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# Deep Learning on a Data Diet: Finding Important Examples Early in Training

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## Abstract

The recent success of deep learning has partially been driven by training increasingly overparametrized networks on ever larger datasets. It is therefore natural to ask: how much of the data is superfluous, which examples are important for generalization, and how do we find them? In this work, we make the striking observation that, on standard vision benchmarks, the initial loss gradient norm of individual training examples, averaged over several weight initializations, can be used to identify a smaller set of training data that is important for generalization. Furthermore, after only a few epochs of training, the information in gradient norms is reflected in the normed error-L2 distance between the predicted probabilities and one hot labels—which can be used to prune a significant fraction of the dataset without sacrificing test accuracy. Based on this, we propose data pruning methods which use only *local information early in training*, and connect them to recent work that prunes data by discarding examples that are rarely forgotten *over the course of training*. Our methods also shed light on how the underlying data distribution shapes the training dynamics: they rank examples based on their importance for generalization, detect noisy examples and identify subspaces of the model’s data representation that are relatively stable over training.

## 1 Introduction

Recently, deep learning has made remarkable progress driven, in part, by training overparameterized models on ever larger datasets. This trend creates new challenges: the large computational resources required pose a roadblock to the democratization of AI. Memory and resource constrained settings, such as on-device computing, require smaller models and datasets. Identifying important training data plays a role in online and active learning. Finally, it is of theoretical interest to understand how individual examples and sub-populations of training examples influence learning.

To address these challenges, we propose a scoring method that can be used to identify important and difficult examples early in training, and prune the training dataset without large sacrifices in test accuracy. We also investigate how different sub-populations of the training data identified by our score affect the loss surface and training dynamics of the model.

Recent work on pruning data [1, 2], can be placed in the broader context of identifying coresets that allow training to approximately the same accuracy as would be possible with the original data [3–7]. These works attempt to identify examples that provably guarantee a small gap in training error on the full dataset. However, due to the nonconvex nature of deep learning, these techniques make conservative estimates that lead to weak theoretical guarantees and are less effective in practice.

A very different approach was recently discovered by Toneva et al. [8]. They track the number of times through training an example transitions from being correctly classified to misclassified, called a “forgetting event”, and find that some examples are rarely forgotten, while others are forgotten repeatedly. Empirically, they observed that training accuracy is not affected by the rarely forgotten training examples and a large fraction of the training data can be removed without any impact on test accuracy. However, since this method relies on collecting forgetting statistics throughout training, the forgetting score is typically calculated in the middle of or at the end of training. Toneva et al. [8] find that, in their example of a ResNet18 trained on CIFAR-10 for 200 epochs, the Spearman rank correlation between early and late scores is good after about 25 epochs and stabilizes after 75 epochs.

Broadly speaking, the ability to prune datasets raises a number of questions: What is the nature of examples that can be removed from the training data without hurting accuracy? How early in training can we recognize such examples? How many examples do we need and how does this depend on the data distribution? These questions may have no generic answers and so, in this work, we begin to pursue them empirically in the context of several standard vision benchmarks and standard network architectures. Answers to these questions may both (1) lead to new methodologies that could dramatically reduce training times and memory requirements, and (2) offer important insights into the training dynamics of deep neural networks, and the role of data.

Our first finding is that *very early in training* (just a few epochs), partial forgetting scores identify large fractions of data that can be pruned. Analyzing this puzzling result with a one gradient step analysis of training suggests a very simple heuristic: use the loss gradient norm of individual examples to identify important examples. While this approach does not work when the loss gradient norms are computed at the weights early in training of a single trajectory, we find that, surprisingly, averaging these norms over multiple weight initializations does produce a ranking that correlates strongly with forgetting scores and allows us to prune a significant fraction of examples early in training. Indeed, even at initialization, we can prune 50% of examples from CIFAR-10 without affecting accuracy, while on the more challenging CIFAR-100 dataset, we can prune 25% of examples with only a 1% drop in accuracy.

Through a series of empirical studies, we have begun to tease apart the properties of important examples and how they can depend on the data distribution. In particular, we find that the examples with the very highest norms become superfluous as the amount of label noise increases. Indeed, even on clean data, we find that in the high pruning regime, the best population excludes the very highest-scoring examples.

### 1.1 Contributions

- We propose to score the importance of each training example  $(x_i, y_i)$  by its expected loss gradient norm (**GraNd score**), which, up to a constant, bounds the expected change in loss for an arbitrary example  $(x, y)$  caused by removing  $(x_i, y_i)$ .
- We show that **pruning training samples with small GraNd scores** at initialization allows one to train on as little as 50% of the training data without any loss in accuracy (CIFAR-10). While the pruning levels are comparable to those provided by other methods [1, 8], our score is the only one that is well-defined at initialization and early in training.
- Our experimental findings suggest that, **within the first few epochs of training, the GraNd score is well-approximated by the norm of the error vector (EL2N score)**, where the error vector is the predicted class probabilities minus one-hot label encoding. In fact, we find that the EL2N score provides even better information for data-pruning across a wide range of data pruning levels, even early in training.
- We study the role of examples with the highest EL2N scores, and find that excluding a small subset of the **very highest scoring examples produces a boost in performance**. This boost in performance is enhanced in a corrupted label regime.
- We introduce a method, based on **linearly connected modes**, for studying the empirical risk surface in terms of the modes of **subsets of data**, allowing us to identify when, in training, the final performance on subpopulations is determined. We demonstrate that the linearly connected mode at-convergence of empirical risk surface **computed on low EL2N score** examples is determined much earlier in training compared to high score examples.

- Finally, we study how an example’s EL2N score connects to the network’s training dynamics. We do so by tracking the data-dependent NTK submatrices corresponding to the low or high score examples, and measuring the rate at which it evolves in a scale-invariant way. We find that the NTK submatrix for the high score examples evolves faster throughout training, supporting our hypothesis that high-scoring examples are the ones driving the learning and the changes in the NTK feature space [9].

## 2 Which samples are important for learning?

### 2.1 Preliminaries

We consider supervised classification, where  $S = \{(x_i, y_i)\}_{i=1}^N$  denotes the training set, drawn i.i.d. from an unknown data distribution  $\mathcal{D}$ , with input vectors  $x \in \mathbb{R}^d$  and one-hot vectors  $y \in \{0, 1\}^K$  encoding labels. For a fixed neural network architecture, let  $f_{\mathbf{w}}(x) \in \mathbb{R}^K$  be the logit outputs of the neural network with weights  $\mathbf{w} \in \mathcal{W} \subseteq \mathbb{R}^D$  on input  $x \in \mathbb{R}^d$ . Let  $\sigma$  be the softmax function given by  $\sigma(z_1, \dots, z_K)_k = \exp\{z_k\} / \sum_{k'=1}^K \exp\{z_{k'}\}$ . Let  $p(\mathbf{w}, x) = \sigma(f(\mathbf{w}, x))$  denote the neural network output in the form of a probability vector. For any probability vector  $\hat{p}$ , let  $\ell(\hat{p}, y) = \sum_{k=1}^K y^{(k)} \log \hat{p}^{(k)}$  denote cross-entropy loss.

Let  $\mathbf{w}_0, \mathbf{w}_1, \mathbf{w}_2, \dots, \mathbf{w}_T$  be the iterates of stochastic gradient descent (SGD), where, for some sequence of minibatches  $S_0, S_1, \dots, S_{T-1} \subseteq S$  of size  $M$ , we have

$$\mathbf{w}_t = \mathbf{w}_{t-1} - \eta \sum_{(x,y) \in S_{t-1}} g_{t-1}(x, y), \quad (1)$$

for  $g_{t-1}(x, y) = \nabla_{\mathbf{w}_{t-1}} \ell(p(\mathbf{w}_{t-1}, x), y)$ , and  $t = 1, \dots, T$ .

