Abstract

We show that the error of iteratively magnitude-pruned networks empirically follows a scaling law with interpretable coefficients that depend on the architecture and task. We functionally approximate the error of the pruned networks, showing it is predictable in terms of an invariant tying width, depth, and pruning level, such that networks of vastly different pruned densities are interchangeable. We demonstrate the accuracy of this approximation over orders of magnitude in depth, width, dataset size, and density. We show that the functional form holds (generalizes) for large scale data (e.g., ImageNet) and architectures (e.g., ResNets). As neural networks become ever larger and costlier to train, our findings suggest a framework for reasoning conceptually and analytically about a standard method for unstructured pruning.

1. Introduction

For decades, neural network pruning—eliminating unwanted parts of a network—has been a popular approach for reducing network sizes or the computational demands of inference (LeCun et al., 1990; Reed, 1993; Han et al., 2015). In practice, pruning can reduce the parameter-counts of contemporary models by 2x (Gordon et al., 2020) to 5x (Renda et al., 2020) with no increase in error. More than 80 pruning techniques have been published in the past decade (Blalock et al., 2020), but, despite this enormous volume of research, there remains little guidance on important aspects of pruning. Consider a seemingly simple question one might ask when using a particular pruning technique:

Given a family of neural networks (e.g., ResNets on ImageNet of various widths and depths), which family member should we prune (and by how much) to obtain the network with the smallest parameter-count such that error does not exceed some threshold \( \epsilon_k \)?

As a first try, we could attempt to answer this question using brute force: we could prune every member of a network family (i.e., perform grid search over widths, depth, and pruned densities) and select the smallest pruned network that satisfies our constraint on error. However, depending on the technique, pruning one network (let alone grid searching) could take days or weeks on expensive hardware.

If we want a more efficient alternative, we will need to make assumptions about pruned networks: namely, that there is some structure to the way that their error behaves. For example, that pruning a particular network changes the error in a predictable way. Or that changing its width or depth changes the error when pruning it in a predictable way. We could then train a smaller number of networks, characterize this structure, and estimate the answer to our question.

We have reason to believe that such structure does exist for certain pruning methods, since there are already techniques that take advantage of it implicitly. For example, Cai et al. (2019) create a single neural network architecture that can be scaled down to many different sizes; to choose which subnetwork to deploy, Cai et al. train an auxiliary, black-box neural network to predict subnetwork performance. Although this black-box approach implies the existence of structure for this pruning method, it does not reveal this structure explicitly or make it possible to reason analytically in a fashion that could answer our research question.

For other aspects of deep learning beyond pruning, such structure has been observed—and, further, codified explicitly—yielding insights and predictions in the form of scaling laws. Tan & Le (2019) design the EfficientNet family by developing a heuristic for predicting efficient tradeoffs between depth, width, and resolution. Hestness et al. (2017) observe a power-law relationship between dataset size and the error of vision and NLP models. Rosenfeld et al. (2020) observe a power-law relationship between dataset size and the error of vision and NLP models. Rosenfeld et al. (2020) observe a power-law relationship between dataset size and the error of vision and NLP models. Rosenfeld et al. (2020) observe a power-law relationship between dataset size and the error of vision and NLP models. Kaplan et al. (2020) develop a similar power law for language models that incorporates the computational cost of training.

Inspired by this work, we address our research question about pruning by finding a scaling law to predict the error of pruned networks. We focus on a pruning method called iterative magnitude pruning (IMP), where weights with the
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We study iterative magnitude pruning (IMP) (Janowsky, 1989; Han et al., 2015). IMP prunes by removing a fraction—typically 20%—of weights without regard to specific layers, resulting in a simple functional form and interpretable parameters. Our scaling law enables a framework for reasoning analytically about IMP, allowing us to answer our motivating question and similar questions about pruning.

2. Experimental Setup

Pruning. We study iterative magnitude pruning (IMP) (Janowsky, 1989; Han et al., 2015). IMP prunes by removing a fraction—typically 20%—of weights without regard to specific layers. We use per-weight magnitude pruning because it is generic, well-studied (Han et al., 2015), and produces state-of-the-art tradeoffs between density and error (Gale et al., 2019; Blalock et al., 2020; Renda et al., 2020).

Pruning weights typically increases the error of the trained network, so it is standard practice to further train after pruning to reduce error. For IMP, we use a practice called weight rewinding (Frankle et al., 2020; Renda et al., 2020), in which the values of unpruned weights are rewound to their values earlier in training (in our case, epoch 10) and the training process is repeated from there to completion. To achieve density levels below 80%, this process is repeated iteratively—rewinding by 20%, rewinding, and retraining—until a desired density level is reached. For a formal statement of this pruning algorithm, see Appendix A.

Datasets. In the main body of the paper, we study the image classification tasks CIFAR-10 and ImageNet. Our scaling law predicts the error when training with the entire dataset and smaller subsamples. We use subsampling because it provides a cost-effective way to collect some of the data for fitting our functional form. To subsample a dataset to a size of \( n \), we randomly select \( n \) of the training examples without regard to individual classes such that in expectation we preserve the original dataset distribution (we always retain the entire test set). When performing iterative pruning, we maintain the same subsample for all pruning iterations. We consider other datasets in Appendix E.

Networks. In the main body of the paper, we study ResNets for CIFAR-10 and ImageNet. We develop a scaling law that predicts the error (when pruned) of an entire family of networks with varying widths and—unlike the CIFAR-10 ResNets—depths. To vary width, we multiply the number of channels in each layer by a width scaling factor. To vary depth of the CIFAR-10 ResNets, we vary the number of residual blocks. We refer to a network by its depth \( l \) (the number of layers in the network, not counting skip connections) and its width scaling factor \( w \). We consider other networks in Appendix E.

Notation and terminology. Throughout the paper, we use the following notation and terminology:

- \( \mathcal{D}_N = \{ (x_i, y_i) \}_{i=1}^N \) is a labeled training set with \( N \) examples. A subsample of size \( n \) is a subset of \( \mathcal{D}_N \) with \( n \) examples selected uniformly at random.
- \( l \) and \( w \) are, respectively, the depth (i.e., the number of layers, excluding skip connections) and the width scaling factor of a particular network.
- Networks that vary by width and depth are a family.

