \((n + 2)N \sim n \). Let \( m \geq 1 \) be an integer, and \( W^n_m \) be the set of all functions of \( n \) variables with continuous partial derivatives of orders up to \( m < \infty \) such that \( ||f|| + \sum_{1 \leq |k| \leq m} \|D^k f\| \leq 1 \), where \( D^k \) denotes the partial derivative indicated by the multi-integer \( k \geq 1 \), and \( |k|_1 \) is the sum of the components of \( k \).

For the hierarchical binary tree network, the analogous spaces are defined by considering the compact set \( W^n_{m,2} \) to be the class of all compositional functions \( f \) of \( n \) variables with a binary tree architecture and constituent functions \( h \) in \( W^n_2 \). We define the corresponding class of deep networks \( D_{N,2} \) to be the set of all deep networks with a binary tree architecture, where each of the constituent nodes is in \( S_{M,2} \), where \( N = |V|M, V \) being the set of non-leaf vertices of the tree. We note that in the case when \( n \) is an integer power of 2, the total number of parameters involved in a deep network in \( D_{N,2} \) is \( 4N \).

The first theorem is about shallow networks.

**Theorem 1** Let \( \sigma : \mathbb{R} \rightarrow \mathbb{R} \) be infinitely differentiable, and not a polynomial. For \( f \in W^n_m \) the complexity of shallow networks that provide accuracy at least \( \epsilon \) is \( N = \mathcal{O}(\epsilon^{-n/m}) \) and is the best possible. \[1\]

The estimate of Theorem 1 is the best possible if the only a priori information we are allowed to assume is that the target function belongs to \( f \in W^n_m \). The exponential dependence on the dimension \( n \) of the number \( \epsilon^{-n/m} \) of parameters needed to obtain an accuracy \( \mathcal{O}(\epsilon) \) is known as the curse of dimensionality. Note that the constants involved in the theorems will depend upon the norms of the derivatives of \( f \) as well as \( \sigma \).

Our second and main theorem is about deep networks with smooth activations (preliminary versions appeared in (6–8)). We formulate it in the binary tree case for simplicity but it extends immediately to functions that are compositions of constituent functions of a fixed number of variables \( d \) (in convolutional networks \( d \) corresponds to the size of the kernel).

**Theorem 2** For \( f \in W^n_{m,2} \) consider a deep network with the same compositional architecture and with an activation function \( \sigma : \mathbb{R} \rightarrow \mathbb{R} \) which is infinitely differentiable, and not a polynomial. The complexity of the network to provide approximation with accuracy at least \( \epsilon \) is \( N = \mathcal{O}((n - 1)\epsilon^{-2/m}) \). \[2\]

The proof is in (25). The assumptions on \( \sigma \) in the theorems are not satisfied by the ReLU function \( x \mapsto x_+ \), but they are satisfied by smoothing the function in an arbitrarily small interval around the origin. The result of the theorem can be extended to non-smooth ReLU (25).

In summary, when the only a priori assumption on the target function is about the number of derivatives, then to guarantee an accuracy of \( \epsilon \), we need a shallow network with \( \mathcal{O}(\epsilon^{-n/m}) \) trainable parameters. If we assume a hierarchical structure on the target function as in Theorem 2, then the corresponding deep network yields a guaranteed accuracy of \( \epsilon \) with \( \mathcal{O}(\epsilon^{-2/m}) \) trainable parameters. Note that Theorem 2 applies to all \( f \) with a compositional architecture given by a graph which correspond to, or is a subgraph of, the graph associated with the deep network – in this case the graph corresponding to \( W^n_{m,2} \).

## 2. The Optimization Landscape of Deep Nets with Smooth Activation Function

The main question in optimization of deep networks is to the landscape of the empirical loss in terms of its global minima and local critical points of the gradient.