### 2.2 Gradient Norm Score and an infinitesimal analysis

Fix a training set  $S$ . Due to training with SGD from a random initialization, the weight vector at time  $t > 0$ ,  $\mathbf{w}_t$ , is a random variable. The expected magnitude of the loss vector is our primary focus:

**Definition 2.1.** The GraNd score of a training example  $(x, y)$  at time  $t$  is  $\chi_t(x, y) = \mathbb{E}_{\mathbf{w}_t} \|g_t(x, y)\|_2$ .

Here we describe conditions under which the GraNd score controls the contribution of a training example to the change in the training loss. In order to simplify our analysis, we approximate the training dynamics as if they were in continuous time.

A key quantity in our analysis is the time derivative of the loss for a generic labeled example  $(x, y)$ :  $\Delta_t((x, y), S_t) = -\frac{d\ell(f_t(x), y)}{dt}$  (where  $f_t(\cdot) = f_{\mathbf{w}_t}(\cdot)$ ), i.e., the instantaneous rate of change in the loss on  $(x, y)$  at time  $t$ , where the gradient is computed on the minibatch  $S_t$ . By the chain rule,

$$\Delta_t((x, y), S_t) = g_t(x, y) \frac{d\mathbf{w}_t}{dt}. \quad (2)$$

This relates to our discrete time dynamics via  $\frac{d\mathbf{w}_t}{dt} \approx \mathbf{w}_{t+1} - \mathbf{w}_t = -\eta \sum_{(x', y') \in S_{t-1}} g_{t-1}(x', y')$ .

Our goal is to understand how removing a training point from minibatch  $S_t$  affects  $\Delta_t((x, y), S_t)$ .

**Lemma 2.2.** Let  $S_{\neg j} = S \setminus (x_j, y_j)$ . Then for all  $(x, y)$ , there exists  $c$  such that

$$\|\Delta_t((x, y), S) - \Delta_t((x, y), S_{\neg j})\| \leq c \|g_t(x_j, y_j)\|. \quad (3)$$

*Proof.* For a given example  $x$ , the chain rule yields  $\Delta_t((x, y), S) = -\frac{d\ell(f_t(x), y)}{dt} = \frac{d\ell(f_t(x), y)}{d\mathbf{w}_t} \frac{d\mathbf{w}_t}{dt}$ . Since the weights are updated using SGD, we have  $\frac{d\mathbf{w}_t}{dt} = -\eta \sum_{(x_j, y_j) \in S_t} g_t(x_j, y_j)$ . Letting  $c = \eta \left\| \frac{d\ell(f_t(x), y)}{d\mathbf{w}_t} \right\|$ , the result follows.  $\square$

At any given training step, given the current location  $\mathbf{w}_t$ , the contribution of a training example  $(x, y)$  to the decrease of loss on any other example, is bounded by Eq. (3). Since the constant  $c$  does not depend on  $(x, y)$ , we only consider the gradient norm term,  $\|g_t(x, y)\|$ . The expected value of this gradient norm is exactly the GraNd score of  $(x, y)$ . In other words, examples with a small

GraNd score in expectation have a bounded influence on learning how to classify the rest of the training data at a given training time<sup>1</sup>. We therefore propose to rank training examples by their GraNd scores, larger norm meaning more important for maintaining  $\Delta_t(x)$ .

For an arbitrary input  $x \in \mathbb{R}^d$ , let  $\psi_t^{(k)}(x) = \nabla_{\mathbf{w}_t} f_t^{(k)}(x)$  denote the  $k$ th logit gradient. Then GraNd can be written as

$$\chi_t(x, y) = \mathbb{E} \left\| \sum_{k=1}^K \nabla_{f_t^{(k)}} \ell(f_t(x), y)^T \psi_t^{(k)}(x) \right\|_2. \quad (4)$$

Under the cross entropy loss,  $\nabla_{f_t^{(k)}} \ell(f_t(x), y)^T = p(\mathbf{w}_t, x)^{(k)} - y_k$ . When  $\{\psi_t^{(k)}(x)\}_k$  are roughly orthogonal across logits, and are of a similar size across logits and training examples  $x$ , then we can approximate GraNd by just the norm of the error vector.

**Definition 2.3.** The EL2N score of a training sample  $(x, y)$  is defined to be  $\mathbb{E}\|p(\mathbf{w}_t, x) - y\|_2$ .

Our experimental results suggest that this approximation becomes accurate after a few epochs of training (see Section 3).

### 2.3 Comparison to forgetting scores

Toneva et al. [8] define a “forgetting event” for a training sample to be a point in training when the classifier switches from making a correct classification decision to an incorrect one. They define an approximate *forgetting score* for each training example as the number of times during training when it was included in a minibatch *and* underwent a forgetting event. Toneva et al. demonstrate that examples with low forgetting score may be completely omitted during training without any noticeable effect on the accuracy of the learned predictor. In Fig. 1 and Appendix D.3, we make an empirical comparison of forgetting scores to our proposed GraNd and EL2N scores.

In Lemma 2.2, we bounded the contribution of a training example to the decrease of the loss of any other sample over a single gradient step. Due to  $\psi_t(\cdot)$ ’s being time-dependent, it is complicated to extend the analysis to multiple steps. However, it is interesting to consider a case when  $\psi_t(x_i) = \psi(x_i)$  for all  $x_i$  in the training set, and  $K = 1$ . Then summing the bound in Eq. (3) on how much a sample  $(x_j, y_j)$  affects the logit output on an arbitrary point at each time  $t \in \{1, \dots, T\}$ , we obtain a score that depends on  $\|\psi(x_j)\| \|\sum_t (p_t(x_j) - y_j)\|$ . For two examples,  $(x, y)$  and  $(x', y')$ , such that  $\|\psi(x')\| \approx \|\psi(x)\|$ , we see that the example that is learned faster and maintains small error over training time will have a smaller GraNd score on average throughout training. Note that  $|(p_t(x_j) - y_j)|$ , if rescaled, is an upper bound on 0–1 loss, and therefore  $\sum_t |(p_t(x_j) - y_j)|$  upper bounds the number of forgetting events during training (after rescaling). In this simplified setting an example with a high number of forgetting events will also have a high GraNd score.

## 3 Empirical Evaluation of GraNd and EL2N Scores via Data Pruning

In the previous section, we motivated GraNd and EL2N scores by quantifying the influence of a training example on the loss of an arbitrary example after one optimization step. In this section, we evaluate these scores empirically, and verify that they identify examples important for generalization. Networks trained on subsets of the data with high scores achieve levels of test accuracy comparable to training on the full dataset and are competitive with other state of the art data pruning methods. Perhaps most remarkably, these scores are effective even when computed early in training and perform significantly better than a random baseline, even at initialization.

**Data pruning experiments.** We train convolutional neural networks of varying depth—ResNet18 and ResNet50 [10]—on standard vision datasets of varying difficulty—CIFAR-10, CIFAR-100 [11], and CINIC-10 [12]. All scores are calculated by averaging the scores from ten independent training runs. After calculating scores and selecting a training subset, final test accuracies are obtained by retraining networks from random initializations on only the selected subset. For each experiment, we report the mean of four independent runs and represent variability across runs by shading the region which spans the 16th to 84th percentile of obtained accuracies. See Appendix B for more implementation details and Appendix D for additional experiments.

<sup>1</sup>Note that the opposite is not necessarily true: examples with large scores may have gradients that cancel out and do not contribute much, meaning that this upper bound is loose.