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1We do not prune biases or BatchNorm, so pruning 20% of weights prunes fewer than 20% of parameters.

2See Appendix B for full details on architectures and hyperparameters. Note that CIFAR-10 and ImageNet ResNets have different architectures (He et al., 2016).
3. Modeling the Error of a Pruned Network

Our goal in this section is to develop a functional form that models the error of a member of a network family as it is pruned (using IMP) based on its unpruned error $\epsilon_{\text{up}}(w, l, n)$. In other words, we wish to find a function $\hat{\epsilon}(\epsilon_{\text{up}}, d | l, w, n)$ that predicts the error at each density $d$ for a network with a specific depth $l$, width scaling factor $w$, and dataset size $n$.

Intuition. Since IMP prunes 20% at a time, it produces pruned networks at intermediate densities $d_K = 0.8^K$ in the process of pruning to density $d_K = 0.8^K$. In Figure 1 (left), we plot the error of these pruned networks for CIFAR-10 ResNets with depth $l = 20$ and different widths $w$. All of these curves follow a similar pattern.\footnote{The same patterns occur for $l$ and $n$ for CIFAR-10 and $w$ and $n$ for ImageNet (see Appendix C). We focus on width for CIFAR-10 here for illustration.}

- $d$ is the density of a pruned network (i.e., the fraction of weights that have not been pruned).
- $\epsilon(d, l, w, n)$ is the test error of a network with the specified density, depth, width scaling, and dataset size.
- $\epsilon_{\text{up}}(l, w, n) = \epsilon(1, l, w, n)$ is the test error of the unpruned network with the specified depth, width scaling, and dataset size. When clear from context, we omit $(w, l, n)$ and write $\epsilon_{\text{up}}$.
- $\hat{\epsilon}(\epsilon_{\text{up}}, d | l, w, n)$ is an estimate of the error of a pruned model for a scaling law that has been fitted to a specific network with the specified depth, width scaling, and dataset size (Section 3).
- $\hat{\epsilon}(\epsilon_{\text{up}}, d, l, w, n)$ is an estimate of the error of a pruned model with the specified depth, width scaling, and dataset size for a scaling law that has been fitted to a network family (Section 4).

Dimensions. In developing our scaling laws, we vary four dimensions: dataset subsample size ($n$) and network degrees of freedom density ($d$), network depth ($l$), and width scaling factor ($w$). In the main body of the paper, we consider the ranges of these values as specified in Table 1. See the caption in Table 1 for full details. We train three replicates of each CIFAR-10 configuration with different seeds.

### 274 ImageNet ResNet points. Note that we use these configurations to find and evaluate the functional form. Once we do so, far fewer configurations are needed to fit the functional form for each setting (see Section 5).

| Network Family | Densities ($d$) | Depths ($l$) | Width Scalings ($w$) | Subsample Sizes ($n$) |
|----------------|----------------|-------------|----------------------|----------------------|
| CIFAR-10 ResNet | $0.8^i, i \in \{0, \ldots, 40\}$ (see below) | $l \in \{8, 14, 20, 26, 30, 98\}$ (see below) | $2^l, i \in \{-4, \ldots, 2\}$ (see below) | $\frac{3}{2}, i \in \{2, 1, 4, 8, 16, 32, 64\}$ |
| ImageNet ResNet | $0.8^i, i \in \{0, \ldots, 30\}$ (see below) | $l \in \{8, 14, 20, 26, 30, 98\}$ (see below) | $2^l, i \in \{-4, \ldots, 0\}$ (see below) | $\frac{3}{2}, i \in \{2, 1, 4\}$ |

Table 1. The ranges of settings we consider in our experiments in the main body of the paper. We consider all densities $d \in \{0.8^i | i \in \mathbb{N}_0\}$ where (1) the network is not disconnected and (2) the network does better than random chance; this value varies based on the configuration ($l, w, n$) and the random seed. Where neither of these conditions apply, we cap and $n$.

- Observation 1: Low-error plateau. The densest networks (right part of curves) have similar error to the unpruned network: $\epsilon_{\text{up}}(w)$. We call this the low-error plateau.
- Observation 2: Power-law region. When pruned further, error increases linearly on the logarithmic axes of the figure. Linear behavior on a logarithmic scale is the functional form of a power law, where error relates to density via exponent $\gamma$ and coefficient $c$: $\epsilon(d, w) \approx cd^{-\gamma}$. In particular, $\gamma$ is the slope of the line on the logarithmic axes.
- Observation 3: High-error plateau. When pruned further, error again flattens; we call this the high-error plateau and call the error of the plateau $\epsilon^3$.

Figure 1 (center) labels these regions for CIFAR-10 ResNet-$20 (w = 1, n = 1)$ and shows an approximation of these regions that is piece-wise linear on logarithmic axes. These observations are our starting point for developing a functional form that estimates error when pruning.

Functional form. Our next task is to find a functional form that captures these observations about the relationship between density and error. In prior work, Rosenfeld et al. (2020) observe that the relationship between width and error shares the same general shape: it has a region of lower error, a power-law region, and region of higher error. However, this relationship is different enough from the one we observe (see Appendix G) to merit an entirely new functional form.

To develop this functional form, we note that the three regions of the curves in Figure 1 (the low-error plateau, the power-law region, and the high-error plateau) can be described by three power laws: two plateaus with exponent $\gamma$ and an intermediate region with exponent $\gamma$. A functional family that arises frequently in systems that exhibit different power-law regions is the rational family. The particular family member we consider is as follows:\footnote{The expression $\left\| \frac{d - j \varphi}{d - j \varphi} \right\|^\gamma$ can be rewritten as $\epsilon_{\text{up}} \left( \frac{d^2 + p^2(\epsilon^1/\epsilon_{\text{up}})^2/\gamma}{d^2 + p^2} \right)^{\gamma/2}$.}

$$\hat{\epsilon}(\epsilon_{\text{up}}, d, l, w, n) = \epsilon_{\text{up}} \left[ \frac{d - j \varphi}{d - j \varphi} \right]^\gamma \left. \epsilon_{\text{up}} \right|_{d = j \varphi} \left( \frac{d^2 + p^2(\epsilon^1/\epsilon_{\text{up}})^2/\gamma}{d^2 + p^2} \right)^{\gamma/2} \quad \text{where} \quad j = \sqrt{-1} \quad (1)$$
This function’s shape is controlled by \( \epsilon_{np} \), \( \epsilon^1 \), \( \gamma \), and \( p \) (visualized in Figure 1, right). \( \epsilon_{np} \) and \( \epsilon^1 \) are the values of the low and high-error plateaus. \( \gamma \) is the slope of the power-law region on logarithmic axes. \( p \) controls the density where the high-error plateau transitions to the power-law region.