### A. Related work

There are many recent papers studying optimization in deep learning. For optimization we mention work based on the idea that noisy gradient descent (27–30) can find a global minimum. More recently, several authors studied the dynamics of gradient descent for deep networks with assumptions about the input distribution or on how the labels are generated. They obtain global convergence for some shallow neural networks (31–36). Some local convergence results have as well been proved (37–39). The most interesting such approach is (36), which focuses on minimizing the training loss and proving that randomly initialized gradient descent can achieve zero training loss (see also (40–42)). In summary, there is by now an extensive literature on optimization that formalizes and refines to different special cases and to the discrete domain our results of (43, 44).

### B. Degeneracy of global and local minima under the exponential loss

The first part of the argument of this section relies on the obvious fact (see (1)), that for RELU networks under the hypothesis of an exponential-type loss function, there are no local minima that separate the data – the only critical points of the gradient that separate the data are the global minima.

Notice that the global minima are at \( \rho = \infty \), when the exponential is zero. As a consequence, the Hessian is identically zero with all eigenvalues being zero. On the other hand any point of the loss at a finite \( \rho \) has nonzero Hessian: for instance in the linear case the Hessian is proportional to \( \sum_n x_n a_n^T \).

The local minima which are not global minima must misclassify. How degenerate are they?

Simple arguments (1) suggest that the critical points which are not global minima cannot be completely degenerate. We thus have the following

**Property 1** Under the exponential loss, global minima are completely degenerate with all eigenvalues of the Hessian (\( W \) of them with \( W \) being the number of parameters in the network) being zero. The other critical points of the gradient are less
are generically (that is with probability one) degenerate on a "solution" of the stochastic differential Langevin equation and deep network, SGD selects with high probability the global dimensionality degenerate; the other critical points – saddles and local minima – are a large number of global zero-error minimizers which are degenerate, with at least one – and typically $N$ – nonzero eigenvalues.

For the general case of non-exponential loss and smooth nonlinearities instead of the RELU the following conjecture has been proposed (1):

**Conjecture 1**: For appropriate overparametrization, there are a large number of global zero-error minimizers which are degenerate; the other critical points – saddles and local minima – are generically (that is with probability one) degenerate on a set of much lower dimensionality.

**C. SGD and Boltzmann Equation.** The second part of our argument (in (44)) is that SGD concentrates in probability on the most degenerate minima. The argument is based on the similarity between a Langevin equation and SGD and on the fact that the Boltzmann distribution is exactly the asymptotic “solution” of the stochastic differential Langevin equation and also of SGDL, defined as SGD with added white noise (see for instance (45)). The Boltzmann distribution is

$$p(f) = \frac{1}{Z} e^{-\frac{L(f)}{T}},$$

where $Z$ is a normalization constant, $L(f)$ is the loss and $T$ reflects the noise power. The equation implies that SGDL prefers degenerate minima relative to non-degenerate ones of the same depth. In addition, among two minimum basins of equal depth, the one with a larger volume is much more likely in high dimensions as shown by the simulations in (44). Taken together, these two facts suggest that SGD selects degenerate minimizers corresponding to larger isotropic flat regions of the loss. Then SDGL shows concentration – because of the high dimensionality – of its asymptotic distribution Equation 3.

Together (43) and (1) suggest the following

**Conjecture 2**: For appropriate overparametrization of the deep network, SGD selects with high probability the global minimizers of the empirical loss, which are highly degenerate.

### 3. Generalization

Recent results by (2) illuminate the apparent absence of “overfitting” (see Figure 4) in the special case of linear networks for binary classification. They prove that minimization of loss functions such as the logistic, the cross-entropy and the exponential loss yields asymptotic convergence to the maximum margin solution for linearly separable datasets, independently of the initial conditions and without explicit regularization. Here we discuss the case of nonlinear multilayer DNNs under exponential-type losses, for several variations of the basic gradient descent algorithm. The main results are:

- classical uniform confidence bounds for generalization suggest a form of complexity control on the dynamics of the weight directions $V_k$; minimize a surrogate loss subject to a unit $L_2$ norm constraint;
- gradient descent on the exponential loss with an explicit $L_2$ unit norm constraint is equivalent to a well-known gradient descent algorithms weight normalization which is closely related to batch normalization;
- unconstrained gradient descent on the exponential loss yields a dynamics with the same critical points as weight normalization: the dynamics implicitly respect a $L_2$ unit constraint on the directions of the weights $V_k$.

