Mitigating deep double descent by concatenating inputs

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Abstract
The double descent curve is one of the most intriguing properties of deep neural networks. It contrasts the classical bias-variance curve with the behavior of modern neural networks, occurring where the number of samples nears the number of parameters. In this work, we explore the connection between the double descent phenomena and the number of samples in the deep neural network setting. In particular, we propose a construction which augments the existing dataset by artificially increasing the number of samples. This construction empirically mitigates the double descent curve in this setting. We reproduce existing work on deep double descent, and observe a smooth descent into the overparameterized region for our construction. This occurs both with respect to the model size, and with respect to the number of epochs.

1. Introduction
Underparameterization and overparameterization are at the heart of understanding modern neural networks. The traditional notion of underparameterization and overparameterization led to the classic U-shaped generalization error curve (Hastie et al., 2001; Geman et al., 1992), where generalization would worsen when the model had either too few (underparameterized) or too many parameters (overparameterized). Correspondingly, it was expected that an underparameterized model would underfit and fail to identify more complex and informative patterns, and an overparameterized model would overfit and identify non-informative patterns.

This view no longer holds for modern neural networks. It is widely accepted that neural networks are vastly overparameterized, yet generalize well. There is strong evidence that increasing the number of parameters leads to better generalization (Zagoruyko and Komodakis, 2016; Huang et al., 2017; Larsson et al., 2016), and models are often trained to achieve zero training loss (Salakhutdinov, 2017), while still improving in generalization error, whereas the traditional view would suggest overfitting.

To bridge the gap, (Belkin et al., 2018) proposed the double descent curve, where the underparameterized region follows the U-shaped curve, and the overparameterized region smoothly decreases in generalization error, as the number of parameters increases further. This results in a peak in generalization error, where a fewer number of samples would counter-intuitively decrease the error. There has been extensive experimental evidence of the double descent curve in deep learning (Nakkiran et al., 2019; Yang et al., 2020), as well as in models such as random forests, and one layer neural networks (Belkin et al., 2018; Ba et al., 2020).

One recurring theme in the definition of overparameterization and underparameterization lies in the number of neural network parameters relative to the number of samples (Belkin et al., 2018; Nakkiran et al., 2019; Ba et al., 2020; Bibas et al., 2019; Muthukumar et al., 2019; Hastie et al., 2019). On a high level, a greater number of parameters than samples is generally considered overparameterization, and fewer is considered underparameterization.

However, this leads to the question “What is a sample?”
In this paper, we revisit the fundamental underpinnings of overparameterization and underparameterization, and stress test when it means to be overparameterized or underparameterized, through extensive experiments of a cleverly constructed input. We artificially augment existing datasets by simply stacking every combination of inputs, and show the mitigation of the double descent curve in the deep neural network setting. We humbly hypothesize that in deep neural networks we can, perhaps, artificially increase the number of samples without increasing the information contained in the dataset, and by implicitly changing the classification pipeline mitigate the double descent curve. In particular, the narrative of our paper obeys the following:

- We propose a simple construction to artificially augment existing datasets of size $O(n)$ by stacking inputs to produce a dataset of size $O(n^2)$.
- We demonstrate that the construction has no impact on
we introduce the concatenated inputs construction with more parameters in the overparameterized region, the size of a dataset can be artificially, but non-trivially, increased. This construction can be applied both to the regression setting and the classification setting. In linear regression, for given input pairs, \((x_1, y_1), (x_2, y_2)\), an augmented dataset can be constructed:

\[
\begin{align*}
\{(x_1, x_1, y_1 + y_2/2), (x_2, x_2, y_1 + y_2/2), \\
(x_1, x_1, y_2/2 + y_1/2), (x_2, x_2, y_2/2 + y_1/2)\}
\end{align*}
\]

where \([\alpha, \beta]\) represents concatenation of the input \(\alpha, \beta\). In the setting of classification, the process is identical, where the targets are produced by element-wise addition and then averaged to sum to 1. The averaging is not strictly necessary even in the deep neural network classification case, where the binary cross entropy loss can be used instead of cross entropy. For test data, we concatenate the same input with itself, and the target is the original target. This way a dataset of size \(O(n)\) is artificially augmented to size \(O(n^2)\).

Our reasons for the concatenated inputs construction are as follows: (i) there is limited injection of information or semantic meaning; (ii) the number of samples is significantly increased. For the purposes of understanding underparameterization, overparameterization and double descent, such a construction tries to isolate the definition of a sample.

2. Related Works

The double descent curve was proposed recently (Belkin et al., 2018), where the authors define over/underparameterization as the proportion of parameters to samples, explained through model capacity. With more parameters in the overparameterized region, there is larger “capacity” (i.e., the model class contains more candidates), and thus may contain better, simpler models by Occam’s Razor rule. The interpolation region is suggested to exist when the model capacity is capable of fitting the data nearly perfectly by overfitting on non-informative features, resulting in higher test error.

Double descent is also observed in deep neural networks (Nakkiran et al., 2019), in addition to epoch-wise double descent. Experimentation is amplified by label noise. With the observation of unimodel variance (Neal et al., 2018), (Yang et al., 2020) decomposes the risk into bias and variance, and posits that double descent arises due to the bell-shaped variance curve rising faster than the bias decreases.

There is substantial theoretical work on double descent, particularly in the least squares regression setting. (Advani and Saxe, 2017) analyses this linear setting and proves the existence of the interpolation region, where the number of parameters equals the number of samples in the asymptotic limit. Another work (Hastie et al., 2019) proves that regularization reduces the peak in the interpolation region. (Belkin et al., 2019) requires only finite samples, where the features and target be jointly Gaussian. Other papers with similar setup include (Muthukumar et al., 2019; Bibas et al., 2019; Mitra, 2019; Mei and Montanari, 2019; Ba et al., 2020; Nakkiran, 2019; Bartlett et al., 2019; Chen et al., 2020).