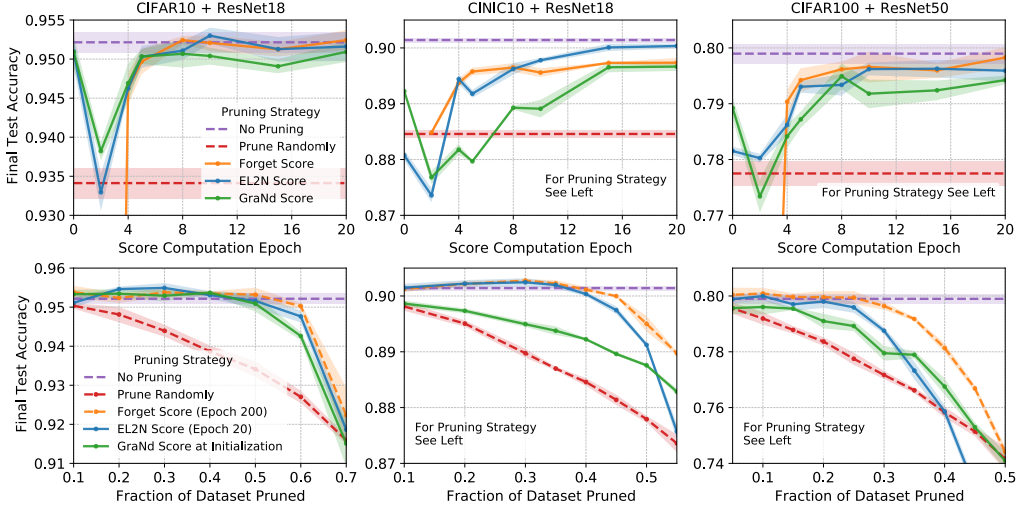


Figure 1: Columns correspond to three different dataset and network combinations (labeled at the top). *First row:* Final test accuracy achieved by training on a subset of training data comprised of examples with maximum forgetting, EL2N and GraNd scores computed at different times early in training. Subsets of a fixed size are used: networks are trained on 50% of training data for CIFAR-10, 60% for CINIC-10 and 75% for CIFAR-100. *Second row:* Final test accuracy achieved by training after different fractions of the dataset are pruned. Here we compare forgetting scores at the end of training, EL2N scores early in training (at epoch 20) and GraNd scores at initialization. In each case, examples with the lowest scores are pruned at initialization. *In all experiments* accuracies achieved by training on the full dataset and on a random subset of the corresponding size are used as baselines.

In Fig. 1, we show the results of two sets of experiments (top and bottom) on three different network and dataset combinations. The first experiment asks, how early in training are forgetting, GraNd and EL2N scores effective at identifying examples important for generalization? We compare the final test accuracy from training on subsets of fixed size but pruned based on scores computed at different times early in training. The second experiment compares how GraNd scores at initialization, EL2N scores early in training and forgetting scores at the end of training negotiate the trade-off between generalization performance and training set size. The training sets are constructed by pruning different fractions of the lowest score examples. In all examples, training on the full dataset and a random subset of the corresponding size are used as baselines. We make the following observations.

**Pruning at initialization.** In all settings, GraNd scores can be used to select a training subset *at initialization* that achieves test accuracy significantly better than random, and in some cases, competitive with training on all the data. This is remarkable because GraNd only contains information about the gradient norm at initialization, averaged over initializations. This suggests that the geometry of the training distribution induced by a random network contains a surprising amount of information about the structure of the classification problem. **EL2N scores**, which only contain information about errors, are not consistently effective at initialization and forgetting scores, which require counting forgetting events over training, are not defined at initialization.

**Pruning early in training.** We find that, after only a few epochs of training, EL2N scores are extremely effective at identifying important examples for generalization. For a wide range of intermediate pruning levels, training on the highest scores performs on par with or better than training on the full dataset. Even at higher pruning levels, **EL2N scores computed using local information early in training are competitive with forgetting scores which integrate information over the training trajectory.** This suggests that the average error vector *a few epochs into training* can identify examples that the network heavily uses to shape the decision boundary *throughout training*.

Interestingly, at extreme levels of pruning with either EL2N or GraNd scores, we observe a sharp drop in performance. We hypothesize that this is because at high levels of pruning, using either

GraNd or EL2N scores leads to bad coverage of the data distribution. By only focusing on the highest error examples, it is likely that an entire subpopulation of significant size that is present in the test data is now excluded from the training set. We only fit a small number of very difficult examples and do not keep enough of a variety of examples for training models with good test error.

**A property of the data.** Two results suggest that the ranking of important examples induced by EL2N and GraNd scores is a property of the dataset and not specific to a network. First, in Appendix D.2, we show that a ResNet18 and a ResNet50 trained on CIFAR-10 have similar performance curves and the same amount of data can be pruned, even though ResNet50 is a much deeper network with more parameters. Additionally, in an analysis of the sensitivity of the scoring methods to hyperparameters in Appendix D.1, we observe that scores calculated on a single network do not perform as well as those averaged across networks. We hypothesize that averaging the gradient or error norms over multiple initializations or training trajectories removes dependence on specific weights, allowing a more accurate distillation of the properties of the dataset.

In the following experiments, we focus on EL2N scores computed early in training, as they appear to more accurately identify important examples.

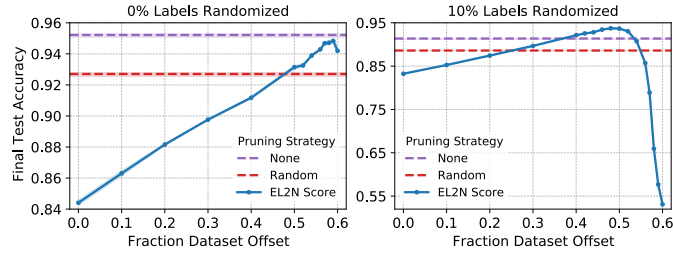
#### 4 Identifying noise examples

In the previous section, we studied the effect of keeping the highest-scoring examples, and found that we could train on only the top 50% of examples by score without a drop in accuracy (CIFAR-10). What is the nature of subpopulations of examples that allow us to reach high accuracy? One hypothesis is that the highest-scoring examples are the most important ones for achieving an accurate classifier. In this section, we refute this hypothesis, and demonstrate the role of label noise.

To test whether the highest-scoring examples are most important for achieving high accuracy, we first sort the examples by increasing EL2N score computed after a small number of training epochs.<sup>2</sup> Then we perform a sliding window analysis by training on a subset of examples with scores within a window from percentile  $f$  to percentile  $f + P$  percentile, always keep  $P\%$  of the data but sliding up  $f$ . As this window slides to higher percentiles, performance increases, except when the window includes examples with the very highest scores Fig. 2 (left). Indeed the optimal sliding window actually excludes approximately 500 of the highest-scoring training examples. These effects are reduced in the low pruning regime (see Appendix E.1). In Appendix C, we visualize some of the images that are excluded from each class.

Before we analyze these results, we first place them into a wider context, where we also change the amount of noise in the underlying label distribution. We repeat the experiment outlined above, but corrupt a random  $K\%$  of labels, replacing them with a random label, mirroring the protocol popularized by Zhang et al. [13]. Fig. 2 reveals that with increased label corruption, the optimal window shifts and excludes a higher number of examples. Therefore, the effect we see in the noiseless case appears to be magnified in the presence of label noise. Appendix E.2 examines how adding label noise influences the distribution of EL2N scores of examples.

These findings have several implications. The most obvious implication is that training with only the highest-scoring samples may not be optimal, especially when there is label noise. When the population has a low Bayes error rate, using only the highest scoring samples yields optimal results. However, without a validation set, one should be cautious in excluding high-score examples.



**Figure 2:** ResNet18 trained on a 40% subset of CIFAR-10 with clean (left) and 10% randomized labels (right). The training subset contains the *lowest* scoring examples *after* examples with scores below the offset are discarded. Scores computed at epoch 10.

<sup>2</sup>In Appendix E.3, we repeat these experiments for the GraNd score.