We observe that several of these results directly apply to kernel machines for the exponential loss under the separability/interpolation assumption, because kernel machines are one-homogeneous.

### A. Related work.

A number of papers have studied gradient descent for deep networks (46–48). Close to the approach summarized here (details are in (1)) is the paper (49). Its authors study generalization assuming a regularizer because they are – like us – interested in normalized margin. Unlike their assumption of an explicit regularization, we show here that commonly used techniques, such as weight and batch normalization, in fact minimize the surrogate loss margin while controlling the complexity of the classifier without the need to add a regularizer or to use weight decay. Surprisingly, we will show that even standard gradient descent on the weights implicitly controls the complexity through an “implicit” unit $L_2$ norm constraint. Two very recent papers ((4) and (3)) develop an elegant but complicated margin maximization based approach which lead to some of the same results of this section (and many more). The important question of which conditions are necessary for gradient descent to converge to the maximum of the margin of $\tilde{f}$ are studied by (4) and (3). Our approach does not need the notion of maximum margin but our theorem 3 establishes a connection with it and thus with the results of (4) and (3). Our main goal here (and in (1)) is to achieve a simple understanding of where the complexity control underlying generalization is hiding in the training of deep networks.

### B. Deep networks: definitions and properties.

We define a deep network with $K$ layers with the usual coordinate-wise scalar activation functions $\sigma(z) : R \rightarrow R$ as the set of functions $f(W;x) = \sigma(W^K \sigma(W^{K-1} \cdots \sigma(W^1 x)))$, where the input is $x \in \mathbb{R}^d$, the weights are given by the matrices $W^k$, one per layer, with matching dimensions. We sometime use the symbol $W$ as a shorthand for the set of $W^k$ matrices $k = 1, \cdots, K$. For simplicity we consider here the case of binary classification in which $f$ takes scalar values, implying that the last layer matrix $W^K$ is $W^K \in \mathbb{R}^{1 \times K_l}$. The labels are $y_n \in \{-1, 1\}$. The weights of hidden layer $l$ are collected in a matrix of size $h_l \times h_{l-1}$. There are no biases apart form the input layer where the bias is instantiated by one of the input dimensions being a constant. The activation function in this section is the ReLU activation.

For ReLU activations the following important positive one-homogeneity property holds $\sigma(z) = \frac{\partial \sigma(z)}{\partial z} z$. A consequence of one-homogeneity is a structural lemma (Lemma 2.1 of (50))
where
\[ f(x) \] is the expected loss, \( \hat{L}(f) \) is the empirical loss, \( \mathbb{R}_N(F) \) is the empirical Rademacher average of the class of functions \( F \), measuring its complexity; \( c_1, c_2 \) are constants that depend on properties of the Lipschitz constant of the loss function, and on the architecture of the network.

Thus minimizing under a constraint on the Rademacher complexity a surrogate function such as the cross-entropy (which becomes the logistic loss in the binary classification case) will minimize an upper bound on the expected classification error because such surrogate functions are upper bounds on the 0 – 1 function. We can choose a class of functions \( \hat{F} \) with normalized weights and write \( f(x) = \rho f(x) \) and \( \mathbb{R}_N(F) = \rho \mathbb{R}_N(\hat{F}) \). One can choose any fixed \( \rho \) as a (Ivanov) regularization-type tradeoff.

In summary, the problem of generalization may be approached by minimizing the exponential loss – more in general an exponential-type loss, such the logistic and the cross-entropy – under a unit norm constraint on the weights matrices, since we are interested in the directions of the weights:

\[ \lim_{\rho \to \infty} \min_{\|V_k\|=1} L(\rho \hat{f}) = \text{max}_{\|V_k\|\le 1} \eta(\tilde{f}) \]

where \( \rho \) is a (Ivanov) regularization-type tradeoff.

