On the Importance of Full Rank Initializations in Deep Neural Networks

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Abstract
Several methods have been proposed over the last few years for initializing the weights of neural networks in order to converge to better solutions or to reduce the time taken for convergence. On the other hand, there have been recent efforts connecting the full rank nature of weight matrices with the optimality of the final converged solution. In this work, we study the connection between popular initialization methods and the conditions necessary at optimal solution using deep linear networks with the squared loss. Through this connection, we attempt to provide a new explanation as to why these different initialization methods work well in practice.

1 Introduction
Over the past few years, several methods for initializing the weights of neural networks have been proposed. Although recent theoretical understanding claims that all local minima are perhaps equivalent [3], practical use of deep neural networks continues to rely on appropriate weight initialization methods. Glorot and Bengio [1] proposed a popular initialization method based on equalizing the variances of outputs of consecutive layers in a deep linear network. He et al [2] extended this work to provide a variant of this initialization. Few years later, Saxe et al [9] proposed their methods based on orthogonal weights. There have been continued efforts towards finding good initializations [7, 4, 10], due to their importance as suggested by Sutskever et al in [11]. On the other hand, there have been efforts connecting the full rank nature of weight matrices with the optimality of the final solution [12, 15, 13].

In this work, we seek to study the connection between well-known initialization methods that work well in practice and the necessary conditions imposed on the weight matrices at optimality. To the best of our knowledge, this is the first such study; although the findings are preliminary, we believe this is an interesting direction of study for the community to understand the training of deep neural networks.

2 Preliminaries
Consider a dataset of \( m \) points consisting of input-output pairs \((x, y)\), where each \( x \) is of dimension \( d_x \) and each \( y \) is of dimension \( d_y \), i.e., the inputs and outputs can be represented as matrices \( X \in \mathbb{R}^{d_x \times m} \) and \( Y \in \mathbb{R}^{d_y \times m} \) respectively. We consider a linear neural network of \( H \) hidden layers. Let the width of these \( H \) layers be \( d_1, d_2, \ldots, d_H \). The width of the input and output layers are \( d_x \) and \( d_y \) respectively, which we call \( d_0 \) and \( d_{H+1} \) for consistency of notation. The weight matrices connecting the \((i-1)\)th and the \(i\)th layers is denoted by \( W_i \in \mathbb{R}^{d_i \times d_{i-1}} \) for \( i = 1, 2, \ldots, H + 1 \). The output of this network can be written as a matrix product \( W_{H+1}W_H \cdots W_1X \), and our task is to minimize the sum-of-squares loss over the entire dataset defined below (note that our setup follows earlier efforts such as [3] and [12]):

\[
L(W) := \frac{1}{2} \| W_{H+1}W_H \cdots W_1X - Y \|_F^2 \tag{1}
\]

where \( W \) denotes the tuple of weight matrices \((W_{H+1}, W_H, \ldots, W_1)\) and \( \| . \|_F \) denotes the Frobenius norm.
Let $W_i^t$ represent the weight matrix $W_i$ after $t(t > 0)$ iterations of gradient descent during training. We denote the initial weight matrix before gradient descent begins by $W_i^0$.

**Assumptions:** Our results follow [3] and [14], and hence use the same assumptions: $d_x \leq m$, $d_y \leq m$, and $XX^T$, $YY^T$ are matrices of full rank. Furthermore, we assume that $d_0^i \geq d_1^i \geq \ldots \geq d_{H^i}^i$, i.e. the network is pyramidal in shape. Studying the results of this work relaxing these assumptions is an important direction of ongoing/future work.

3 Primary Claim

For convenience, we refer to commonly used initialization methods including Xavier initialization [1]. He initialization [2], zero-mean (truncated) Gaussian initialization and Gaussian variants of Le Cun initialization [6] as ‘well-known initialization methods’ in this work.

**Claim 1 (Informal).** Well-known initialization methods converge to a global minimum for deep linear neural networks satisfying assumptions in Section 2 when trained using Gradient Descent (GD).

**Justification.** Claim 1 will be shown as a result of the following sub-claims and results.

**Sub-Claim 1.** Well-known initialization methods initialize the matrices to be full rank with high probability.

**Proof.** The rank of a matrix of size $N \times n$ whose entries are sampled from a sub-Gaussian distribution with mean $0$ is $\min\{N, n\}$ (i.e., the matrix is full rank) with high probability. The complete result can be found in [8]. Hence, initializing the matrix using zero-mean (truncated) Gaussian initialization and Gaussian variants of Le Cun initialization [6], Xavier initialization [1] and He initialization [2] result in full-rank initialization.