Feldman [14] discusses memorization in a noisy-label setup and gives conditions under which one should memorize in order to not misclassify singleton examples (examples in the training data that are the sole representatives of a subpopulation). For example, if the subpopulation appears with a frequency  $\Omega(1/N)$ , memorizing such examples can improve generalization. In practice, we may not know whether our data fits these conditions. However, our analysis in Fig. 2 suggests a simple and powerful method to prune data for optimal performance by optimizing just two hyperparameters of a sliding window using a validation set.

## 5 Optimization landscape and the training dynamics

### 5.1 Evolution of the data-dependent NTK

The dynamics of neural-network training in the infinite-width limit are now well understood [15, 16]: for an appropriate scaling of the learning rate and initial weights, the neural network behaves like a linear model in which the data is transformed by the Neural Tangent Kernel (NTK) at initialization, which is defined as the product of the Jacobians of the logits at initialization. In the limit, neural network training implements kernel regression with the fixed NTK as the kernel.

However, finite neural networks outperform their infinite-width limits [17] and have different dynamics early in training [18]. In fact, rather than being constant, the data-dependent NTK, defined by Fort et al. [9] as the Gram matrix of the logit Jacobian, evolves with high velocity in the initial phase of training. Then, around the time of onset of linear mode connectivity, the NTK velocity stabilizes at a smaller value and remains nearly constant for the rest of the high learning rate training time.

Here we seek to understand which training samples contribute to the NTK gram matrix evolution. To empirically approximate the velocity of a NTK submatrix corresponding to a subset of images in a scale invariant way, we follow [9]. We compute the cosine distance between two NTK gram matrices on the given subset, one computed at epoch  $t$ , and another one at epoch  $t+1$ , one epoch later (see Appendix B.3). We look at submatrices of a fixed size, formed by examples with contiguous EL2N scores. Fig. 3 shows that higher EL2N scores lead to higher velocities. This relationship is not affected by the time at which both are computed.

Interestingly, the kernel velocity drops off sharply for examples with the very highest scores when label noise is introduced. In Section 4, we showed that dropping these examples boosts the accuracy of the final predictor. We hypothesize that, while the kernel velocity is higher for harder examples that the model is actively trying to fit, the kernel velocity drops off for the very highest scoring examples that might be too difficult to learn, perhaps because they are unrepresentative samples or they have label noise.

### 5.2 Connections to the Linear Mode Connectivity

We now examine how the ranking of the examples by EL2N connects to the geometry of the loss surface. In particular, Frankle et al. [19] studied the effect of minibatch randomness on the training trajectory, focusing on identifying the point in training when two networks, starting from the same weights, but trained with independent minibatches, converge to the same “linearly connected” mode. They find that, for standard vision datasets, the onset of this “linear mode connectivity” (LMC) happens early in training.

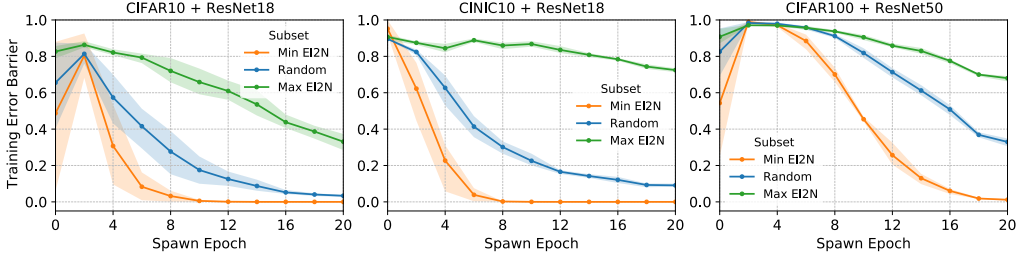


Figure 4: The final training error barrier between children on subsets of a 1000 highest (green) and lowest (orange) EL2N score examples, and randomly selected training subset (blue) as a function of the spawning time. *Left to right:* different dataset and network combinations.

More precisely, let  $w_1, w_2, \dots, w_T$  be the training trajectory of a *parent* network, fix a *spawning time*  $t^*$ , and let  $v_{t^*}, v_{t^*+1}, v_{t^*+2}, \dots, v_T$  be an independent training trajectory (i.e., with independent minibatches), beginning at  $v_{t^*} = w_{t^*}$ . We call  $v_T$  the child network and  $v_{t^*}, v_{t^*+1}, \dots$  the child trajectory. The (training) error barrier between two weights  $w$  and  $w'$ , denoted  $\text{err}(w, w'; S)$ , is the maximum deviation of the training error surface  $\hat{R}_S(\cdot)$  above the line connecting the empirical risk at  $w$  and  $w'$ . That is,

$$\text{err}(w, w'; S) = \sup_{\alpha \in [0,1]} \{\hat{R}_S(\alpha w + (1 - \alpha) w') - \alpha \hat{R}_S(w) - (1 - \alpha) \hat{R}_S(w')\}. \quad (5)$$

We then define the *mean (training) error barrier, spawning at  $t^*$ , at time  $t$* , for  $t^* \leq t \leq T$ , denoted  $\text{err}_t^{t^*}(S)$ , to be the expected error barrier between  $w_t$  and  $v_t$  on the data  $S$ . That is,

$$\text{err}_t^{t^*}(S) = \mathbb{E}_{w_{t^*+1:t}, v_{t^*+1:t}} [\text{err}(w_t, v_t; S)], \quad (6)$$

where the expectation is taken over the randomness in the trajectories of  $w$  and  $v$  *after*  $t^*$  due to the choice of minibatches, conditional on the initial trajectories up through time  $t^*$ . (Note that, at the end of training  $t = T$ , the supremum in  $\text{err}(w_T, v_T; S)$  is often achieved near  $\alpha = 1/2$ , and so this is a cheap approximation used in practice.) The “onset” of linear mode connectivity is the earliest spawning time  $t^*$  at which point  $\text{err}_T^{t^*}(S) \approx 0$ , where  $S$  is the whole training set. In our work, we instead compute the error barrier on *subsets of the training set*, which allows us to compare the training dynamics and modes on *subpopulations*.

In Fig. 4, we measure the mean error barrier  $\text{err}_t^{t^*}(S')$  as a function of the spawning time  $t^*$ , in the cases where  $S'$  are either 1) the training examples with the smallest scores, 2) the largest scores, or 3) a random subset of training examples. We find that the error barrier falls close to zero very rapidly for examples that have low EL2N scores, and stays high for high score examples. These findings suggest that the loss landscape derived from restricted subsets of examples with low and high EL2N behave very differently. The loss landscape derived from easy subsets of examples with low scores is quite flat, in the sense that error barriers between children as a function of spawn time rapidly diminish. On the other hand, the loss landscape derived from harder subsets of examples with higher scores is rougher, with higher error barriers that persist for longer in the spawn time. Further, this result is in agreement with the results presented in Section 5.1, showing that most of the learning happens in the high EL2N score examples.

## 6 Related Work

As we have already discussed, our work is closely related to an empirical study by Toneva et al. [8], which examines the frequency with which correct classification decisions are forgotten during training. The authors observe that examples that are rarely forgotten are also ones that do not contribute much to the final accuracy of the predictor. In particular, if we retrain from initialization after having removed these rarely forgotten examples from the training data, we achieve the same accuracy. Similar to our work, this work analyzes the dynamics of training in deep learning through the lens of training examples, and demonstrates that standard vision datasets have superfluous information. However, unlike forgetting scores, our proposed methods use only local information, bringing to light that the local ordering of examples is roughly preserved throughout training.

Coleman et al. [1] use a small proxy network in combination with other training data selection methods to find a small subset of important-for-training examples, that can then be used to train a

large state-of-the-art (SOTA) deep neural network. In their empirical study, they observe that most important examples selected via a proxy model are also important for training a SOTA network. In addition, they study a proxy which reuses SOTA network’s architecture but is trained for a shorter time. The authors observe that selecting the important examples after at least 50 epochs of training works better than selecting them at random, but not as well as after the full training run. They do not study shorter training times for proxies or relate it to the training dynamics in any other way.