C. Uniform convergence bounds: minimizing a surrogate loss under norm constraint. Classic generalization bounds for regression (51) suggest that minimizing the empirical loss of a loss function such as the cross-entropy subject to constrained complexity of the minimizer is a way to attain generalization, that is an expected loss which is close to the empirical loss:

\[ L(f) \le \hat{L}(f) + c_1 \mathbb{R}_N(F) + c_2 \sqrt{\frac{\ln(\frac{1}{\delta})}{2N}} \]  

D. Minimization under unit norm constraint: weight normalization. The approach is then to minimize the loss function

\[ L(f(w)) = \sum_{n=1}^N e^{-f(V;x_n,y_n)} \]  

with \( \rho = \prod \rho_k \) subject to \( \|V_k\| = 1 \forall k \), that is under a unit norm constraint for the weight matrix at each layer (if \( p = 2 \) then \( \|V_k\|_F^2 = 1 \) is the Frobenius norm), since \( V_k \) are the directions of the weights which are the relevant quantity for classification. The minimization is understood as a sequence of minimizations for a sequence of increasing \( \rho_k \). Clearly these constraints imply the constraint on the norm of the product of weight matrices for any \( p \) norm (because any induced operator norm is a sub-multiplicative matrix norm). The standard choice for a loss function is an exponential-type loss such the cross-entropy, which for binary classification becomes the logistic function. We study here the exponential because it is simpler and retains all the basic properties.

There are several gradient descent techniques that given the unconstrained optimization problem transform it into a constrained gradient descent problem. To provide the background let us formulate the standard unconstrained gradient descent problem for the exponential loss as it is used in practical training of deep networks:

\[ W_{k}^{i,j} = -\frac{\partial L}{\partial W_{k}^{i,j}} = \sum_{n=1}^N y_n \frac{\partial f(x_n;W)}{\partial W_{k}^{i,j}} e^{-y_nf(x_n;W)} \]
where $W_k$ is the weight matrix of layer $k$. Notice that, since the structural property implies that at a critical point we have $\sum_{n=1}^{N} y_n f(x_n; w)e^{-y_n f(x_n; W)} = 0$, the only critical points of this dynamics that separate the data (i.e. $y_n f(x_n; w) > 0 \forall n$) are global minima at infinity. Of course for separable data, while the loss decreases asymptotically to zero, the norm of the weights $\rho_k$ increases to infinity, as we will see later. Equations 7 define a dynamical system in terms of the gradient of the exponential loss $L$.

The set of gradient-based algorithms enforcing a unit-norm constraints (53) comprises several techniques that are equivalent for small values of the step size. They are all good approximations of the true gradient method. One of them is the Lagrange multiplier method; another is the tangent gradient method based on the following theorem:

**Theorem 4 (53)** Let $||u||_p$ denote a vector norm that is differentiable with respect to the elements of $u$ and let $g(t)$ be any vector function with finite $L_2$ norm. Then, calling $\nu(t) = \frac{\partial ||u||_p}{\partial u} u(t)$, the equation

$$\dot{u} = h(t) = S g(t) = (I - \nu v^T ||v||^2)g(t) \quad [8]$$

with $||u(0)|| = 1$, describes the flow of a vector $u$ that satisfies $||u||_p = 1$ for all $t \geq 0$.

In particular, a form for $g$ is $g(t) = \mu(t) \nabla_u L$, the gradient update in a gradient descent algorithm. We call $S g(t)$ the the tangent gradient transformation of $g$. In the case of $p = 2$ we replace $\nu$ in Equation 8 with $u$ because $\nu(t) = \frac{\partial ||u||_2}{\partial u} = u$. This gives $S = I - \frac{u u^T}{||u||^2}$ and $\dot{u} = S g(t)$.

