RIGGING THE LOTTERY: 
MAKING ALL TICKETS WINNERS

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ABSTRACT

Sparse neural networks have been shown to be computationally efficient networks with improved inference times. There is a large body of work on training dense networks to yield sparse networks for inference (Molchanov et al., 2017; Zhu & Gupta, 2018; Louizos et al., 2017; Li et al., 2016; Guo et al., 2016). This limits the size of the largest trainable sparse model to that of the largest trainable dense model. In this paper we introduce a method to train sparse neural networks with a fixed parameter count and a fixed computational cost throughout training, without sacrificing accuracy relative to existing dense-to-sparse training methods. Our method updates the topology of the network during training by using parameter magnitudes and infrequent gradient calculations. We show that this approach requires less floating-point operations (FLOPs) to achieve a given level of accuracy compared to prior techniques. We demonstrate state-of-the-art sparse training results with ResNet-50, MobileNet v1 and MobileNet v2 on the ImageNet-2012 dataset. Finally, we provide some insights into why allowing the topology to change during the optimization can overcome local minima encountered when the topology remains static.

1 INTRODUCTION

The parameter and FLOP efficiency of sparse neural networks is now well demonstrated on a variety of problems (Han et al., 2015; Srinivas et al., 2017). Some work has even shown inference time speedups are possible on Recurrent Neural Networks (RNNs) (Kalchbrenner et al., 2018) and Convolutional Neural Networks (ConvNets) (Park et al., 2016). Currently, the most accurate sparse models are obtained with techniques that require, at a minimum, the cost of training a dense model in terms of memory and FLOPs (Zhu & Gupta, 2018; Guo et al., 2016), and sometimes significantly more (Molchanov et al., 2017). This paradigm has two main limitations:

1. The maximum size of sparse models is limited to the largest dense model that can be trained. Even if sparse models are more parameter efficient, we can’t use pruning to train models that are larger and more accurate than the largest possible dense models.

2. It is inefficient. Large amounts of computation must be performed for parameters that are zero valued or that will be zero during inference.

Additionally, it remains unknown if the performance of the current best pruning algorithms are an upper bound on the quality of sparse models. Gale et al. (2019) found that three different algorithms all achieve about the same sparsity / accuracy trade-off. However, this is far from conclusive proof that no better performance is possible. In this work we show the surprising result that dynamic sparse training, which includes the method we introduce below, can find more accurate models than the current best approaches to pruning initially dense networks. Importantly, our method does not change the FLOPs required to execute the model during training, allowing one to decide on a specific inference cost prior to training.

The Lottery Ticket Hypothesis (Frankle & Carbin, 2019) hypothesized that if we can find a sparse neural network with iterative pruning, then we can train that sparse network from scratch, to the same level of accuracy, by starting from the original initial conditions. In this paper we introduce a
new method for training sparse models without the need of a “lucky” initialization; for this reason, we call our method “The Rigged Lottery” or RigL. We show that this method is:

- **Memory efficient**: It requires memory only proportional to the size of the sparse model. It never requires storing quantities that are the size of the dense model. This is in contrast to [Dettmers & Zettlemoyer, 2019] which requires storing the momentum for all parameters, even those that are zero valued.
- **Computationally efficient**: The amount of computation required to train the model is proportional to the number of nonzero parameters in the model.
- **Accurate**: The performance achieved by the method matches and sometimes exceeds the performance of pruning based approaches.

Our method works by infrequently using instantaneous gradient information to inform a re-wiring of the network. We show that this allows the optimization to escape local minima where it would otherwise become trapped if the sparsity pattern were to remain static. Crucially, as long as the full gradient information is needed less than every $\frac{1}{1 - \text{sparsity}}$ iterations, then the overall work remains proportional to the model sparsity.

## 2 Related Work

Research on finding sparse neural networks dates back at least three decades, where [Mozer & Smolensky, 1989] concluded that pruning weights based on magnitude was a simple and powerful technique. [Ström, 1997] later introduced the idea of retraining the previously pruned network to increase accuracy. [Han et al., 2016b] went further and introduced multiple rounds of magnitude pruning and retraining. This is, however, relatively inefficient, requiring ten rounds of retraining when removing 20% of the connections to reach a final sparsity of 90%. To overcome this problem, [Narang et al., 2017] introduced gradual pruning, where connections are slowly removed over the course of a single round of training. [Zhu & Gupta, 2018] refined the technique to minimize the amount of hyper-parameter selection required.