While these methods generate full-rank matrices with high probability, other popular methods such as orthogonal initialization [9] and identity initialization [5] are full-rank certainly, by construction.

**Lemma 1.** Consider two weight matrices $W_i$ and $W_{i+1}$. If both of these matrices are full rank, then the product $W_{i+1}W_i$ is full rank.

**Proof.** We use the rank of product and Sylvester’s inequalities to prove this result. Note that $W_{i+1} \in \mathbb{R}^{d_{i+1} \times d_i}$ and $W_i \in \mathbb{R}^{d_i \times d_{i-1}}$. From our assumption on the width of the network, $d_{i+1} \leq d_i \leq d_{i-1}$, and hence rank($W_{i+1}$) = $d_{i+1}$ and rank($W_i$) = $d_i$, since both these matrices are full rank.

From the rank of product inequality, rank($W_{i+1}W_i$) $\leq \min\{d_{i+1}, d_i\} = d_{i+1}$. Using Sylvester’s inequality, rank($W_{i+1}W_i$) $\geq d_{i+1} + d_i - d_i = d_{i+1}$. From the two inequalities, we get rank($W_{i+1}W_i$) = $d_{i+1}$.

**Corollary 1.** In well-known initialization methods, the product of initialized weight matrices is full-rank with high probability, for deep linear networks satisfying the assumptions in Section 2.

**Proof.** This corollary directly follows from Lemma 1 and Sub-Claim 1.

Using the corollary above, we conclude that rank($W_H^0W_{H-1}^0 \ldots W_1^0$) = $d_y$.

**Sub-Claim 2.** The product of rank of weight matrices, rank($W_HW_{H-1} \ldots W_1$), follows a non-decreasing path over the iterations of Gradient Descent (GD). In other words, rank($W_H^tW_{H-1}^{t+1} \ldots W_1^{t+1}$) $\geq$ rank($W_H^{t+1}W_{H-1}^{t+1} \ldots W_1^{t+1}$) for all $t > 0$.

**Justification.** We justify this claim empirically at this time, and conducted an exhaustive set of experiments with the assumptions considered earlier. We trained deep linear neural networks with variable number of layers (ranging from 2 to 5) and of different sizes (ranging from 500 to 50), and plotted the variation of the rank of the product of weight matrices with iterations while training with GD. Owing to space constraints, we present herewith (Figures 1 and 2) sample plots supporting our claim. Our comprehensive experiments suggested that these claim hold almost all the time under the aforementioned assumptions, and proving this theoretically is part of our ongoing/future efforts.
Lemma 2. Under the assumptions and constructions stated so far, \( \text{rank}(W^t W^t_{H-1} \ldots W^t_1) = d_y \) for all \( t \geq 0 \), when \( \text{rank}(W^0_H W^0_{H-1} \ldots W^0_1) = d_y \).

Proof. Combining the facts that \( \text{rank}(W^0_H W^0_{H-1} \ldots W^0_1) = d_y \), \( \text{rank}(W^t H W^t_{H-1} \ldots W^t_1) \leq d_y \) for any \( t \), and that \( \text{rank}(W^{t+1}_H W^{t+1}_{H-1} \ldots W^{t+1}_1) \geq \text{rank}(W^t H W^t_{H-1} \ldots W^t_1) \) for all \( t \geq 0 \), we get that \( \text{rank}(W^t H W^t_{H-1} \ldots W^t_1) = d_y \).

Continuing from Corollary 1 and Lemma 2, we infer that the product of the weight matrices remain full rank during training using GD with well-known initialization methods in these settings. To summarize, we hypothesize that the listed well-known methods initialize networks with weight matrices in order that the product of weight matrices is full-rank (not necessarily with this objective though) and since GD follows a non-decreasing path of rank of product of weight matrices, the product of weight matrices remains full-rank throughout training.

We now connect the above observations to a result from [14] which provides necessary and sufficient conditions to check whether a given optimization method has converged to a global optimum. Under certain conditions, the result classifies the set of critical points into global optima or saddles based on the rank of product of weight matrices as specified below:

Theorem 1 (Thm 2.1, [14]). If \( k = \min\{d_x, d_y\} \), consider \( V_1 := \{(W_1, \ldots, W_{H+1}) : \text{rank}(W_{H+1} W_H \ldots W_1) = k\} \). Every critical point of \( L(W) \) in \( V_1 \) is a global minimum and every critical point of \( L(W) \) in \( V_1^c \) is a saddle point.