Consider now the empirical loss $L$ written in terms of $V_k$ and $\rho_k$ instead of $W_k$, using the change of variables defined by $W_k = \rho_k V_k$ but without imposing a unit norm constraint on $V_k$. The flows in $\rho_k, V_k$ can be computed as $\dot{\rho}_k = \frac{\partial L}{\partial \rho_k} = V_k^T \frac{\partial L}{\partial W_k}$ and $\dot{V}_k = \frac{\partial W_k}{\partial V_k} \frac{\partial L}{\partial W_k} = \rho_k \frac{\partial L}{\partial W_k}$ given by Equations 7.

We now enforce the unit norm constraint on $V_k$ by using the tangent gradient transform on the $V_k$ flow. This yields

$$\dot{\rho}_k = V_k^T \frac{\partial L}{\partial W_k} \dot{V}_k = S_k \rho_k \frac{\partial L}{\partial W_k} \quad [9]$$

Notice that the dynamics above follows from the classical approach of controlling the Rademacher complexity of $f$ during optimization (suggested by bounds such as Equation 4). The approach and the resulting dynamics for the directions of the weights may seem different from the standard unconstrained approach in training deep networks. It turns out, however, that the dynamics described by Equations 9 is the same dynamics of Weight Normalization.

The technique of Weight normalization (54) was originally proposed as a small improvement on standard gradient descent “to reduce covariate shifts”. It was defined for each layer in terms of $w = g(||w||)$, as

$$\dot{g} = \frac{v}{||v||} \frac{\partial L}{\partial w} = \frac{g}{||v||} S \frac{\partial L}{\partial w} \quad [10]$$

with $S = I - \frac{v v^T}{||v||^2}$. It is easy to see that Equations 9 are the same as the weight normalization Equations 10, if $||v|| = 1$. We now observe, multiplying Equation 9 by $v^T$, that $v^T \dot{v} = 0$ because $v^T S = 0$, implying that $||v||^2$ is constant in time with a constant that can be taken to be 1. Thus the two dynamics are the same.

**E. Generalization with hidden complexity control.** Empirically it appears that GD and SGD converge to solutions that can generalize even without batch or weight normalization. Convergence may be difficult for quite deep networks and generalization may not be as good as with batch normalization but it still occurs. How is this possible?

We study the dynamical system $W_k^{\nu^j}$ under the parametrization $W_k^{\nu^j} = \rho_k V_k^{\nu^j}$ with $||V_k^{\nu^j}|| = 1$. We consider for each weight matrix $W_k$ the corresponding “vectorized” representation in terms of vectors $W_k^{\nu^j} = W_k$. We use the following definitions and properties (for a vector $w$):

- Define $\bar{w} = \frac{w}{||w||_2} \bar{w}$; thus $w = ||w||_2 \bar{w}$ with $||\bar{w}||_2 = 1$. Also define $S = I - \bar{w} \bar{w}^T = I - \frac{w w^T}{||w||^2}$.
- The following relations are easy to check:
  1. $\frac{\partial ||w||_2}{\partial w} = \bar{w}$
  2. $\frac{\partial \bar{w}}{\partial w} = \frac{S}{||w||}$
  3. $Sw = S\bar{w} = 0$
  4. $S^2 = S$

The gradient descent dynamic system used in training deep networks for the exponential loss is given by Equation 7. Following the chain rule for the time derivatives, the dynamics for $W_k$ is exactly (see (1)) equivalent to the following dynamics for $||W_k|| = \rho_k$ and $V_k$:

$$\dot{\rho}_k = \frac{\partial ||W_k||}{\partial W_k} \dot{W}_k = V_k^T W_k \quad [11]$$

and

$$\dot{V}_k = \frac{\partial V_k}{\partial W_k} \frac{\partial L}{\partial W_k} = \frac{S_k}{\rho_k} W_k \quad [12]$$

where $S_k = I - V_k V_k^T$. We used property 1 in 4 for Equation 11 and property 2 for Equation 12.