Another line of related work is on coresets (see, e.g., [4, 5, 7, 20–22], and many others). The term *coresets* generally refers to a possibly weighted subset of training data. Much of the work on coresets is focused on identifying small training data subsets that provably yield an  $\epsilon$ -approximate solution to the original objective (on all the training data). Most guarantees require the problem to have special structure, such as convexity. For nonconvex problems, like training deep neural networks, guarantees are provided for very conservative proxies, e.g., based on Lipschitz constants or smoothness. While coreset selection comes with nice theoretical guarantees, in our opinion, the utility of these methods is best considered an empirical question.

Coresets have also been studied in the active learning community. Here, the goal is to select a small set of examples to label at any given iteration of training (see, e.g., [23–27], and references therein). Coreset selection has also been proposed as a way to increase model robustness [28].

Informally, removing a training example from the training data and not hurting the generalization error suggests that the example has small “influence” on the test data. Influence of the training examples on test examples is studied in sample-based explainability [29–31]. On the theory side, Feldman [14] recently proposed to model data as a mixture of populations and study the role of memorization when the data distribution is long-tailed. Feldman demonstrates conditions under which memorization is necessary for good generalization. In doing so, he proposes a definition of example memorization and influence, which can be interpreted as a leave-one-out notion of stability. In an empirical study following this work, Feldman and Zhang [32] demonstrate that classifiers trained on computer vision benchmarks benefit from memorization. In particular, training without high-memorization-value examples comes at a cost of accuracy of the learned neural network classifier. In Appendix F, we compare GraNd, EL2N, forgetting scores, and memorization values on CIFAR-100-trained Resnet50 networks; memorization values do not correlate with the other scores.

## 7 Discussion

In summary, our work both (1) introduces methods to significantly prune data without sacrificing test accuracy using *only* local information *very early* in training (Fig. 1), sometimes even at initialization, and (2) uses the resulting methods to obtain new scientific insights into how different subsets of training examples drive the dynamics of deep learning. We start from a principled approach by asking how much on average each training example influences the loss reduction of other examples, and from that starting point, we obtain 2 scores, namely gradient norm (GraNd) and error norm (EL2N) that bound or approximate this influence, with higher scores indicating higher potential influence. We find that examples with higher scores tend to be harder to learn, in the sense that they are forgotten more often over the entire course of training. We also find that the very highest scoring examples tend to be either unrepresentative outliers of a class, have non standard backgrounds or odd angles, are subject to label noise, or are otherwise difficult. This observation yields a simple and powerful sliding window method (Fig. 2) to prune data by keeping examples within a range of scores, where the start and the end of the range constitute just 2 hyperparameters that can be tuned via a validation set. Furthermore, we find that high-scoring examples primarily drive feature learning by maximally supporting the velocity of the NTK, whereas learning dynamics might actually give up on the very highest scoring examples that may correspond to unrepresentative examples or noise (Fig. 3). Finally we show that higher (lower) scoring subsets of examples contribute to a rougher (smoother) loss landscape (Fig. 4). Overall this decomposition of both loss landscape geometry and learning dynamics into differential contributions from different types of examples constitutes an exciting new methodology for analyzing deep learning. A deeper understanding of the differential role played by different subsets of examples could aid not only in data pruning, but also in curriculum design, active learning, federated learning with privacy, and analysis of fairness and bias.

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## References

- [1] C. Coleman, C. Yeh, S. Mussmann, B. Mirzasoleiman, P. Bailis, P. Liang, J. Leskovec, and M. Zaharia. *Selection via proxy: Efficient data selection for deep learning*. 2019. arXiv: [1906.11829](#).
- [2] M. Hwang, Y. Jeong, and W. Sung. “Data distribution search to select core-set for machine learning”. In: *Proc. 9th Int. Conf. Smart Media & Appl. (SMA 2020), Jeju, Korea*. 2020, pp. 17–19.
- [3] S. Har-Peled and A. Kushal. “Smaller coresets for k-median and k-means clustering”. *Discrete & Computational Geometry* 37.1 (2007), pp. 3–19.
- [4] J. H. Huggins, T. Campbell, and T. Broderick. “Coresets for scalable bayesian logistic regression” (2016). arXiv: [1605.06423](#).
- [5] T. Campbell and T. Broderick. “Bayesian coreset construction via greedy iterative geodesic ascent”. In: *Int. Conf. Machine Learning*. PMLR. 2018, pp. 698–706.
- [6] I. W. Tsang, J. T. Kwok, P.-M. Cheung, and N. Cristianini. “Core vector machines: Fast SVM training on very large data sets.” *Journal of Machine Learning Research* 6.4 (2005).
- [7] A. Munteanu, C. Schwiegelshohn, C. Sohler, and D. P. Woodruff. *On coresets for logistic regression*. 2018. arXiv: [1805.08571](#).
- [8] M. Toneva, A. Sordoni, R. T. d. Combes, A. Trischler, Y. Bengio, and G. J. Gordon. *An empirical study of example forgetting during deep neural network learning*. 2018. arXiv: [1812.05159](#).
- [9] S. Fort, G. K. Dziugaite, M. Paul, S. Kharaghani, D. M. Roy, and S. Ganguli. “Deep learning versus kernel learning: an empirical study of loss landscape geometry and the time evolution of the Neural Tangent Kernel”. In: *Advances in Neural Information Processing Systems*. 2020. arXiv: [2010.15110](#).
- [10] K. He, X. Zhang, S. Ren, and J. Sun. “Deep residual learning for image recognition”. In: *Proceedings of the IEEE conference on computer vision and pattern recognition*. 2016, pp. 770–778.
- [11] A. Krizhevsky, G. Hinton, et al. “Learning multiple layers of features from tiny images” (2009).
- [12] L. N. Darlow, E. J. Crowley, A. Antoniou, and A. J. Storkey. “CINIC-10 is not ImageNet or CIFAR-10”. *CoRR* abs/1810.03505 (2018). arXiv: [1810.03505](#).
- [13] C. Zhang, S. Bengio, M. Hardt, B. Recht, and O. Vinyals. “Understanding deep learning requires rethinking generalization”. In: *Int. Conf. Representation Learning (ICLR)*. 2017. arXiv: [1611.03530v2](#).
- [14] V. Feldman. “Does learning require memorization? a short tale about a long tail”. In: *Proc. 52nd Ann. ACM SIGACT Symp. Theory of Comput. (STOC)*. 2020, pp. 954–959.
- [15] A. Jacot, F. Gabriel, and C. Hongler. “Neural tangent kernel: Convergence and generalization in neural networks”. In: *Advances in Information Processing Systems (NeurIPS)*. 2018. arXiv: [1806.07572](#).
- [16] J. Lee, L. Xiao, S. S. Schoenholz, Y. Bahri, R. Novak, J. Sohl-Dickstein, and J. Pennington. *Wide neural networks of any depth evolve as linear models under gradient descent*. 2019. arXiv: [1902.06720](#).
- [17] S. Arora, S. S. Du, Z. Li, R. Salakhutdinov, R. Wang, and D. Yu. *Harnessing the power of infinitely wide deep nets on small-data tasks*. 2019. arXiv: [1910.01663](#).
- [18] A. Lewkowycz, Y. Bahri, E. Dyer, J. Sohl-Dickstein, and G. Gur-Ari. *The large learning rate phase of deep learning: the catapult mechanism*. 2020. arXiv: [2003.02218](#).
- [19] J. Frankle, G. K. Dziugaite, D. Roy, and M. Carbin. “Linear mode connectivity and the lottery ticket hypothesis”. In: *Int. Conf. Machine Learning (ICML)*. 2020, pp. 3259–3269. arXiv: [1912.05671](#).
- [20] P. K. Agarwal, S. Har-Peled, K. R. Varadarajan, et al. “Geometric approximation via coresets”. *Combinatorial and computational geometry* 52 (2005), pp. 1–30.