A diversity of approaches not based on magnitude based pruning have also been proposed. [LeCun et al., 1990] and [Hassibi & Stork, 1993] are some early examples, but impractical for modern neural networks as they use information from the Hessian to prune a trained network. More recent work includes $L_0$ Regularization [Christos Louizos, 2018], Variational Dropout [Molchanov et al., 2017], Dynamic Network Surgery [Guo et al., 2016] and Sensitivity Driven Regularization [Tartaglione et al., 2018]. [Gale et al., 2019] examined magnitude pruning, $L_0$ Regularization and Variational Dropout and concluded that they all achieve about the same accuracy versus sparsity tradeoffs on ResNet-50 and Transformer architectures.

Training techniques that allow for sparsity throughout the entire training process were, to our knowledge, first introduced in Deep Rewiring (DeepR) [Bellec et al., 2017]. In DeepR, the standard Stochastic Gradient Descent (SGD) optimizer is augmented with a random walk in parameter space. Additionally, connections have a pre-defined sign assigned at random; when the optimizer would normally flip the sign, the weight is set to 0 instead and new weights are activated at random.

Sparse Evolutionary Training (SET) [Mocanu et al., 2018] proposed a simpler scheme where weights are pruned according to the standard magnitude criterion used in pruning and added back at random. The method is simple and achieves reasonable performance in practice. Dynamic Sparse Reparameterization (DSR) [Mostafa & Wang, 2019] introduces the idea of allowing the parameter budget to shift between different layers of the model, allowing for non-uniform sparsity. This allows the model to put parameters where they are most effective. Unfortunately, the models under consideration are mostly ConvNets, so the result of this parameter reallocation (which is to decrease the sparsity of early layers and increase the sparsity of later layers) has the overall effect of increasing the FLOP count because the spatial size is largest at the beginning. Sparse Networks from Scratch (SNFS) [Dettmers & Zettlemoyer, 2019] introduces the idea of using the momentum of each parameter as the criterion to be used for growing weights and demonstrates it leads to an improvement in test accuracy. Like DSR, they allow the sparsity of each layer to change and focus on a constant

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* Pronounced "riggle".
Table 1: Comparison of different sparse training techniques. Drop and Grow columns correspond to the strategies used during the mask update. Selectable FLOPs is possible if the cost of training the model is fixed at the beginning of training.

<table>
<thead>
<tr>
<th>Method</th>
<th>Drop</th>
<th>Grow</th>
<th>Selectable FLOPs</th>
<th>Space &amp; FLOPs</th>
</tr>
</thead>
<tbody>
<tr>
<td>SNIP</td>
<td>$\min(</td>
<td>w</td>
<td>)$</td>
<td>none</td>
</tr>
<tr>
<td>DeepR</td>
<td>stochastic</td>
<td>random</td>
<td>yes</td>
<td>sparse</td>
</tr>
<tr>
<td>SET</td>
<td>$\min(</td>
<td>w</td>
<td>)$</td>
<td>random</td>
</tr>
<tr>
<td>DSR</td>
<td>$\min(</td>
<td>w</td>
<td>)$</td>
<td>random</td>
</tr>
<tr>
<td>SNFS</td>
<td>$\min(</td>
<td>w</td>
<td>)$</td>
<td>momentum</td>
</tr>
<tr>
<td>RigL (ours)</td>
<td>$\min(</td>
<td>w</td>
<td>)$</td>
<td>gradient</td>
</tr>
</tbody>
</table>

There has also been a line of work investigating the Lottery Ticket Hypothesis (Frankle & Carbin, 2019). Frankle et al. (2019) showed that the formulation must be weakened to apply to larger networks such as ResNet-50 (He et al., 2015). In large networks, instead of the original initialization, the values after thousands of optimization steps must be used for initialization. Zhou et al. (2019) showed that lottery tickets obtain non-random accuracies even before the training has started. Though the possibility of training sparse neural networks with a fixed sparsity mask using lottery tickets is intriguing, it remains unclear whether it is possible to generate such initializations – both of masks and parameters – de novo.