The key point here is that the dynamics of $V_k$ includes a unit $L_2$ norm constraint: using the tangent gradient transform will not change the equation because $S^2 = S$.

As separate remarks, notice that if for $t > t_0, f$ separates all the data, $\frac{d}{dt} \rho_k > 0$, that is $\rho$ diverges to $\infty$ with $\lim_{t \to \infty} \dot{\rho} = 0$. In the 1-layer network case the dynamics yields $\rho \approx \log t$ asymptotically. For deeper networks, this is different. (1) shows (for one support vector) that the product of weights at each layer diverges faster than logarithmically, but each individual layer diverges slower than in the 1-layer case. The norm of the each layer grows at the same rate $\rho^2_k$, independent of $k$. The $V_k$ dynamics has stationary or critical points given by

$$\sum \alpha_n(\rho) t \left( \frac{\partial f(x_n)}{\partial V_k^{\nu^j}} - V_k^{\nu^j} f(x_n) \right), \quad [13]$$

where $\alpha_n = e^{-y_n \rho(f(x_n))}$. We examine later the linear one-layer case $f(x) = v^T x$ in which case the stationary points of the gradient are given by $\sum \alpha_n(\rho)(x_n - v v^T x_n)$ and of course coincide with the solutions obtained with Lagrange multipliers. In the general case the critical points correspond
for $\rho \to \infty$ to degenerate zero “asymptotic minima” of the loss.

To understand whether there exists an implicit complexity control in standard gradient descent of the weight directions, we check whether there exists an $L_p$ norm for which unconstrained normalization is equivalent to constrained normalization.

From Theorem 4 we expect the constrained case to be given by the action of the following projector onto the tangent space:

$$
S_p = I - \frac{\nu v^T}{\|v\|^2} \quad \text{with} \quad \nu_v = \frac{\partial \|w\|}{\partial w_i} = \text{sign}(w_i) \circ \left( \frac{|w_i|}{\|w\|_p} \right)^{p-1}.
$$

The constrained Gradient Descent is then

$$
\dot{\rho}_k = V_k^T W_k \quad \dot{V}_k = \rho_k S_p \dot{W}_k. \tag{14}
$$

On the other hand, reparametrization of the unconstrained dynamics in the $p$-norm gives (following Equations 11 and 12)

$$
\dot{\rho}_k = \frac{\partial \|W_k\|_p}{\partial W_k} \frac{\partial W_k}{\partial t} = \text{sign}(W_k) \circ \left( \frac{|W_k|}{\|W_k\|_p} \right)^{p-1} \cdot \dot{W}_k
$$

and

$$
\dot{V}_k = \frac{\partial V_k}{\partial W_k} \frac{\partial W_k}{\partial t} = \frac{I - \text{sign}(W_k) \circ \left( \frac{|W_k|}{\|W_k\|_p} \right)^{p-1} W_k^T}{\|W_k\|_p^{p-1}} \cdot \dot{W}_k. \tag{16}
$$

These two dynamical systems are clearly different for generic $p$ reflecting the presence or absence of a regularization-like constraint on the dynamics of $V_k$.

As we have seen however, for $p = 2$ the 1-layer dynamical system obtained by minimizing $L$ in $\rho_k$ and $V_k$ with $W_k = \rho_k V_k$ under the constraint $\|V_k\|_2 = 1$, is the weight normalization dynamics

$$
\dot{\rho}_k = V_k^T W_k \quad \dot{V}_k = S \rho_k \dot{W}_k, \tag{17}
$$

which is quite similar to the standard gradient equations

$$
\dot{\rho}_k = V_k^T W_k \quad \dot{v} = \frac{S \rho_k}{\rho_k} \dot{W}_k. \tag{18}
$$

The two dynamical systems differ only by a $\rho_k^2$ factor in the $\dot{V}_k$ equations. However, the critical points of the gradient for the $\dot{V}_k$ flow, that is the point for which $\dot{V}_k = 0$, are the same in both cases since for any $t > 0$ $\rho(t) > 0$ and thus $V_k = 0$ is equivalent to $S W_k = 0$. Hence, gradient descent with unit $L_2$-norm constraint is equivalent to the standard, unconstrained gradient descent but only when $p = 2$. Thus