Fitting. To fit \( \hat{\epsilon}(\epsilon_{np}, d | l, w, n) \) to actual data \( \epsilon(d, l, w, n) \), we estimate values for the free parameters \( \epsilon^1 \), \( \gamma \), and \( p \) by minimizing the relative error \( \delta \equiv \frac{\epsilon_{np}(d, l, w, n) - \epsilon(d, l, w, n)}{\epsilon(d, l, w, n)} \) using least squares regression. The fit is performed separately for each configuration \((l, w, n)\) for all densities, resulting in per-configuration estimates of \( \hat{\epsilon}^1 \), \( \hat{\gamma} \), and \( \hat{p} \).

Evaluating fit. For a qualitative view,\(^5\) we plot the actual error\(^6\) \( \epsilon(d, l, w, n) \) and the estimated error \( \hat{\epsilon}(\epsilon_{np}, d | l, w, n) \) as a function density for CIFAR-10 ResNets of varying widths (Figure 2, left). Our estimated error appears to closely follow the actual error. The most noticeable deviations occur at large densities, where the error decreases slightly on the low-error plateau whereas we treat it as flat (see Section 6).

Quantitatively, we measure the extent to which estimated error departs from the actual error using the mean \( \mu \) and standard deviation \( \sigma \) of the relative deviation \( \delta \). Figure 2 (center) compares the estimated and actual errors for the networks in Figure 2 (left); Figure 2 (right) shows the same comparison for all configurations of \( l, w, \) and \( n \) on CIFAR-10 and the more than 4,000 pruned ResNets that result. The relative deviation on all configurations has mean \( \mu < 2\% \) and standard deviation \( \sigma < 4\% \); this means that, if the actual error is 10\%, the estimated error is 9.8±0.4\% \((\hat{\epsilon} = (1-\delta)\epsilon)\).

4. Jointly Modeling Error For All Dimensions

In Section 3, we found a functional form \( \hat{\epsilon}(\epsilon_{np}, d | l, w, n) \) (Eq. 1) that accurately predicts the error when pruning a specific member of a network family (with depth \( l \) and width \( w \)) trained with a dataset of size \( n \). The parameters governing Equation 1 \((\epsilon^1, p, \) and \( \gamma \) ) varied between and depended on the specific configuration of \( l, w, n \). However, we are interested in a single joint scaling law \( \hat{\epsilon}(\epsilon_{np}, d, l, w, n) \) that, given the unpruned network error \( \epsilon_{np}(l, w, n) \), accurately predicts error across all dimensions: all members of a network family that vary in depth and width, all densities, and all dataset sizes. Importantly, the parameters of this scaling law must be constants as a function of all dimensions. In this section, we develop this joint scaling law.

Intuition: the error-preserving invariant. Our desired scaling law \( \hat{\epsilon}(\epsilon_{np}, d, l, w, n) \) will be a four-dimensional function of \( d, w, l, \) and \( n \). To develop an intuition for

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\(^5\)Error is a 4-dimensional function, so we can only qualitatively examine 2D projections. All such projections are in Appendix D.

\(^6\)We compute the error as the mean across three replicates with different random seeds and dataset subsamples.
We refer to the expression \( l^p w^q d = \nu \) which network error is \((\text{Figure 3, left}). The constant-error contours are linear except canceling, we arrive at the following expression:

\[
\hat{\epsilon}(\epsilon_{np}, d, l, w, n) = \epsilon_{np} \left( \frac{m^{*} - gp}{m^{*} - gp^{*}} \right)^{\frac{1}{p}} = \epsilon_{np} \left( \frac{w^{*} - dp}{w^{*} - dp^{*}} \right)^{\frac{1}{q}}
\]

which is the joint functional form \( \hat{\epsilon}(\epsilon_{np}, d, l, w, n) \) of all four dimensions \( d, l, w, \) and \( n \). Critically, for this to be a useful joint form, the free parameters \( \epsilon^{1}, p^{*}, \) and \( \gamma \) must be constants shared across all possible values of \( d, l, w, \) and \( n \). We will assume this is the case and directly quantify how well this assumption holds in the evaluation section below.

For qualitative intuition as to why this is a reasonable assumption, consider the relationship between \( m^{*} \) and the test error of pruned networks as we vary depth, width, and dataset size (Figure 4). Across all projections, the annotated \( \epsilon^{1} \) (error of the high-error plateau), \( \gamma \) (slope of the power-law region) and \( p^{*} \) (value of \( m^{*} \) where the high-error plateau transitions to the power-law region) appear the same.

The preceding discussion addresses how we handle \( l, w, \) and \( d \) in our joint scaling law. We address dataset size \( n \) in Eq. 2 implicitly through the way that it affects \( \epsilon_{np}, \) and we validate that this is a reasonable choice through the evaluation below. We retain the explicit form \( \hat{\epsilon}(\epsilon_{np}, d, l, w, n) \) to stress that the lack of explicit dependency on \( n \) is non-trivial and was not known prior to our work.

**Fitting.** To fit \( \hat{\epsilon}(\epsilon_{np}, d, l, w, n) \) to the actual data \( \epsilon(d, l, w, n) \), we estimate values for the free parameters \( \epsilon^{1}, \gamma, p^{*}, \phi \) and \( \psi \) by minimizing the relative error \( \delta \triangleq \frac{\hat{\epsilon}(\epsilon_{np}, d, l, w, n) - \epsilon(d, l, w, n)}{\epsilon(d, l, w, n)} \) using least squares regression. The fit is performed jointly over all configurations of \( d, l, w, \) and \( n, \) resulting in joint estimates of \( \hat{\epsilon}^{1}, \hat{\gamma}, \hat{p}, \phi, \) and \( \psi \). One can also perform a partial fit for a subset of dimensions (e.g., just \( d, l, \) and \( n \)) by omitting \( \phi \) and/or \( \psi \) (see Appendix D).