3 RIGGING THE LOTTERY

Our method, RigL, is illustrated in Figure 1. At regularly spaced intervals our method removes a fraction of connections based on weight magnitudes and activates new ones using instantaneous gradient information. After updating the connectivity, training continues with the updated network until the next update. The main parts of our algorithm, Sparsity Distribution, Update Schedule, Drop Criterion, Grow Criterion, and the various options we considered for each, are explained below. The improved performance of RigL is due to two reasons: the use of a new method for activating connections that is efficient and more effective than choosing at random, and the use of a natural extension to an existing method for distributing parameters statically among convolutional layers.

(0) Notation. Given a dataset $D$ with individual inputs $x_i$ and targets $y_i$, one can train a neural network to minimize the loss function $\sum_i L(f_\theta(x_i), y_i)$, where $f_\theta(x)$ is the neural network with parameters $\theta$ of length $N$. The vector $\theta$ can be decomposed into parameters $\theta^l$, of length $N^l$, for
each layer \( l \). A sparse network keeps only a fraction \( D \in (0, 1) \) of all connections, resulting in a \textbf{sparsity} of \( S = 1 - D \). More precisely, denoting the sparsity of individual layers by \( s^l \), the total parameter count of the sparse neural network satisfies \( \sum_l (1 - s^l)N^l = (1 - S) * N \).

\( \text{(1) Sparsity Distribution.} \) There are many ways of distributing the non-zero weights across the layers while satisfying the equality above. We avoid re-allocating parameters between layers during the training process as it makes it difficult to target a specific final FLOP budget, which is important for many inference applications. We consider the following three strategies:

1. \textit{Uniform}: The sparsity \( s^l \) of each individual layer is the same as the total sparsity \( S \), except for the first and last layers; following the strategy of [Gale et al., 2019], we fix the sparsity of the last layer to 0.8 and keep the first layer as a dense layer.

2. \textit{Erdős-Rényi}: As introduced in [Mocanu et al., 2018], \( s^l \) scales proportionally to the ratio \( \frac{n^l_{i+1} + n^l_{i}}{n^l_{i}} \), where \( n^l_{i} \) denotes number of neurons at layer \( l \). This enables the number of connections in a sparse layer to scale with the number of output channels.

3. \textit{Erdős-Rényi-Kernel (ERK)}: This method modifies the original Erdős-Rényi formulation by including the kernel dimensions in the scaling factors. In other words, the number of parameters of the sparse convolutional layers are scaled by \( \frac{w^{l_{i+1}} + w^{l_i} + h^{l_i} + h^{l_{i+1}}}{w^{l_i} + w^{l_{i+1}} + h^{l_i} + h^{l_{i+1}}} \), where \( w^l \) and \( h^l \) are the width and the height of the \( l \)th convolutional kernel. Sparsity of the fully connected layers scale as in the original Erdős-Rényi formulation. Similar to Erdős-Rényi, ERK allocates higher sparsities to the layers with more parameters while allocating lower sparsities to the smaller ones.

In all methods, the bias and batch-norm parameters are kept dense.

\( \text{(2) Update Schedule.} \) The update schedule is defined by the following parameters: (1) the number of iterations between sparse connectivity updates (\( \Delta I \)), (2) the iteration at which to stop updating the sparse connectivity (\( I_{\text{end}} \)), (3) the initial fraction of connections updated (\( \alpha \)) and (4) a function \( f_{\text{decay}} \), invoked every \( \Delta I \) iterations until \( I_{\text{end}} \), possibly decaying the fraction of updated connections over time. For the latter we choose to use \textit{cosine} annealing, as we find it slightly outperforms the other methods considered.

\[
f_{\text{decay}}(t) = \frac{\alpha}{2} \left( 1 + \cos \left( \frac{t\pi}{I_{\text{end}}} \right) \right)
\]

Alternatives to cosine annealing like a \textit{constant} schedule and \textit{inverse power} annealing are studied in the Appendix D.

\( \text{(3) Drop criterion.} \) Over the course of training, we drop the lowest magnitude weights according to the update schedule. Specifically, we drop the connections given by \( \text{TopK}(\cdot[\theta^l], f_{\text{decay}}(t)(1 - s^l)N^l) \).