**Fact 1** The standard dynamical system used in deep learning, defined by $\dot{W}_k = -\rho_k \frac{\partial L}{\partial W_k}$, implicitly respects a unit $L_2$ norm constraint on $V_k$ with $\rho_k V_k = W_k$. Thus, under an exponential loss, if the dynamics converges, the $V_k$ represent the minimizer under the $L_2$ unit norm constraint.

Thus standard GD implicitly enforces the $L_2$ norm constraint on $V_k = \frac{W_k}{\|W_k\|_2}$ consistently with Srebro’s results on implicit bias of GD. Other minimization techniques such as coordinate descent may be biased towards different norm constraints.

### F. Linear networks and rates of convergence

The linear $(f(x) = \rho v^T x)$ networks case (2) is an interesting example of our analysis in terms of $\rho$ and $v$ dynamics. We start with unconstrained gradient descent, that is with the dynamical system

$$
\dot{\rho} = \frac{1}{\rho} \sum_{n=1}^{N} e^{-\rho v^T x_n} v^T x_n \quad \dot{v} = \frac{1}{\rho} \sum_{n=1}^{N} e^{-\rho v^T x_n} (x_n - v^T x_n). \tag{19}
$$

If gradient descent in $v$ converges to $\dot{v} = 0$ at finite time, $v$ satisfies $v^T x = x$, where $x = \sum_{j \neq k} a_j x_j$ with positive coefficients $a_j$ and $x_j$ are the $C$ support vectors (see (1)). A solution $v^T = \|x\| x$ then exists (1), the pseudoinverse of $x$, since $x$ is a vector, is given by $x^T = \frac{x^T}{\|x\|^2}$. On the other hand, the operator $T$ in $v(t + 1) = T(v(t))$ associated with equation 19 is non-expanding, because $\|v\| = 1$, $\forall t$. Thus there is a fixed point $v \propto x$ which is independent of initial conditions (56) and unique (in the linear case).

The rates of convergence of the solutions $\rho(t)$ and $v(t)$, derived in different way in (2), may be read out from the equations for $\rho$ and $v$. It is easy to check that a general solution for $\rho$ is of the form $\rho \propto C \log t$. A similar estimate for the exponential term gives $e^{-\rho v^T x_n} \propto \frac{1}{\rho}$. Assume for simplicity a single support vector $x$. We claim that a solution for the error $\epsilon = v - x$, since $v$ converges to $x$, behaves as $\frac{1}{\rho \log t}$. In fact we write $v = x + \epsilon$ and plug it in the equation for $v$ in 20. We obtain (assuming normalized input $\|x\| = 1$)

$$
\dot{\epsilon} = \frac{1}{\rho} e^{-\rho v^T x} (x - (x + \epsilon) (x + \epsilon)^T) \propto \frac{1}{\rho} e^{-\rho v^T x} (x - x - x \epsilon^T - x \epsilon^T), \tag{20}
$$

which has the form $\dot{\epsilon} = \frac{1}{\rho \log t} (2x \epsilon^T)$. Assuming $\epsilon$ of the form $\epsilon \propto \frac{1}{\rho \log t}$ we obtain $-\frac{1}{\rho \log t} = -B \frac{1}{\rho \log t}$. Thus the error indeed converges as $\epsilon \propto \frac{1}{\rho \log t}$.

A similar analysis for the weight normalization equations 17 considers the same dynamical system with a change in the equation for $v$, which becomes

$$
\dot{v} \propto e^{-\rho} \rho (I - vv^T)x. \tag{21}
$$

This equation differs by a factor $\rho^2$ from equation 20. As a consequence equation 21 is of the form $\dot{\epsilon} = -\frac{1}{\rho \log t} \epsilon$, with a general solution of the form $\epsilon \propto t^{-\frac{1}{2} \log t}$. In summary, GD with weight normalization converges faster to the same equilibrium than standard gradient descent: the rate for $\epsilon = v - x$ is $t^{-\frac{1}{2} \log t}$ vs $\frac{1}{\rho \log t}$.