**Evaluating fit.** In Figure 5, we plot the actual error \( \epsilon(d, l, w, n) \) and the estimated error \( \hat{\epsilon}(\epsilon_{np}, d, l, w, n) \) for the CIFAR-10 ResNets (all widths, depths and dataset sizes) and ImageNet ResNets (all widths and dataset sizes for depth 50). As in Section 3, our estimated error appears to closely follow the actual error. Deviations arise mainly at high densities where error decreases below \( \epsilon_{np} \) and low densities approaching high error saturation.

We again quantify the fit of the estimated error using the mean \( \mu \) and standard deviation \( \sigma \) of the relative deviation \( \delta \). The relative deviation on the joint scaling laws for the CIFAR-10 and ImageNet networks has a mean \( \mu < 2\% \) and standard deviation of \( \sigma < 6\% \).

To contextualize these results, Figure 5 (right) quantifies the variation in error we see over multiple replicates of the CIFAR-10 experiments due to using different random seeds. It plots the minimum, maximum, and mean errors across the three replicates we ran.\(^7\) The variation across trials has a standard deviation of \( \sigma = 3.4\% \), sizeable relative to the estimation error of \( \sigma = 5.8\% \) for the joint scaling law. This

\(^7\)We only ran a single replicate of the ImageNet experiments due to the significant cost of collecting data.
indicates that a significant portion of our error may stem from measurement noise.

The functional form has just five parameters and obtains an accurate fit on 4,301 points on CIFAR-10 and 274 points on ImageNet, suggesting it is a good approximation. In Appendix E, we show that it achieves a similarly good fit for additional architectures and datasets. In Section 5, we show that, although we use a large number of points to develop and evaluate our functional form here, it is possible to get a good fit with far fewer points and the fit has low sensitivity to the choice of points.

5. Sensitivity of Fit to Number of Points

In Section 4, we showed that our scaling law was accurate when we fit it on all of the available data. Now that we possess the functional form and know that it can accurately model the behavior of IMP, we study the amount of data necessary to obtain a stable,\(^8\) accurate fit. This question is especially relevant when the functional form is applied to new settings—new networks, datasets, or pruning algorithms—and we must collect new data to do so. The functional form has only five parameters, suggesting that few experiments will be necessary to obtain an accurate fit.

Experiments. To evaluate the effect of the number of points on the stability and accuracy of the fit, we randomly sample varying numbers of points, fit the scaling law to those points, and evaluate the quality of the fit over all points. We sample these points in two ways.

Experiment 1. Randomly sample \(T\) networks \((w, l, n, d)\). This experiment evaluates the stability and accuracy of the fit when naively varying the number of points.

Experiment 2. Randomly sample \(T\) network configurations \((w, l, n)\) and include all densities \(d\) for each configuration. This experiment captures the specific use case of IMP, where obtaining data at density \(d\) requires obtaining all densities \(d' > d\). As such, we anticipate that data will be obtained by iteratively pruning a small number of configurations \((w, l, n)\) to low density.

Results. We perform each experiment for many different values of \(T\) on the CIFAR-10 ResNets pruned with IMP. We repeat the experiment at each value of \(T\) 30 times with different samples of points and report the mean and standard deviation of \(\mu\) and \(\sigma\) for the fit. Experiments 1 and 2 respectively appear in Figure 6 left and right. The shaded
areas represent one standard deviation from the mean in each direction. On Experiment 1, when just 40 networks \((w, l, d, n)\) are available, the standard deviation on both \(\mu\) and \(\sigma\) is just one percentage point. On Experiment 2, when just 15 random configurations of \((w, l, n)\) are available at all densities, we similarly achieve standard deviation below 1%. In both cases, as the number of networks increases, the standard deviation decreases further.

These results show that, now that our scaling law is known, it is possible to obtain an accurate (and stable) estimation using far less data than we used to evaluate the quality of the fit in Section 4. This implies that translating our scaling law to new settings will be far less data-intensive than developing and evaluating it in the first place. Moreover, the results in this section reflect a particularly naive way of selecting points: doing so randomly; we made no effort to ensure that, for example, the networks represented a diverse range of widths, depths, dataset sizes, and densities. By selecting these networks in a strategic way, it may be possible to further reduce the number of networks necessary to obtain a similarly accurate fit.

6. Discussion: Selecting a Functional Form

We have shown that our proposed functional form \(\hat{\epsilon}(d, l, w, n)\) accurately approximates the error when pruning families of neural networks. In this section, we discuss some of the key criteria that led us to select this particular functional form. We intend this section to provide insight into our choices in the context of the broader design space and to highlight opportunities for further refinement.

**Criterion 1: Transitions.** In Section 3, we observe that, when pruning a neural network with IMP, error has a low-error plateau, a power-law region, and a high-error plateau. Between these regions are *transitions* where error varies smoothly from one region to the next. Matching the shape of these transitions was a key consideration for selecting our function family. To illustrate the importance of properly fitting the transitions, Figure 6 shows two possible functional families for fitting the relationship between density and error for the CIFAR-10 ResNets. Actual error is in black, and the functional form from Section 3 is in blue. In red is the fit for a functional form adapted from the one that Rosenfeld et al. (2020) use to model the relationship between width and error. The difference between these functional families is the way they model transitions, and the one we choose in this paper better models the transitions in our setting. For further discussion of this comparison, see Appendix G.

**Criterion 2: Few, interpretable parameters.** Selecting a functional form is not merely a curve-fitting exercise. We seek the underlying structure that governs the relationships between \(d, l, w, n\), and error in a manner akin to a law of physics. As such, our functional form should have a small number of parameters that are *interpretable*. In our functional form (Eq. 2), each parameter has a clear meaning. The parameters \(\epsilon^\ast, p', \gamma\) control the high-error plateau, the transition to the power-law region, and the slope of the power-law region. \(\phi\) and \(\psi\) control the interchangeability of width and depth with density. We approximate error over multiple orders of magnitude and 4,301 configurations of ResNet-20 on CIFAR-10 with just five parameters, indicating we have distilled key information about the behavior of pruning into our functional form.