- [21] D. Feldman, M. Schmidt, and C. Sohler. “Turning big data into tiny data: Constant-size coresets for k-means, PCA, and projective clustering”. *SIAM J. Computing* 49.3 (2020), pp. 601–657.
- [22] E. Tolochinsky and D. Feldman. *Coresets for monotonic functions with applications to deep learning*. 2018. arXiv: [1802.07382](#).
- [23] K. Wei, R. Iyer, and J. Bilmes. “Submodularity in data subset selection and active learning”. In: *Int. Conf. Machine Learning*. PMLR. 2015, pp. 1954–1963.
- [24] O. Sener and S. Savarese. *Active learning for convolutional neural networks: A core-set approach*. 2017. arXiv: [1708.00489](#).
- [25] K. Killamsetty, D. Sivasubramanian, G. Ramakrishnan, and R. Iyer. *GLISTER: Generalization based Data Subset Selection for Efficient and Robust Learning*. 2020. arXiv: [2012.10630](#).
- [26] B. Mirzasoleiman, J. Bilmes, and J. Leskovec. “Coresets for data-efficient training of machine learning models”. In: *Int. Conf. Machine Learning (ICML)*. PMLR. 2020, pp. 6950–6960. arXiv: [1906.01827](#).
- [27] Y. Shen, H. Yun, Z. C. Lipton, Y. Kronrod, and A. Anandkumar. *Deep active learning for named entity recognition*. 2017. arXiv: [1707.05928](#).
- [28] B. Mirzasoleiman, K. Cao, and J. Leskovec. “Coresets for Robust Training of Neural Networks against Noisy Labels” (2020). arXiv: [2011.07451](#).
- [29] P. W. Koh and P. Liang. “Understanding black-box predictions via influence functions”. In: *Int. Conf. Machine Learning*. PMLR. 2017, pp. 1885–1894.
- [30] E. Barshan, M.-E. Brunet, and G. K. Dziugaite. “Relatif: Identifying explanatory training samples via relative influence”. In: *Int. Conf. Artificial Intelligence and Statistics (AISTATS)*. 2020, pp. 1899–1909.
- [31] G. Pruthi, F. Liu, S. Kale, and M. Sundararajan. “Estimating Training Data Influence by Tracing Gradient Descent”. In: *Advances in Neural Information Processing Systems*. 2020.
- [32] V. Feldman and C. Zhang. *What neural networks memorize and why: Discovering the long tail via influence estimation*. 2020. arXiv: [2008.03703](#).
- [33] J. Bradbury, R. Frostig, P. Hawkins, M. J. Johnson, C. Leary, D. Maclaurin, G. Necula, A. Paszke, J. VanderPlas, S. Wanderman-Milne, and Q. Zhang. *JAX: composable transformations of Python+NumPy programs*. Version 0.2.5. 2018.
- [34] J. Heek, A. Levskaya, A. Oliver, M. Ritter, B. Rondepierre, A. Steiner, and M. van Zee. *Flax: A neural network library and ecosystem for JAX*. Version 0.3.4. 2020.

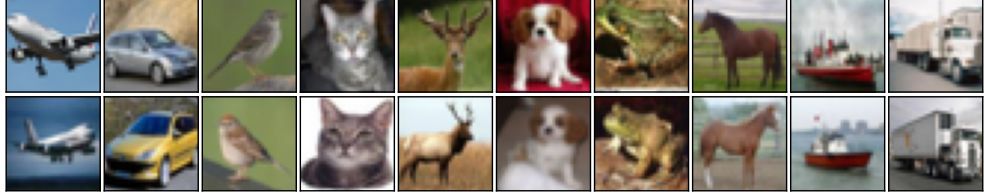


Figure 5: Examples with the smallest (first row) and second smallest (second row) GraNd scores for each class (columns, from left to right: airplane, automobile, bird, cat, deer, dog, frog, horse, ship, truck) from a ResNet18 trained on CIFAR-10. GraNd scores were calculated at initialization.



Figure 6: Examples with the second largest (first row) and largest (second row) GraNd scores for each class (columns, from left to right: airplane, automobile, bird, cat, deer, dog, frog, horse, ship, truck) from a ResNet18 trained on CIFAR-10. GraNd scores were calculated at initialization.

## A Ethical and societal consequences

This work raises several ethical considerations. Being, an empirically driven work, it consumed considerable energy. However, we hope that it will enable advancements in theory that will more efficiently guide experiments. Also, we focus mostly on accuracy as a metric, which tends to hide disparate effects on marginalized groups. But since this work attempts to explicitly uncover the influence of training examples and sub-populations, we hope that it will lead to methods that will decrease bias in the training procedure, especially if marginalized groups are under-represented in the dataset and are thus difficult to learn.

## B Implementation Details

The code is made available at [https://github.com/mansheej/data\\_diet](https://github.com/mansheej/data_diet)

### B.1 Resources used

We run all experiments on a single 16GB NVIDIA Tesla V100 GPU. The entire project (from early exploration to final paper) used about 15000 GPU hours. We used an internal cluster at ServiceNow.

### B.2 Training details

**Deep Learning Frameworks.** We used JAX [33] and Flax [34] in our implementations.

**Data.** We use CIFAR-10, CIFAR-100 [11], and CINIC-10 [12]. CIFAR-10 and CIFAR-100 are used in their standard format. For CINIC-10, we combine the training and validation sets into a single training set with 180000 images. The standard test set of 90000 images is used for testing. Each dataset is normalized by its per channel mean and standard deviation over the training set. All datasets get the same data augmentation: pad by 4 pixels on all sides, random crop to  $32 \times 32$  pixels, and left-right flip image with probability half.



Figure 7: Examples with the smallest (first row) and second smallest (second row) EL2N scores for each class (columns, from left to right: airplane, automobile, bird, cat, deer, dog, frog, horse, ship, truck) from a ResNet18 trained on CIFAR-10. EL2N scores were calculated at epoch 10.



Figure 8: Examples with the second largest (first row) and largest (second row) EL2N scores for each class (columns, from left to right: airplane, automobile, bird, cat, deer, dog, frog, horse, ship, truck) from a ResNet18 trained on CIFAR-10. EL2N scores were calculated at epoch 10.

**Models.** We use ResNet18-v1 and ResNet50-v1. Our implementation is based on the example in Flax [34] designed for larger high resolution images. Since CIFAR and CINIC images are  $32 \times 32$  pixels only, we use the low-resolution variant of these networks: the first two layers (a convolution layer with  $7 \times 7$  kernel and  $2 \times 2$  stride, and a max pooling layer with  $3 \times 3$  kernel and  $2 \times 2$  stride) are replaced with a single convolution layer with  $3 \times 3$  kernel and  $1 \times 1$  stride.

**Training hyperparameters.** All networks are trained with the Stochastic Gradient Descent (SGD) optimizer, learning rate = 0.1, nesterov momentum = 0.9, weight decay = 0.0005. For CIFAR-10 and CIFAR-100, we use batch size = 128, and for CINIC-10, we use batch size = 256. The learning rate is decayed by a factor of 5 after 60, 120 and 160 epochs and all networks are trained for a total of 200 epochs (for the full dataset, i.e. 78000 steps for CIFAR-10 and CIFAR-100, and 140600 steps for CINIC-10). When using a pruned dataset, to allow for a fair comparison with the full dataset, we keep the number of iterations and schedule fixed for different pruning levels.