Our goal was to find $\lim_{\rho \to \infty} \arg \min_{\|V_k\|_2 = 1 \forall k} L(\rho \hat{f})$. We have seen that various forms of gradient descent enforce different paths in increasing $\rho$ that empirically have different effects on convergence rate. It will be an interesting theoretical and practical challenge to find the optimal way, in terms of generalization and convergence rate, to grow $\rho \to \infty$.

Our analysis of simplified batch normalization (1) suggests that several of the same considerations that we used for weight normalization should apply (in the linear one layer case BN is identical to WN). However, BN differs from WN in the multilayer case in several ways, in addition to weight normalization: it has for instance separate normalization for each unit, that is for each row of the weight matrix at each layer.
Fig. 3. The top left graph shows testing vs training cross-entropy loss for networks each trained on the same data sets (CIFAR10) but with a different initializations, yielding zero classification error on training set but different testing errors. The top right graph shows the same data, that is testing vs training loss for the same networks, normalized by dividing each weight by the Frobenius norm of its layer. Notice that all points have zero classification error at training. The red point on the top right refers to a network trained on the same CIFAR-10 data set but with randomized labels. It shows zero classification error at training and test error at chance level. The top line is a square-loss regression of slope 1 with positive intercept. The bottom line is the diagonal at which training and test loss are equal. The networks are 3-layer convolutional networks. The left can be considered as a visualization of Equation 4 when the Rademacher complexity is not controlled. The right hand side is a visualization of the same relation for normalized networks that is conditions for \( N \) and for the architecture of the network the terms \( c_1 \mathcal{R}_N(\tilde{f}) + c_2 \sqrt{\frac{\ln(\frac{1}{\delta})}{N}} \) represent a small offset. From (55).

4. Discussion

A main difference between shallow and deep networks is in terms of approximation power or, in equivalent words, of the ability to learn good representations from data based on the compositional structure of certain tasks. Unlike shallow networks, deep local networks – in particular convolutional networks – can avoid the curse of dimensionality in approximating the class of hierarchically local compositional functions. This means that for such class of functions deep local networks represent an appropriate hypothesis class that allows good approximation with a minimum number of parameters. It is not clear, of course, why many problems encountered in practice should match the class of compositional functions. Though we and others have argued that the explanation may be in either the physics or the neuroscience of the brain, these arguments are not rigorous. Our conjecture at present is that compositionality is imposed by the wiring of our cortex and, critically, is reflected in language. Thus compositionality of some of the most common visual tasks may simply reflect the way our brain works.

Optimization turns out to be surprisingly easy to perform for overparametrized deep networks because SGD will converge with high probability to global minima that are typically much more degenerate for the exponential loss than other local critical points.

More surprisingly, gradient descent yields generalization in classification performance, despite overparametrization and even in the absence of explicit norm control or regularization, because standard gradient descent in the weights enforces an implicit unit \((L_2)\) norm constraint on the directions of the weights in the case of exponential-type losses.

In summary, it is tempting to conclude that the practical success of deep learning has its roots in the almost magic synergy of unexpected and elegant theoretical properties of several
aspects of the technique: the deep convolutional network architecture itself, its overparameterization, the use of stochastic gradient descent, the exponential loss, the homogeneity of the RELU units and visual cortex.

Of course many problems remain open on the way to develop a full theory and, especially, in translating it to new architectures. More detailed results are needed in approximation theory, especially for densely connected networks. Our framework for optimization is missing at present a full theory, especially for densely connected networks. The analysis of generalization should include an analysis of the trade-off between the weights for multilayer networks (see (4) and (3)). A full theory would also require an analysis of the trade-off between approximation and estimation error, relaxing the separability assumption.

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