**Sources of systemic error and limitations.** By seeking to minimize the number of parameters in our functional form, we leave some phenomena unmodeled. In particular, there are two phenomena we have chosen not to model that introduce systemic error. First, the low-error plateau is not a plateau. Error often improves slightly at high densities before returning to \(\epsilon_{np}\) during the transition to the power-law region. Our model treats the region as flat and treats error as monotonically increasing as density decreases. This source of error accounts for a bias of \(\approx 1\%\) relative error in our estimation (Appendix H). Second, we model both transitions (between the power-law region and each plateau) with a single shape and the same transition rate. If we treated them separately and used higher-order terms in the rational form, we could potentially reduce some of the residual error in our estimation at the cost of additional complexity.

7. Implications and Conclusions

Our main contribution is a functional form \(\hat{\epsilon}(\epsilon_{np}, d, l, w, n)\) that accurately predicts the error when pruning members of a network family using IMP. There are several broader implications of our ability to characterize pruning in this way. The mere existence of this functional form means there is indeed structure to the way pruning affects error. Although prior work (Cai et al., 2019) has implicitly relied on the existence of structure for a different pruning method, we are the first to explicitly describe such structure. This
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Figure 7. Estimated error from our scaling law (blue) and the scaling law adapted from Rosenfeld et al. (2020) (red) for CIFAR-10 ResNet as \( w \) varies. Actual error is in black. Our scaling law better captures transitions between the low-error plateau and the power-law region.

Figure 8. Estimated error as width varies for the CIFAR-10 ResNets (left). Actual error as width varies for the CIFAR-10 ResNets (right). The dotted black line is the minimal number of parameters necessary to reach each error \( \epsilon_k \) among all of the pruned networks. Reaching this point requires starting with a particular lower-error network (purple) and pruning until error increases to \( \epsilon_k \). Starting too large (pink) will miss this point.

functional form enables a framework in which we can reason conceptually and analytically about pruning. In doing so, we can make new observations about pruning that are non-obvious or costly to exhaustively demonstrate empirically. For example, recall our motivating question:

*Given a family of neural networks, which should we prune (and by how much) to obtain the network with the smallest possible parameter-count such that its error does not exceed some threshold \( \epsilon_k \)?*

This is an increasingly common question as the community begins to think about pruning as a way of seeking the optimal pruned member of a family of networks rather than as a technique applied to a specific network in isolation. At its heart, the question is really an optimization problem: find the configuration of \( d, l, \) and \( w \) that minimizes parameter-count \( m \) subject to an error constraint: 

\[
\arg\min_{l,w,d} m \text{ s.t. } \epsilon = \epsilon_k.
\]

For ResNets, the parameter-count \( m \) is proportional to \((dw)^2\).\(^4\) Hence, this yields the following optimization problem:

\[
l, w, d = \arg\min_{l,w,d} lw^2d \quad \text{s.t.} \\
\epsilon_k \left\| \frac{1}{l^\gamma w^\gamma d - j} \left( \frac{e^\gamma}{\epsilon_k} \right) \right\| = \epsilon_k
\]

This optimization problem is solvable directly without running any further experiments.

Studying this optimization problem reveals a useful insight about—in this case—the CIFAR-10 ResNets. In the pruning literature, it is typical to report the minimum density where the pruned network matches the error \( \epsilon_{np}(l, w) \) of the unpruned network (Han et al., 2015). However, our scaling law suggests this is not the smallest model to achieve error \( \epsilon_{np}(l, w) \). Instead, it is better to train a larger network with depth \( l' \) and width \( w' \) and prune until error reaches \( \epsilon_{np}(l, w) \), despite the fact that error will be higher than \( \epsilon_{np}(l', w') \). This analytic result parallels and extends the findings of Li et al. (2020) on NLP tasks. However, unlike Li et al., our scaling law suggests starting too large is detrimental for the CIFAR-10 ResNets, leading to a higher parameter-count at error \( \epsilon_k \).

Figure 8 (left) illustrates this behavior concretely: it shows the error predicted by our scaling law for CIFAR-10 ResNets with varying widths. The dotted black line shows the minimal parameter-count at which we predict it is possible to achieve each error. Importantly, none of the low-error plateaus intersect this black dotted line, meaning a model cannot be minimal until it has been pruned to the point where it increases in error. This occurs because the transitions of our functional form are gradual. On the other hand, if we start with a model that is too large, it will no longer be on the black line when it has been pruned to the point where its error reaches \( \epsilon_{np}(l, w) \); this behavior occurs because error decreases as a function of the invariant \( m^* \) rather than the parameter-count \( m \) and because \( m \propto m^* \).

In Figure 8 (right), we plot the same information from the actual CIFAR-10 data and see the same phenomena occur in practice. The difference between the estimated and actual optimal parameter count is no more than 25%.

Looking ahead, there are many directions for future work. Further studying sources of systematic error (transition shape and error improvements on the low-error plateau) is a promising avenue for making it possible to extrapolate from small-scale settings to large-scale settings (see Appendix F for a forward-looking discussion). Furthermore, while we focus on CIFAR-10 and ImageNet ResNets in the
main body, it is important to understand the generality of our functional form for other networks and tasks (see Appendix E). Finally, now that we have described the structure of the error of IMP-pruned networks, it will be valuable to study the nature of scaling laws that capture the behavior of the plethora of other pruning methods that achieve different tradeoffs between parameter-count and error.