\( \text{(4) Grow criterion.} \) The novelty of our method lies in how we grow new connections. We grow the connections with highest magnitude gradients, \( \text{TopK}_{\theta^l_{\text{active}}}(|\text{grad}(\theta^l)|, f_{\text{decay}}(t)(1 - s^l)N^l) \), where \( \theta^l_{\text{active}} \) is the set of active connections after the drop step. Newly activated connections are initialized to zero. This procedure can be applied to each layer in sequence and the dense gradients can be discarded immediately after selecting the top connections. If a layer is too large to materialize the full gradient with respect to the weights, then we can further reduce the memory requirements by performing an iterative calculation:

1. Initialize the set \( TK = \{\} \).
2. Materialize a subset of size \( M \) of the full gradient, which we denote \( G_{i+M} \).
3. Update \( TK \) to contain the Top-K elements of \( G_{i+M} \) concatenated with \( TK \).
4. Repeat steps 1 through 3 until all of the gradients have been materialized. The final set \( TK \) contains the connections we wish to grow.

As long as \( \Delta I > \frac{1}{I_{\text{end}}} \) the total work in calculating dense gradients is amortized and still proportional to \( S \). This is in contrast to the method of [Dettmers & Zettlemoyer, 2019], which requires calculating and storing the full gradients at each optimization step.

\( ^{\dagger}\text{TopK}(v,k) \text{ returns the indices and values of the top-k elements of vector } v. \)
Our experiments focus on the ImageNet-2012 dataset and we use the TensorFlow Model Pruning library (Zhu & Gupta, 2018) to sparsify our networks. A Tensorflow (Abadi et al., 2015) implementation of our method along with three other baselines (SET, SNFS, SNIP) will be open sourced. In all experiments, we use SGD with momentum as our optimizer. We set the momentum of the optimizer to 0.9, $L_2$ regularization coefficient to 0.0001, and label smoothing to 0.1. The learning rate schedule starts with a linear warm up reaching its maximum value of 1.6 at epoch 5 which is then dropped by a factor of 10 at epochs 30, 70 and 90. We train our networks with a batch size of 4096 for 32000 steps which roughly corresponds to 100 epochs of training. When we increase the training steps by a factor $x$, the anchor epochs of the learning rate schedule are also scaled by the same factor; we indicate this scaling with a subscript (e.g. RigL$_{5 \times}$). Our training pipeline uses standard data augmentation, which includes random flips and crops.

### 4 Empirical Evaluation

Figure 2-left summarizes the performance of various methods training on an 80% sparse ResNet-50. We also train small dense networks with equivalent parameter count. All sparse networks use the constant sparsity distribution and a cosine update schedule ($\alpha = 0.1, \Delta I = 50$). Overall, we observe that the performance of all methods improves with training time; thus, for each method we run extended training with up to $5 \times$ the training steps of the original. The calculation of the number of FLOPs required by each method are detailed in Appendix E.

As noted by Gale et al. (2019), Evci et al. (2019), Frankle et al. (2019), and Mostafa & Wang (2019), training a fixed sparsity network from scratch (Static) leads to inferior performance. Training a small dense network with the same number of parameters gets better results than Static, but it fails to match the performance of dynamic sparse models. All dynamic sparse methods are able to achieve high accuracies, but RigL achieves the highest accuracy and does so while consistently requiring fewer FLOPs than the other methods.

Given that different applications or scenarios might require a limit on the number of FLOPs for inference, we investigate the performance of our method at various sparsity levels. As mentioned previously, one strength of our method is that its resource requirements are constant throughout training and can be chosen before training begins. In Figure 2-right we show the performance of our method at different sparsities and compare them with the pruning results of Gale et al. (2019), which uses 1.5x training steps, relative to the original 32k. To make a fair comparison with regards to FLOPs, we scale the learning schedule of all other methods by 5x. Note that even after extending the training, it takes less FLOPs to train sparse networks (except for the 80% sparse RigL-ERK) compared to the pruning method.
### Table 2: Performance and cost of sparse training methods on training 80% and 90% sparse ResNet-50s.

FLOPs needed for training and test are normalized with the FLOPs of a dense model. See Appendix E for further explanation on how FLOPs are calculated. Methods with a subscript indicate a rescaled training time, whereas '*' indicates results obtained from another work. (ERK) corresponds to the sparse networks with Erdős-Renyi-Kernel sparsity distribution. RigL\(_{5,\times}\) (ERK) achieves 76.89% Top-1 Accuracy using only 20% of the parameters of a dense model and 42% of its FLOPs.