### B.3 Experimental details

**Reporting results.** For every quantity we plot, we do 4 independent runs (independent model initialization and SGD noise) and report the mean and the 16th to 84th percentile of obtained accuracies for representing variability across runs. The mean is reported as lines and the variability is reported as shading in the plots.

**Calculating scores.** All scores (EL2N, GraNd and forgetting scores) are calculated by averaging the scores across 10 independent runs.

**Random label experiments.** For the random label experiments, at the beginning of training, we pick 10% of the examples randomly and permute their labels (to keep overall label statistics fixed). The subsets are selected as follows:

1. score all examples and sort them in ascending order by score;
2. drop the set of images with the smallest scores that make up a fraction of the dataset equal to the specified offset;

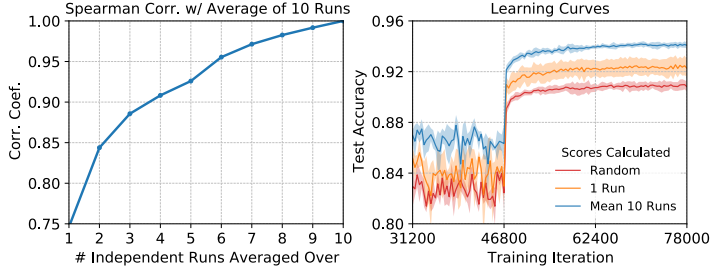


Figure 9: The effect of averaging GraNd scores calculated at initialization of a ResNet18 trained on CIFAR-10. *Left:* The Spearman rank correlation coefficient between GraNd scores obtained by averaging over a given number of independent runs (x-axis) to those obtained by averaging over 10 independent runs. *Right:* Training accuracy curves for ResNet18 trained on a 50% subset of CIFAR-10. The training subset is obtained by either random sampling or keeping the examples with the largest GraNd scores. We compare the test accuracy obtained using GraNd scores from 1 initialization and from averaging over 10 independent initializations. We zoom in to the end of training to highlight the differences between the learning curves.

3. keep the next set of images based on the given subset size;
4. drop all the following images.

**Kernel velocity experiments.** The kernel velocities are calculated with 100 examples. The examples are picked as follows: first, score all examples and sort them in ascending order by score; then, for every example index in the figure, calculate the NTK gram matrix velocity (described in Section 5.1) for 100 contiguous sorted images starting at that example index.

The NTK submatrix velocity on a subset of examples at a particular point in training is defined as follows. Let the examples be  $x_1, x_2, \dots, x_m$ . Let the time at which the NTK submatrix is calculated be  $t$ , the parameters of the network  $f$  is  $w_t$ . Let  $C$  be the number of classes and  $N$  the number of parameters in the network. Using the notation in Section 2.1, the  $k$ th logit gradient of the model on example  $i$  is  $\psi_t^{(k)}(x_i) = \nabla_{w_t} f_t^{(k)}(x_i)$ . We cast these into a  $mC \times N$  matrix  $\Psi_t$  where the rows run over each logit gradient of each image and the columns run over the parameters of the model. The NTK submatrix is a  $mC \times mC$  matrix given by  $K_t = \Psi_t \Psi_t^T$ . The kernel velocity is calculated as

$$v = 1 - \frac{\langle K_t, K_{t+1} \rangle}{\|K_t\| \|K_{t+1}\|} \quad (7)$$

where  $\langle \cdot, \cdot \rangle$  is the Frobenius inner product and  $\|\cdot\|$  is the Frobenius norm.

**Linear mode connectivity experiments.** The training error barrier between children is calculated by following [9, 19]. In addition to estimating the error barrier on 1000 random images, we also estimate the error barrier on 1000 images with the largest and smallest EL2N score. The EL2N score used is calculated at epoch 10 of the parent run.

## C Example Images

In this section, we examine the examples with small and large GraNd and EL2N scores for a ResNet18 trained on CIFAR-10. GraNd scores were computed at initialization and EL2N scores at epoch 10. We show two examples from each class with both minimum and maximum EL2N and GraNd scores in Figs. 5 to 8. The examples with the minimum GraNd and EL2N scores tend to be simple, canonical representations of each class pictured from very typical angles. The examples with maximum scores are harder to identify; they are blurrier, from strange angles or have unexpected backgrounds or other artifacts.

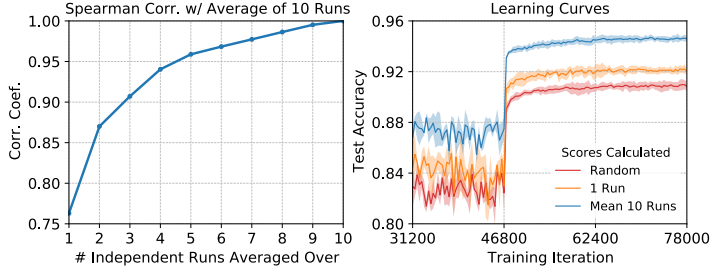


Figure 10: The effect of averaging EL2N scores calculated at epoch 10 of a ResNet18 trained on CIFAR-10. *Left:* The Spearman rank correlation coefficient between EL2N scores obtained by averaging over a given number of independent runs (x-axis) to those obtained by averaging over 10 independent runs. *Right:* Training accuracy curves for ResNet18 trained on a 50% subset of CIFAR-10. The subset is selected by either random sampling or keeping the examples with the largest EL2N scores. We compare the test accuracy obtained when pruning based on EL2N scores from a single run and from averaging over 10 independent runs. We zoom in to the end of training to highlight the differences between the learning curves.

## D Additional Experiments

### D.1 Sensitivity analysis of GraNd and EL2N scores

In all our experiments, GraNd and EL2N scores are averaged over 10 independent initializations or runs. This turns out to be essential for successful pruning using the scores.

In Fig. 9 (right) we show the effect of averaging on GraNd scores computed at initialization on a ResNet18, CIFAR-10. On average, GraNd scores for any individual run have a Spearman rank correlation of about 0.75 with the GraNd scores averaged over 10 runs.

We also compare learning curves when training using 50% of the training data selected as follows: using high GraNd scores at a single initialization; using high GraNd scores averaged over 10 initializations; randomly (baseline). As seen in Fig. 9 (right), using scores averaged over 10 initializations performs significantly better. In Fig. 10, we show similar results for EL2N scores calculated at epoch 10 on a ResNet18 trained on CIFAR-10.

These results suggest that GraNd and EL2N scores represent properties of the dataset rather than of a specific network weights. To get an accurate ranking of example importance, we need to average out the effects of individual initializations/weights. Empirically, we find that averaging over 10-20 runs suffices, and averaging over more runs has insignificant additional benefit.

### D.2 Comparison between scores from different architectures on the same dataset

In this section, we examine how the choice of network architecture affects pruning with EL2N and GraNd scores. Specifically, we repeat the experiment in Fig. 1 bottom row, but for a ResNet18 and a ResNet50 trained on CIFAR-10. The results are shown in Fig. 11. Both networks share the same overall patterns: pruning by GraNd or EL2N scores before training results in roughly the same accuracy for different levels of pruning. Further, independently of the network architecture tested, data pruning by EL2N scores computed at epoch 20 is competitive with pruning based on forgetting scores computed at epoch 200. When trained on either of the networks, pruning based on GraNd scores does significantly better than the random baseline. Overall, these results suggest that network depth has a small effect on our data-pruning results.

Surprisingly, GraNd scores at initialization for ResNet18 seem to be better for pruning than those for ResNet50. In future work, we hope to better understand the relationship between the level of pruning and network depth and width.