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Appendix H. How we computed the effect of error dips on our estimator in Section 6.
A. Formal Statement of Iterative Magnitude Pruning

**Algorithm 1** Iterative Magnitude Pruning (IMP) with weight rewinding to epoch 10 and $N$ iterations.

1: Create a neural network with randomly initialized weights $W_0 \in \mathbb{R}^d$ and initial pruning mask $m = 1^d$
2: Train $W_0$ to epoch 10, resulting in weights $W_{10}$
3: for $n \in \{1, \ldots, N\}$ do
4:    Train $m \odot W_{10}$ (the element-wise product of $m$ and $W_{10}$) to final epoch $T$ and weights $m \odot W_{T,n}$
5:    Prune the 20% of weights in $m \odot W_{T,n}$ with the lowest magnitudes. $m[i] = 0$ if $W_{T,n}[i]$ is pruned
6: Return $m$ and $W_{T,n}$
B. Experimental Details

B.1. ResNets

We study the residual networks (ResNets) designed by He et al. (2016) for CIFAR-10 and ImageNet. ResNets for CIFAR-10 are composed of an initial convolutional layer, three sets of $B$ residual blocks (each with two convolutional layers and a skip connection), and a linear output layer. The sets of blocks have 16, 32, and 64 convolutional channels, respectively.

ResNets for ImageNet and TinyImageNet are composed of an initial convolutional layer, a max-pooling layer, four sets of residual blocks (each with three convolutional layers and a skip connection), and a linear output layer. The sets of blocks have 64, 128, 256, and 512 convolutional channels, respectively. On ImageNet, we use a ResNet with 50 layers. On TinyImageNet, we use a ResNet with 18 layers. Both choices are standard for these datasets.

We place batch normalization before the ReLU activations.

To vary the width of the networks, we multiply the number of convolutional channels by the width scaling factor $w$. To vary the depth of the CIFAR-10 ResNets, we vary the value of $B$. The depth $l$ of the network is the total number of the layers in the network, not counting skip connections.

B.2. VGG Networks

We study the VGG-16 variant of the VGG networks for CIFAR-10 as provided by the OpenLTH repository. The network is divided into five sections, each of which is followed by max pooling with kernel size 2 and stride 2. The sections contain 3x3 convolutional layers arranged as follows:

| Section | Width | Layers |
|---------|-------|--------|
| 1       | 64    | 2      |
| 2       | 128   | 2      |
| 3       | 256   | 3      |
| 4       | 512   | 3      |
| 5       | 512   | 3      |

The network has ReLU activations and batch normalization before each activation. To vary the width of VGG-16, we multiply each of the per-segment widths by the width scaling factor $w$.

B.3. DenseNets

We study the densely connected residual networks (DenseNets) designed by He et al. (2016) for CIFAR-10. DenseNets for CIFAR-10 are composed of an initial convolutional layer, four sets of dense blocks, and a linear output layer. Between the sets of blocks are transition layers of 1x1 convolutions and an average pooling operation that downsamples the image by 2x. Each block comprises a 1x1 convolution that increases the channel count by $4x$ and a 3x3 block that decreases it to a fixed constant size $g$; this output is then concatenated to the input of the block. As such, if the input to the block has $n$ channels, the output of the block has $n + g$ channels. We use DenseNet-121, which has sets of 6, 12, 24, and 16 blocks. $g$ is set to 16 but is multiplied by the width scaling factor $w$ to modify the width.

B.4. Training Hyperparameters

We train CIFAR-10 and SVHN ResNets and VGG-16 for 160 epochs with a batch size of 128. The initial learning rate is 0.1, and it drops by an order of magnitude at epochs 80 and 120. We optimize using SGD with momentum (0.9). We initialize with He uniform initialization. CIFAR-10 data is augmented by normalizing, randomly flipping left and right, and randomly shifting by up to four pixels in any direction (and cropping afterwards). SVHN data is not augmented.

We train CIFAR-10 DenseNets with the same hyperparameters but for 200 epochs (with learning rate drops at 130 and 165 epochs). We train SVHN DenseNets with the same hyperparameters but for 100 epochs (with learning rate drops at 70 and 85 epochs).

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10github.com/facebookresearch/open_lth
We train ImageNet ResNets for 90 epochs with a batch size of 1024. The initial learning rate is 0.4, and it drops by an order of magnitude at epochs 30, 60, and 80. We perform linear learning rate warmup from 0 to 0.4 over the first 5 epochs. We optimize using SGD with momentum (0.9). We initialize with He uniform initialization. Data is augmented by normalizing, randomly flipping left and right, selecting a random aspect ratio between 0.8 and 1.25, selecting a random scaling factor between 0.1 and 1.0, and cropping accordingly.

We train TinyImageNet ResNets identically except for that we train them for 200 epochs (with learning rate drops at 100 and 150 epochs) and a learning rate of 0.2. Augmentation is identical to ImageNet.

### B.5. Dimensions

We use the following dimensions for the additional experiments. We select configurations using the same methodology as in Table 1.

| Network Family          | Densities ($d$) | Depths ($l$)       | Width Scalings ($w$) | Subsample Sizes ($n$) |
|-------------------------|-----------------|--------------------|----------------------|------------------------|
| CIFAR-10/SVHN ResNet    | $0.8^i, i \subseteq \{0, \ldots, 40\}$ | $l \subseteq \{8, 14, 20, 26, 50, 98\}$ | $2^i, i \subseteq \{-4, \ldots, 2\}$ | $\frac{c}{n}, i \in \{1, 2, 4, 8, 16, 32, 64\}$ |
| CIFAR-10/SVHN VGG       | $0.8^i, i \subseteq \{0, \ldots, 37\}$ | 16                  | $2^i, i \subseteq \{-4, \ldots, 0\}$ | $\frac{c}{n}, i \in \{1\}$ |
| CIFAR-10/SVHN DenseNet  | $0.8^i, i \subseteq \{0, \ldots, 50\}$ | 121                 | $2^i, i \subseteq \{-4, \ldots, 1\}$ | $\frac{c}{n}, i \in \{1\}$ |
| TinyImageNet ResNet     | $0.8^i, i \subseteq \{0, \ldots, 50\}$ | 18                  | $2^i, i \subseteq \{-6, \ldots, 0\}$ | $\frac{c}{n}, i \in \{1\}$ |
| ImageNet ResNet         | $0.8^i, i \subseteq \{0, \ldots, 30\}$ | 50                  | $2^i, i \subseteq \{-4, \ldots, 0\}$ | $\frac{c}{n}, i \in \{1, 2, 4\}$ |
In this appendix, we show that our observations from Section 3 hold when varying all dimensions (depth, width, and dataset size) on both the CIFAR-10 and ImageNet ResNets for IMP. Figure 9 shows the error versus density when changing width (left) depth (center) and data (right). In Figure 10, we similarly show the dependency of the error on density for Imagenet when varying width (left) and dataset size (right).