RigL\(_{5,\times}\), our method with constant sparsity distribution, follows pruning very closely in all sparsity levels. Sparse networks that use Erdős-Renyi-Kernel (ERK) sparsity distribution, however, obtain a remarkable performance that exceeds the performance of pruning based models. As observed earlier, training either smaller dense models (with the same number of parameters) or static sparse models fails to perform at a comparable level.

A more fine grained comparison of sparse training methods is presented in Table 2. Methods using uniform sparsity distribution and whose FLOP/memory footprint scales directly with the sparsity of the model are placed in the first sub-group of the table. The second sub-group includes DSR and networks with ERK sparsity distribution which require a higher number of FLOPs for inference. The final sub-group includes methods that require larger storage and the work of a training dense model.

#### 4.2 MobileNet

MobileNet is a compact architecture that performs remarkably well in resource constrained settings. Due to its compact nature with separable convolutions it is known to be notoriously difficult to sparsify (Zhu & Gupta, 2018). In this section we apply our method to MobileNet-v1 (Howard et al., 2017) and MobileNet-v2 (Sandler et al., 2018). Due to its low parameter count we keep the first layer dense, and evaluate using ERK and Uniform sparsity distribution to sparsify the remaining layers.

The performance of sparse MobileNets trained with RigL as well as the baselines are shown in Figure 3. We do extended training (5x of the original number of steps) for all runs in this section. Although MobileNets are more sensitive to sparsity compared to the ResNet-50 architecture, RigL successfully trains sparse MobileNets at high sparsities and exceeds the performance of previously reported pruning results.
Figure 3: (left) RigL significantly improves the performance of Sparse MobileNets on ImageNet-2012 dataset and exceeds the pruning results reported by Zhu & Gupta (2018). (right) Performance of sparse MobileNet-v1 architectures presented with their inference FLOPs. Networks with ERK distribution get better performance with the same number of parameters but take more FLOPs to run. Training wider sparse models with RigL (Big-Sparse) yields a significant performance improvement over dense model even though both have same number of parameters and FLOPs.

To demonstrate the advantages of sparse models we train wider sparse MobileNets while keeping the FLOPs and total number of parameters the same as the dense baseline. A sparse MobileNet v1 with width multiplier 1.98 and constant sparsity has the same FLOP and parameter count as the dense baseline. This yields a 3.8% absolute improvement in Top-1 Accuracy.

4.3 Analyzing the Performance of RigL

In this section we study the effect of sparsity distributions, update schedules, and dynamic connections on the performance of our method. The results for SET and SNFS are similar and are discussed in Appendices A and C.

Effect of Mask Initialization: Figure 4-left shows how the sparsity distribution affects the final test accuracy of sparse ResNet-50s trained with RigL. Erdős-Rényi-Kernel (ERK) performs significantly better than the other two initializations overall. Recall that uniform distribution keeps the first layer dense and has a fixed sparsity for the last dense layer. We scale the uniform sparities so that the overall parameter count is the same across all distributions considered. Though performing better, ERK distribution requires around 2x FLOPs to run compared to uniform. This highlights an interesting trade-off between accuracy and computational efficiency.

Effect of Update Schedule and Frequency: In Figure 4-right, we evaluate the performance of our method on update intervals $\Delta I \in [50, 100, 500, 1000]$ and initial drop fractions $\alpha \in [0.1, 0.3, 0.5]$. Even when we update the sparse connectivity every 1000 iterations, RigL performs above 72.5%.

Figure 4: (left) Performance of RigL at different sparsities using different sparsity masks (right) Ablation study on cosine schedule. Other methods are in the appendix.
Figure 5: (left) Training loss evaluated at various points on interpolation curves between a magnitude pruning model (0.0) and a model trained with static sparsity (1.0). (right) Training loss of RigL and Static methods starting from the static sparse solution, and their final accuracies.