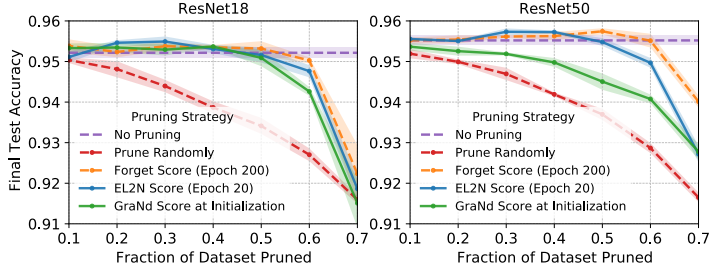


Figure 11: Experimental setup similar to Fig. 1. We compare CIFAR10 data pruning on ResNet18 and ResNet50. The y-axis indicates the final test accuracy achieved by training after pruning different fractions of the dataset (x-axis). Compare forgetting scores at the end of training, EL2N scores early in training (at epoch 20) and GraNd scores at initialization. In each case, examples with the lowest scores are pruned and then the networks are trained from initialization on the data that was not pruned. In both plots accuracy achieved by training on the full dataset and on a random subset of the corresponding size dataset are used as baselines.

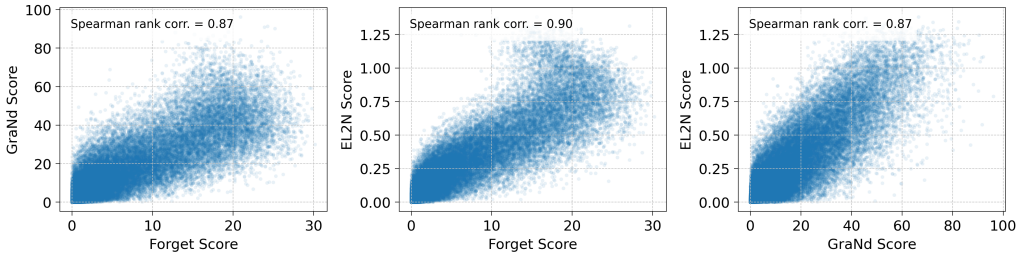


Figure 12: For a ResNet18 trained on CIFAR-10, we compare the values and Spearman rank correlations for pairs of scores. The scores compared are GraNd scores at initialization, EL2N scores at epoch 20 and forgetting scores at epoch 200.

### D.3 Correlations between scores

As discussed in previous sections, we find that different scores lead to similar pruning results. In Fig. 12, for a ResNet18 trained on CIFAR-10, we compare the values and Spearman rank correlations for pairs of scores. GraNd scores are computed at initialization, EL2N scores at epoch 20 and forgetting scores at epoch 200. The scores have high Spearman rank correlation with each other. EL2N and forgetting scores, which have the most similar performance, have the highest Spearman rank correlation.

## E Noise

### E.1 Noisy Examples in Low Pruning Regime

We repeat the noisy labels experiment of Fig. 2 in a different regime; instead of pruning 60% of the data, we prune just 50% of the data. Results are shown in Fig. 13. For the experiment with no labels randomized, at this lower pruning level we no longer see a boost in performance from dropping the highest score examples. However we do see decreasing marginal gains. This suggests that when we have enough data, keeping the high score examples, which are often noisy or difficult, does not hurt performance and can only help.

### E.2 Scores for Noisy Examples

We now examine how adding noise affects the EL2N scores of examples. This analysis is done for the experiment in Fig. 2. See Section 4 for details. Results are shown in Fig. 14. Two results suggest that EL2N scores can be used to identify images with corrupted labels. First, the EL2N

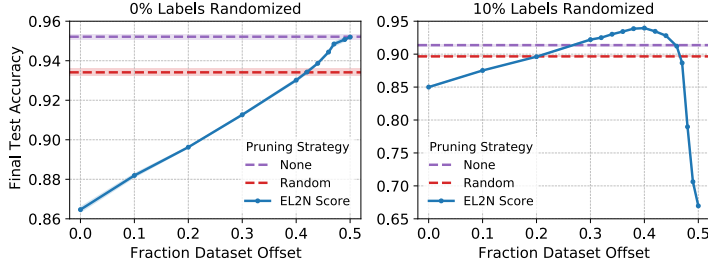


Figure 13: ResNet18 trained on a 50% subset of CIFAR-10 with clean (*left*) and 10% randomized labels (*right*). The training subset contains the *lowest* scoring examples *after* examples with scores below the offset (x-axis) are discarded. Scores computed at epoch 10.

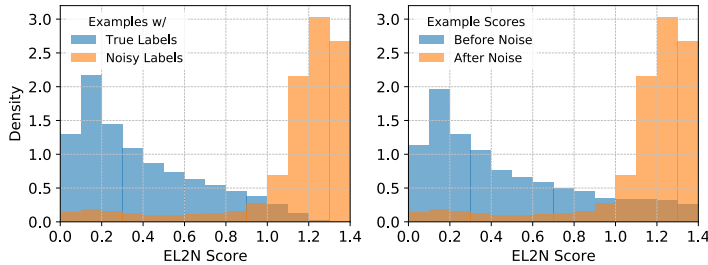


Figure 14: Scores for examples in the noisy label experiment of Fig. 2 *Left*. Distribution of EL2N scores for examples with true labels and corrupted labels. *Right*. Distribution of EL2N scores for examples that are corrupted before and after noise is added.

scores of images with corrupted labels tend to be higher than those with regular labels. Second, after an example’s label is corrupted, its EL2N scores tends to be larger than before.

### E.3 Noise and GraNd Scores

We repeat the experiment from Fig. 2 (Section 4) except, instead of using EL2N scores at epoch 10, we use GraNd scores at initialization for pruning. As seen in Fig. 15, when none of the labels are corrupted, the results are similar to the previous variant of this experiment; keeping examples with larger GraNd scores in the training subset leads to better generalization performance. However, in the case where 10% of the labels are corrupted, we see the opposite trend: a model trained on larger GraNd scores ends up with lower accuracy, performing even worse than the random subset baseline. Note that the GraNd scores are calculated at initialization with a randomly initialized network. These results suggest that GraNd scores at initialization successfully find important examples only because of some favorable properties of the data distribution; adding noise to samples selected uniformly over the training set cripples the method. In future work, we further explore how high Bayes error rate effects GraNd scores at initialization.

## F Comparison to memorization threshold

In Fig. 16 we compare EL2N scores to the memorization values defined in [32]. The memorization values for 1015 CIFAR100 examples are provided by the authors. We replicate their setting by training a ResNet50 on CIFAR-100 and compute the EL2N scores at epoch 20 for the 1015 examples they provide. As seen from Fig. 16, memorization values and EL2N scores do not appear to be correlated.

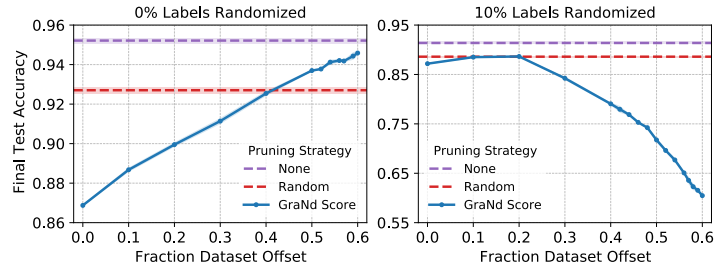


Figure 15: ResNet18 trained on a 40% subset of CIFAR-10 with clean (*left*) and 10% randomized labels (*right*). The training subset contains the *lowest* scoring examples *after* examples with scores below the offset are discarded. Use GraNd scores computed at initialization.

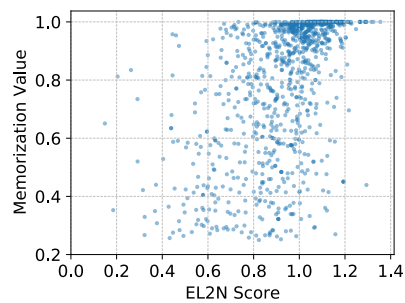


Figure 16: Comparison of EL2N scores and Memorization values from [32]