In Figure 9, we observe that all curves have a similar slope in the power-law region. In Equation 1, this implies that while $\gamma$ is allowed to vary with $l, w$ and $n$, it is in practice approximately a constant. Similarly, the high-error plateau $\epsilon^\uparrow$ is also shared across curves such that it too is approximately constant. In contrast, the transition from high-error plateau to the power-law region is not constant as a function of density. Section 4 finds exactly this dependency of the transition parameter $p$.

**Figure 9.** Relationship between density and error when pruning CIFAR-10 ResNets and varying $w$ (left, $l = 20, n = N$), $l$ (center, $w = 1, n = N$), $n$ (right, $l = 20, w = 1$)

**Figure 10.** Relationship between density and error when pruning ImageNet ResNet-50 and varying $w$ (left, $n = N$), and $n$ (right, $w = 1$)
D. Partial (Projections) Fit Results for Section 4

In Section 4, we fit the error jointly as a function of all dimensions showing that Equation 2 provides a good approximation to the error in practice. In this appendix, we consider important sub-cases, such as the case when one wishes to scale only one degree of freedom while pruning. This serves both a practical scenario, but also allows for a qualitative visualization of the fit (and typical sources of error), which is otherwise difficult to perform over all dimensions jointly. From a practical standpoint, in this case one need not estimate the parameters associated with the fixed degree of freedom.

Recall that, given the non-pruned network error $\epsilon_{np}$, all dependencies on the individual structural degrees of freedom $l, w$ are captured by the invariant $m^* \triangleq l^\phi w^\psi d$. This means that, if one wishes to estimate the error while pruning when holding width fixed, we need not estimate $\psi$. Similarly if depth is held constant, we need not estimate $\phi$.

Figure 11 shows these partial fits. Shown from left to right are the fits done while pruning and varying width, depth and data respectively. Correspondingly, these fits omit separately $\psi$ or $\phi$ or omit both when depth nor width are scaled. The fits were performed with all available density points for each dimension. For CIFAR-10: 7 widths, 224 points for the width partial fit; 7 dataset fractions, 240 points for the data partial fit; 4 depths, 164 points for the depth partial fit. For ImageNet: 5 widths, 83 points for the width partial fit; 3 dataset fractions, 86 points for the data partial fit.

This exercise, apart from its practical implications, highlights the fact that there are in effect two groups of parameters comprising the estimation. The first are the parameters $\epsilon^l, \gamma$ and $\rho^l$ which control the dependency as a function of density (or more generally, as a function of the invariant). The second are $\phi$ and $\psi$ which are properties of the architectural degrees of freedom captured by the invariant. Moreover, within the first group of parameters $\epsilon^l, \gamma$, can be isolated and found from a single pruning curve, as they are not a function of $l, w, n$.

![Figure 11](image-url)

*Figure 11.* Top row: CIFAR-10. Bottom row: ImageNet. Left: varying width. Center: varying dataset size. Right: varying depth. Lines are the actual error and dots are the estimated error.
E. Additional Architectures and Datasets

In this appendix, we show that our functional form applies to additional pairs of networks and datasets: (CIFAR-10 ResNet, SVHN), (VGG-16, CIFAR-10), (VGG-16, SVHN), (DenseNet-121, CIFAR-10), (DenseNet-121, SVHN), (ImageNet ResNet-18, TinyImageNet).

In general, we obtain good fits on CIFAR-10 and TinyImageNet. On SVHN, fits are worse but the networks suffer from high measurement error (i.e., accuracy varies greatly between multiple runs at the same density) at low densities; nevertheless, fits are often better than measurement error because they average out some of the error.

We add these additional comparisons in the following Figures:

- Figure 12: ResNet-20 on SVHN with IMP as width varies. $|\mu| < 2\%, \sigma < 8\%$. Notably, the measurement error in this case is large ($\sigma \sim 9.5\%$), dominating (over the approximation error) the total fit error. The fit averages out some of this error, resulting in a fit error which is lower than the measurement error. In general, experiments on SVHN are quite noisy, leading to significant measurement error.

- Figure 13: VGG-16 on CIFAR-10 with IMP as width varies. $|\mu| < 3\%, \sigma < 7\%$ (compared to measurement error 12%).

- Figure 14: VGG-16 on SVHN with IMP as width varies. $|\mu| < 4\%, \sigma < 15\%$ (compared to measurement error 20%—measurement error is large at very low densities).

- Figure 15: DenseNet-121 on CIFAR-10 with IMP as width varies. $|\mu| < 1\%, \sigma < 8\%$. Bias is evident in the transition as we discuss in Section 6.

- Figure 16: DenseNet-121 on SVHN with IMP as width varies. $|\mu| < 4\%, \sigma < 19\%$ (compared to measurement error 16%—measurement error is large at very low densities).

- Figure 17: ResNet-18 on TinyImageNet with IMP as width varies. $|\mu| < 1\%, \sigma < 1.3\%$ (compared to measurement error 1%).

![Figure 12. Fit for ResNet-20 on SVHN with IMP pruning.](image)
On the Predictability of Pruning Across Scales

Figure 13. Fit for VGG-16 on CIFAR-10 with IMP pruning.

Figure 14. Fit for VGG-16 on SVHN with IMP pruning.

Figure 15. Fit for DenseNet-121 on CIFAR-10 with IMP pruning.
Figure 16. Fit for DenseNet-121 on SVHN with IMP pruning.

Figure 17. Fit for ResNet-18 on TinyImageNet with IMP pruning.
F. Towards Extrapolation

**Background.** In the main body, we showed that our scaling law accurately fits the error of pruned neural networks. As, such it has predictive power, allowing us to reason in a principled manner about pruning trade-offs. Similarly, it allows to make predictions about what would happen at larger model and data scales than explored here. Importantly, only a few experiments need be performed to find the coefficients for the scaling law (see Appendix 5).

However, we could ask, how accurately can we estimate the scaling law parameters from even smaller scales? That is, is it possible to fit our scaling law to data from networks with deliberately smaller depths, widths, and dataset sizes and accurately predict the error of larger-scale models? If so, we could make informed decisions about pruning large-scale models through small-scale experiments alone, saving the costs associated with large scale training and pruning.