**Effect of Dynamic connections:** Frankle et al. (2019) and Mostafa & Wang (2019) observed that static sparse training converges to a solution with a higher loss than dynamic sparse training. In Figure 5-left we examine the loss landscape lying between a solution found via static sparse training and a solution found via dynamic sparse training. Performing a linear interpolation between the two reveals a high-loss barrier which may be insurmountable with a fixed topology. Following Garipov et al. (2018), we attempt to find quadratic and cubic Bézier curves between the two solutions. Surprisingly, even with a cubic curve, we fail to find a path without a high-loss barrier. These results suggest that static sparse training can get stuck at local minima that are isolated from improved solutions. On the other hand, when we optimize the quadratic Bézier curve across the full dense space we find a near-monotonic path to the improved solution, suggesting that allowing new connections to grow lends dynamic sparse training greater flexibility in navigating the loss landscape. In Figure 5-right we train RigL starting from the sub-optimal solution found by static sparse training, demonstrating that it is able to escape the local minimum, whereas re-training with static sparse training cannot.

5 DISCUSSION & CONCLUSION

In this work we introduced ‘Rigged Lottery’ or RigL, an algorithm for training sparse neural networks efficiently. For a given computational budget RigL achieves higher accuracies than existing dense-to-sparse and sparse-to-sparse training algorithms. RigL is useful in three different scenarios: (1) To improve the accuracy of sparse models intended for deployment; (2) To improve the accuracy of large sparse models which can only be trained for a limited number of iterations; and (3) Combined with sparse primitives to enable training of extremely large sparse models which otherwise would not be possible.

The third scenario is unexplored due to the lack of hardware and software support for sparsity. Nonetheless, work continues to improve the performance of sparse networks on current hardware [Hong et al. 2019] [Merrill & Garland, 2016], and new types of hardware accelerators will have better support for parameter sparsity [Wang et al. 2018] [Mike Ashby, 2019] [Liu et al. 2018] [Han et al. 2016a]. RigL provides the tools to take advantage of, and motivation for, such advances.

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### A Effect of Sparsity Distribution on Other Methods

In Figure 6-left we show the effect of sparsity distribution choice on 4 different sparse training methods. The ordering among the three distribution is the same across different methods.

### B Effect of Momentum Coefficient for SNFS

In Figure 6-right we show the effect of the momentum coefficient on the performance of SNFS. Our results suggest that the accumulated values are not important since setting the coefficient to 0 brings...
the best performance. This surprising result might be due to the large batch size we are using (4096), but it still motivates using RigL and instantaneous gradient information only when needed, instead of accumulating them.

C  Effect of Update Schedules on Other Dynamic Sparse Methods

In Figure 7 we repeat the hyper-parameter sweep done for RigL in Figure 4-right, but for SET and SNFS. A cosine schedule with $\Delta I = 50$ and $\alpha = 0.1$ seems to work best across all methods. For this reason, we used these values in the experiments presented in the main text. An interesting observation is that higher drop fractions ($\alpha$) seem to work better with longer intervals $\Delta I$. For example, SET with an update interval of 1000 seems to work quite well with a starting drop fraction $\alpha = 0.5$.

D  Trying Different Update Schedules

In Figure 8 we share the results of using two alternative annealing functions for the update fractions:

1. **Constant**: $f_{\text{decay}}(t) = \alpha$.
2. **Inverse Power**: The fraction of weights updated decreases similarly to the schedule used in Zhu & Gupta (2018): $f_{\text{decay}}(t) = \alpha (1 - \frac{t}{I_{\text{end}}})^k$. In our experiments we tried $k = 1$ which is the linear decay and their default $k = 3$.

*Constant* seems to perform well with low initial drop fractions like $\alpha = 0.1$, but it starts to perform worse with increasing $\alpha$. *Inverse Power* for $k=3$ and $k=1$ (*Linear*) seems to perform similarly for low $\alpha$ values. However, the performance drops noticeably for $k=3$ when we increase the update interval. As reported by Dettmers & Zettlemoyer (2019) linear ($k=1$) seems to provide similar results as the cosine schedule.

E  Calculating FLOPs of Models and Methods

In order to calculate FLOPs needed for a single forward pass of a sparse model, we count the total number of multiplications and additions layer by layer for a given layer sparsity $s^l$. The total FLOPs is then obtained by summing up all of these multiply and adds.

Different sparsity distributions require different number of FLOPs to compute a single prediction. For example Erdős-Renyi-Kernel distributions usually cause earlier layers to be less sparse than the later layers (see Appendix G). The inputs of earlier layers have greater spatial dimensions, so a convolutional kernel that works on such inputs will require more FLOPs to compute the output features compared to later layers. Thus, having earlier layers which are less sparse results in a higher total number of FLOPs required by a model.