The sub-claims above and Theorem 1 allow us to infer that when using GD to train deep linear networks with the above assumptions, the training converges to a critical point (i.e. model) whose product of weight matrices is full rank, and hence results in a global minimum.

Claim 2. Any initialization that ensures that \( \text{rank}(W^0_H W^0_{H-1} \ldots W^0_1) = d_y \) in a linear network satisfying assumptions of Section 2 would lead to a convergence to a global minimum if GD converges.

Proof. The proof is straightforward from Sub-claim 2 and Theorem 1. Sub-claim 2 and Lemma 2 together imply that \( \text{rank}(W^t_H W^t_{H-1} \ldots W^t_1) = d_y \) for \( t \geq 0 \). And so when GD converges, the product of weight matrix is full rank, which in conjunction with theorem 1 implies that GD would converge to global minima.
The above claim gives a sufficient condition for any initialization to be good enough to lead to global convergence for deep linear networks.

We studied this hypothesis using a comprehensive set of experiments again, with a sample result shown in Figure 3. These empirical studies support our hypothesis that well-known initialization methods ensure that \( \text{rank}(W_0 H W_0 H^{-1} \ldots W_1^H) = d_y \), although not explicitly derived keeping this condition in mind. Beyond the objectives considered for these methods, we surmise that the connection that leads to their empirical success is perhaps that they all satisfy the above-mentioned sufficient condition (Claim 2).

Figure 3: Variation of rank with iterations for three networks and different initialization schemes. (Left:) Network architecture 100 x 75 x 50 with Gaussian/normal variant of Xavier’s initialization [1]. (Middle:) Network architecture 200 x 175 x 150 x 125 x 100 with Gaussian/normal variant of He’s initialization [2]. (Right:) Network architecture 500 x 416 x 333 x 250 with Saxe’s initialization [9]. Note that in all the cases, the initialization results in a full rank product and there is no decrease of rank over the iterations.

3.1 Some remarks on the claims

We now present some remarks to dispel misconceptions that might lead from our results. Firstly, we do not claim that GD does not converge to global minima if \( \text{rank}(W_0 H W_0 H^{-1} \ldots W_1^H) < d_y \). We observe empirically that the rank may increase over iterations, such that \( \text{rank}(W_t^H W_{t-1}^H \ldots W_1^H) = d_y \) after a few iterations (which is in accordance with Sub-claim 2), which can result in global convergence. We observe this phenomenon in some of our experiments, presented below in Figure 4 (first two plots). In fact, it seems that when the initialization is such that \( \text{rank}(W_0 H W_0 H^{-1} \ldots W_1^H) \approx d_y \), it is almost always the case that \( \text{rank}(W_t^H W_{t-1}^H \ldots W_1^H) = d_y \) after a finite number of iterations.

Figure 4: The first two plots show the variation of rank with iterations when starting with a rank close to full rank for architectures 100 x 75 x 50 and 100 x 83 x 66 x 50 (full rank: 50 in both scenarios), while the last two show the variation when starting with a low rank for the same architectures. In the first two plots, the rank becomes full i.e., 50, while it saturates at 10 (which is not full rank) in the last two plots.

We also add that although it might seem from the above experiments that if GD were run for a very large number of iterations, it will always lead to product of weight matrices being full rank; this however is not the case. In fact, if the initialization is such that \( \text{rank}(W_0 H W_0 H^{-1} \ldots W_1^H) \) is not very close to \( d_y \), the rank of the product of weight matrices, \( \text{rank}(W_t^H W_{t-1}^H \ldots W_1^H) \), saturates after a certain number of iterations without reaching full rank. The last two plots in Figure 4 show this phenomenon. It is clear from Theorem 1 that in this case as \( \text{rank}(W_t^H W_{t-1}^H \ldots W_1^H) \neq d_y \), the network has not converged to global minimum, but to a local minimum or a saddle point.

Conclusions and Future Work. In this work, we studied the connection between weight initialization methods and the rank of the product of the weight matrices, and showed interesting observations that could have impact on the training of deep neural networks. Our future work includes studying the aforementioned claims relaxing the assumptions made, as well as the theoretical justification of claims shown empirically in this work.
References