Outside the context of pruning, the scaling laws of (Rosenfeld et al., 2020) (for both language models and image classification) and (Kaplan et al., 2020) (for predicting the expected performance of GPT-3 (Brown et al., 2020) at very large scale) have been shown to extrapolate successfully in this manner.

**Results on CIFAR-10.** In Figure 18, we show the result of extrapolating from small-scale networks on CIFAR-10 \((w = \frac{1}{8}, \frac{1}{4}; l = 14, 20)\) to all widths and depths on CIFAR-10. Extrapolation prediction is still accurate: \(\mu < 7\%, \sigma < 6\%\) (vs. \(\mu < 1\%, \sigma < 6\%\) in the main body).

**Future work.** However, extrapolation is particularly sensitive to systemic errors. Specifically, the transitions and the error dips can lead to large deviations when extrapolating. For ImageNet, the error dips (especially on small dataset sizes) are especially pronounced, preventing stable extrapolation. In order to improve extrapolation performance, future work should explore the challenges we discuss in Section 6: approaches to either model or mitigate these dips and to improve the fit of the transitions.

*Figure 18.* Extrapolation results from four pruned networks on CIFAR10 \(w = \frac{1}{8}, \frac{1}{4}; l = 14, 20\) to all larger networks \((n = 1)\). Fit results are in blue, extrapolation in red, actual in black. Error versus number of non-zero parameters (left). Estimated versus actual errors (right).
G. Comparison of Pruning and Non-pruning Scaling Laws

In this appendix, we contrast the behavior of the error when pruning with the behavior of the error in the non-pruning setting. Hestness et al. (2017) show the the error follows a saturating power-law form when scaling data (with both low and high-error plateaus) but does not model them. Rosenfeld et al. (2020) unify the dependency on data and model size while approximating the transitions between regions; they propose the following form:

\[ \tilde{\epsilon}(m, n) = a n^{-\alpha} + b m^{-\beta} + c_{\infty} \]  

(3)

\[ \hat{\epsilon}(m, n) = \epsilon_0 \| \tilde{\epsilon}(m, n) \| \]  

(4)

where \( m \) is the total number of parameters and \( n \) is the dataset size. \( a, b, \alpha, \beta, c_{\infty} \), and \( \eta \) are constants, and \( \epsilon_0 \) plays the role of \( \epsilon_{\uparrow} \) in our notation.

Rosenfeld et al. model the upper transition—from power-law region to the high-error plateau—by a rational form in a fashion similar to the approach we take. The key difference is that we consider a power of the polynomials in the numerator and denominator of the rational form, where in Eq. 3 the power is hidden in the term \( \tilde{\epsilon} \).

The biggest difference arises when considering the lower transition (between the low-error plateau and the power-law region). This transition is captured by Eq. 3. Considering either the width or depth degrees of freedom \( x \in \{ w, l \} \), Eq. 3 can be re-written as:

\[ \tilde{\epsilon}(x) = b_x x^{-\beta_x} + c_x \]  

(5)

Where \( b_x \) and \( \beta_x \) are constants and \( c_x \) is a constant as a function of \( x \) (it is only a function of the data size \( n \)).

Figure 19 (right) shows the error versus depth for different dataset sizes. In grey is the actual error, while in red is the best fit when approximating the error by Eq. 5. Qualitatively, one sees that the fit using Eq. 5 does indeed closely match the error in practice.

Recall that we are interested in comparing the errors as a function of the density. A requirement from any functional form used to model the dependency on the density is to degenerate to the error of the non pruned model \( \epsilon_{np} \) at \( d = 1 \). We adapt Eq. 5 by solving the relation between \( b_x \) and \( c_x \) meeting this constraint, to arrive at:

\[ \tilde{\epsilon}(x) = b_x x^{-\beta_x} + \epsilon_{np} - b_x \]  

(6)

Contrast Eq. 5 with the functional form we propose in Eq. 1, re-written here for convenience:

\[ \hat{\epsilon}(d, \epsilon_{np}, l, w, n) = \epsilon_{np} \left( \frac{d - j p \left( \frac{\epsilon_{np}}{\epsilon_{\uparrow} \epsilon_{np}} \right)^{\frac{1}{2}}} {d - j p} \right)^\gamma \]  

where \( j = \sqrt{-1} \)  

(7)

This can be simplified to capture only the lower transition—far enough from the upper transition \( (d \gg p) \)—to:

\[ \hat{\epsilon}(d, \epsilon_{np}, l, w, n) = \epsilon_{np} \left( \frac{d - j p \left( \frac{\epsilon_{np}}{\epsilon_{\uparrow} \epsilon_{np}} \right)^{\frac{1}{2}}} {d} \right)^\gamma \]  

(8)

Figure 19 (left) shows error versus density for different widths. In blue is the fit with Eq. 8 which follows closely the actual error (black) while in red is the fit with Eq. 6 which deviates noticeably in comparison.
We have seen that in practice that the form of Eq. 6 does not match well the pruning case, where the mismatch originates from lower transition shape. We have thus reached a phenomenological observation distinguishing the pruning and non-pruning forms; we leave the study of the origins of this phenomenon for future work.
H. The effect of error dips on estimation bias

In this appendix, we consider the effect of the error dips on our estimator as discussed in Section 4. As we mention in that section, when pruning a network, error often dips below $\epsilon_{np}$ during the low-error plateau.

Recall that we find the parameters in our estimator (Equation 2) by minimizing the MSE of relative error $\delta$. Our estimation has bias if $\mathbb{E}(\hat{\epsilon} - \epsilon) \neq 0$ where the expectation is over all model and data configurations. Equivalently, the relative bias is $\mu \triangleq \mathbb{E}\delta = 0$ iff the estimator is unbiased. The Estimator captured by the joint form in Equation 2 is a monotonically increasing function of the density. It is also constrained such that at density $d = 1$ it is equal to the non-pruned error $\epsilon_{np}$. It thus, can not reduce The MSE to zero, as it can not decrease to match the actual error dips. This results in the bias of the relative error $\mu$ which in practice is $\sim 1\%$. 