![Figure 7: Cosine update schedule hyper-parameter sweep done using dynamic sparse training methods SET (left) and SNFS (right).](image-url)
Training a neural network consists of 2 main steps:

1. **forward pass:** Calculating the loss of the current set of parameters on a given batch of data. During this process layer activations are calculated in sequence using the previous activations and the parameters of the layer. Activation of layers are stored in memory for the backward pass.

2. **backward pass:** Using the loss value as the initial error signal, we back-propagate the error signal while calculating the gradient of parameters. During the backward pass each layer calculates 2 quantities: the gradient of the activations of the previous layer and the gradient of its parameters. Therefore in our calculations we count backward passes as two times the computational expense of the forward pass. We omit the FLOPs needed for batch normalization and cross entropy.

Dynamic sparse training methods require some extra FLOPs to update the connectivity of the neural network. We omit FLOPs needed for dropping the lowest magnitude connections in our calculations. For a given dense architecture with FLOPs $f_D$ and a sparse version with FLOPs $f_S$, the total FLOPs required to calculate the gradient on a single sample is computed as follows:

- **Static Sparse and Dense.** Scales with $3 * f_S$ and $3 * f_D$ FLOPs, respectively.

- **Snip.** We omit the initial dense gradient calculation since it is negligible, which means Snip scales in the same way as Static methods: $3 * f_S$ FLOPs.

- **SET.** We omit the extra FLOPs needed for growing random connections, since this operation can be done on chip efficiently. Therefore, the total FLOPs for SET scales with $3 * f_S$.

- **SNFS.** Forward pass and back-propagating the error signal needs $2 * f_S$ FLOPs. However, the dense gradient needs to be calculated at every iteration. Thus, the total number of FLOPs scales with $2 * f_S + f_D$.

- **RigL.** Iterations with no connection updates need $3 * f_S$ FLOPs. However, at every $\Delta I$ iteration we need to calculate the dense gradients. This results in the average FLOPs for RigL given by $\frac{(3 * f_S + 2 * f_S + f_D) * \Delta I}{\Delta I + 1}$.

## F Effect of Mask Updates on the Energy Landscape

To update the connectivity of our sparse network, we first need to drop a fraction $d$ of the existing connections for each layer independently to create a budget for growing new connections. Following the recipe of magnitude based pruning (Han et al., 2015), we order parameters at layer $i$ by magnitude $|\theta_i|$ and drop the $N_i * (1 - s) * d$ parameters with lowest magnitude. The effectiveness of this simple criteria can be explained through the first order Taylor approximation of the loss $L$ around the current set of parameters $\theta$.

$$
\Delta L = L(\theta + \Delta \theta) - L(\theta) = \nabla_\theta L(\theta) \Delta \theta + R(||\Delta \theta||^2)
$$

The main goal of dropping connections is to remove parameters with minimal impact on the function and therefore on its performance. Since removing a connection corresponds to setting it to
zero, removing the parameter $\theta_i$ incurs a change of $\Delta \theta = -\theta_i$ in the parameters and a change of $\Delta L_i = -\nabla_{\theta_i} \bar{L}(\theta)\theta_i + R(\theta^2_i)$ in the loss, where the first term is usually defined as the saliency of a connection. Though using saliency to remove connections has been used as a criteria for removing connections (Molchanov et al., 2016), it has been shown to produce inferior results compared to magnitude based removal, especially when used to remove multiple connections at once (Evci, 2018). In contrast, picking the lowest magnitude connections ensures a small remainder term in addition to a low saliency, limiting the damage we make to the network while dropping connections. In other words, dropped connections are likely to be connections which don’t affect the output of the neural network.

After removal of these connections, we add new connections with the highest expected gradients. Note that although we initialize the added connections to zero, they are guaranteed to have a high gradient in the first iteration after the mask update. A direction of non-zero gradient means that the energy landscape on that dimension is not flat.

G  SPARSITY OF INDIVIDUAL LAYERS FOR SPARSE RESNET-50

Sparsity of ResNet-50 layers defined by the Erdős-Rényi-Kernel sparsity distribution plotted in Figure 9
Figure 9: Sparsities of individual layers of the ResNet-50.