3. Methods

We introduce the concatenated inputs construction. This way the size of a dataset can be artificially, but non-trivially, increased. This construction can be applied both to the regression setting and the classification setting. In linear regression, for given input pairs, \((x_1, y_1), (x_2, y_2)\), an augmented dataset can be constructed:

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4. Results

In this section, we reproduce settings from benchmark double descent papers, add the concatenated inputs construction and analyze the findings.

![Figure 1.](image)

**Figure 1.** Left: The standard case. Right: The concatenated inputs construction. Plots of the Test MSE versus the number of samples (pre-concatenation) for min-norm ridgeless regression, where \(d = 30\). Following (Nakkiran, 2019), inputs are drawn \(x \sim N(0, I_d)\), target \(y = \theta x + N(0, \sigma^2)\), where \(\theta\) are the parameters, \(||\theta||_2 = 1\), \(\sigma = 0.1\). \(\theta = X^\dagger y\). As expected, the concatenated inputs construction does not affect the double descent curve, and the peak occurs in the exact same location.

4.1. Linear regression

The linear regression setting has been a fruitful testbed for empirical work in double descent, as well as yielding substantial theoretical understanding. The concatenated inputs construction is applied similarly here, however with different motivation. Namely, we wish to motivate that the concatenated inputs construction is not expected to add any information and is therefore not expected to impact the double descent curve.

We reproduce the linear regression setting from (Nakkiran, 2019), given in Figure 1. For the concatenated inputs construction, we first draw the number of samples before concatenation and construction of the augmented dataset. We observe that, by construction, the concatenated inputs...
In this section, we follow (Yang et al., 2020) and decompose the loss into bias and variance. Namely, let $\mathcal{CE}$ denote the cross entropy loss, $T$ a random variable representing the training set, $\pi$ is the true one-hot label, $\bar{\pi}$ is the average log-probability after normalization, and $\hat{\pi}$ is the output of the neural network. Then,

$$
E_T[\mathcal{CE}(\pi, \hat{\pi})] = D_{KL}(\pi||\bar{\pi}) + E_T[D_{KL}(\bar{\pi}||\hat{\pi})]
$$

where the first component is the bias and the second component is the variance. On a high level, the variance can then be estimated by training separate same capacity models on different splits of the dataset, and then measuring the difference in outputs on the same test set. The bias is then computed by subtracting the empirical variance from the empirical risk. For finer details, see (Yang et al., 2020)
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For training, we follow (Yang et al., 2020) and train a ResNet-34 (He et al., 2016) on CIFAR10, with stochastic gradient descent (learning rate = 0.1, momentum = 0.9). The learning rate is decayed a factor of 0.1 every 200 epochs, with a weight decay of 5e-4. The width $k$ of the network is varied suitably between 1 and 64. We also make 5 splits of 10,000 training samples for the calculation of bias/variance.

We present results in Figure 4. The concatenated inputs construction significantly delays and smoothens the increase in variance relative to the standard case, where the unimodal variance is significantly sharper. This impacts the shape of the test error, where in this setting we see a shifted bump in test error for the concatenated inputs construction. One possible explanation is in the case of deep neural networks the concatenated inputs construction is a form of implicit regularization for small models, which controls overfitting and leads to a smoother variance curve.

5. Discussion

While the understanding of over/underparameterization and double descent is strongly tied to the number of samples, it appears that given a fixed number of unique samples it is possible to manipulate over/underparameterization by artificially boosting the number of samples by the concatenated inputs construction.

A possible explanation in the literature for double descent is that the model is being forced to fit the training data as perfectly as possible, and at some model capacity it is possible to fit the training data perfectly by overfitting on non-existent, or weakly present, features. This results in overfitting and the double descent curve. Interestingly, the concatenated inputs construction generally mitigates double descent, even though it is possible to build models for the concatenated inputs construction from models for the standard setting. This suggests a possible route to improve the understanding of the relationship to the model capacity.

The experimental results in this work support that neural network behavior can change with respect to the number of samples, even if the majority of samples add limited information, via the concatenated inputs construction. In this view, the concatenated inputs construction creates possibly a huge dataset, e.g. 50,000² samples for the originally 50,000 samples CIFAR10 where 50,000² = 2,500,000,000 is far larger than any neural network for CIFAR10. Yet, there is no noticeable underfitting. Namely, the concatenated inputs construction quickly achieves comparable performance to the standard one-hot vector deep learning setting. This suggests we may need to rethink the relationship between underfitting, the number of parameters, samples, and model capacity.

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Figure 3. Top Left: Model Size Double Descent on CIFAR10. Orange line is validation error, blue line is train error. Top Right: Model Size Double Descent on CIFAR100. Orange line is validation error, blue line is train error. Bottom Left: Epoch-wise Double Descent on CIFAR10. Bottom Right: Epoch-wise Double Descent on CIFAR100. For each subplot, Left: Standard one-hot vector setup, Right: Concatenated inputs construction. Bottom plots legends’ represent the width $k$ of the ResNet18. Models are ResNet18 architecture on CIFAR-10 and CIFAR-100 datasets.

Figure 4. Left: Bias and Variance against width $k$ of ResNet-34 in the standard case. Middle: Bias and Variance against width $k$ of ResNet-34 in the concatenated inputs construction. Right: Test error against width $k$ of ResNet-34 in the standard case and concatenated inputs construction.
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