PHEW: Constructing Sparse Networks that Learn Fast and Generalize Well Without Training Data

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Abstract

Methods that sparsify a network at initialization are important in practice because they greatly improve the efficiency of both learning and inference. Our work is based on a recently proposed decomposition of the Neural Tangent Kernel (NTK) that has decoupled the dynamics of the training process into a data-dependent component and an architecture-dependent kernel – the latter referred to as Path Kernel. That work has shown how to design sparse neural networks for faster convergence, without any training data, using the Synflow-L2 algorithm. We first show that even though Synflow-L2 is optimal in terms of convergence, for a given network density, it results in sub-networks with “bottleneck” (narrow) layers – leading to poor performance as compared to other data-agnostic methods that use the same number of parameters. Then we propose a new method to construct sparse networks, without any training data, referred to as Paths with Higher-Edge Weights (PHEW). PHEW is a probabilistic network formation method based on biased random walks that only depends on the initial weights. It has similar path kernel properties as Synflow-L2 but it generates much wider layers, resulting in better generalization and performance. PHEW achieves significant improvements over the data-independent SynFlow and SynFlow-L2 methods at a wide range of network densities.

1. Introduction

Generating sparse neural networks through pruning has recently led to a major reduction in the number of parameters, while having minimal loss in performance. Conventionally, pruning methods operate on pre-trained networks. Generally, such methods use an edge scoring mechanism for eliminating the less important connections. Popular scoring mechanisms include weight magnitudes (Han et al., 2015b; Janowsky, 1989; Park et al., 2020), loss sensitivity with respect to units (Mozer & Smolensky, 1989) and with respect to weights (Karlin, 1990), Hessian (LeCun et al., 1990; Hassibi & Stork, 1993), and first and second order Taylor expansions (Molchanov et al., 2019b;a). More recent approaches use more sophisticated variants of these scores (Han et al., 2015a; Guo et al., 2016; Carreira-Perpinán & Idelbayev, 2018; Yu et al., 2018; Dong et al., 2017; Guo et al., 2016).

Further analysis of pruning has shown the existence of sparse subnetworks at initialization which, when trained, are capable of matching the performance of the fully-connected network (Frankle & Carbin, 2018; Frankle et al., 2019; Liu et al., 2019; Frankle et al., 2020). However, identifying such “winning ticket” networks requires expensive training and pruning cycles. More recently, SNIP (Lee et al., 2019b), (You et al., 2019) and GraSP (Wang et al., 2019) showed that it is possible to find “winning tickets” prior to training – but still having access to at least some training data to compute initial gradients. Furthermore, other work has shown that such subnetworks generalize well across datasets and tasks (Morcos et al., 2019).

Our goal is to identify sparse subnetworks that perform almost as well as the fully connected network without any training data. The closest methods that tackle the same problem with our work are SynFlow (Tanaka et al., 2020) and SynFlow-L2 (Gebhart et al., 2021). The authors of (Tanaka et al., 2020) introduced the concept of “layer collapse” in pruning – the state when all edges in a layer are eliminated while there are edges in other layers that can be pruned. They also proved that iterative pruning based on positive gradient-based scores avoids layer collapse and introduced an iterative algorithm (SynFlow) and a loss function that conserves information flow and avoids layer collapse.

A branch of recent work focuses on the convergence and generalization properties of deep neural networks using linear approximations of the training dynamics (Jacot et al., 2018; Lee et al., 2019a; Arora et al., 2019). Under the infinite-width assumption, (Jacot et al., 2018) showed how
to predict at initialization output changes during training using the Neural Tangent Kernel (NTK). More recently, (Gebhart et al., 2021) decomposed the NTK into two factors: one that is only data-dependent, and another that is only architecture-dependent. This decomposition decoupled the effects of network architecture (including sparsity and selection of initial weights) from the effect of training data on convergence. The architecture-dependent factor can be thought of as the “path covariance” of the network and is referred to as Path Kernel. The authors of (Gebhart et al., 2021) show that the training convergence of a network can be accurately predicted using the path kernel trace. That work concluded with a pruning algorithm (SynFlow-L2) that designs sparse networks with maximum path kernel trace – aiming to optimize at least the architectural component of the network’s convergence speed.

In this work, we first show that even though SynFlow and Synflow-L2 are optimal in terms of convergence for a given network density, they result in sub-networks with “bottleneck layers” (very small width) – leading to poor performance as compared to other data-agnostic methods that use the same number of parameters. This issue is observed even at moderate density values. This is expected given the recent results of (Golubeva et al., 2021), for instance, showing that increasing the width of sparse networks, while keeping the number of parameters constant, generally improves performance.

We then present a method, referred to as PHEW (Paths with Higher Edge Weights), which aims to achieve the best of both worlds: high path kernel trace for fast convergence, and large network width for better generalization performance. Given an unpruned initialized network, and a target number of learnable parameters, PHEW selects a set of input-output paths that are conserved in the network and it prunes every remaining connection. The selection of the conserved paths is based strictly on their initial weight values – and not on any training data. Further, PHEW induces randomness into the path selection process using random walks biased towards higher weight-magnitudes. The network sparsification process does not require any data, and the pruned network needs to be trained only once.

We show that selecting paths with higher edge weights forms sub-networks that have higher path kernel trace than uniform random walks – close to the trace obtained through SynFlow-L2. We also show that the use of random walks results in sub-networks having high per-layer width – similar to that of unpruned networks. Further, PHEW avoids layer-collapse by selecting and conserving input-output paths instead of individual units or connections. We compare the performance of PHEW against several pruning before-training methods and show that PHEW achieves significant improvements over SynFlow and SynFlow-L2. Additionally, we conduct a wide range of ablation studies to evaluate the efficacy of PHEW.

2. Background

2.1. Neural Tangent Kernel

Let \( (\mathcal{X}, \mathcal{Y}) \) denote the training examples, \( \mathcal{L} \) the loss function, and \( f(\mathcal{X}, \theta) \in \mathbb{R}^{NK} \) the network’s output, where \( N \) is the number of examples and \( K \) is the output dimension. Under the gradient flow assumption, and denoting the learning rate by \( \eta \), the output of the network at time \( t \) can be approximated using the first-order Taylor expansion,

\[
f(\mathcal{X}, \theta_{t+1}) = f(\mathcal{X}, \theta_t) - \eta \Theta(t)(\mathcal{X}, \mathcal{X}) \nabla f(\mathcal{X}, \theta_t)\]

where the matrix \( \Theta(t)(\mathcal{X}, \mathcal{X}) = \nabla f(\mathcal{X}, \theta_t) \nabla f(\mathcal{X}, \theta_t)^T \in \mathbb{R}^{NK \times NK} \) is the Neural Tangent Kernel (NTK) at time \( t \) (Jacot et al., 2018). Under the additional assumption of infinite width, the NTK has been shown to remain constant throughout training, and it allows us to exactly predict the evolution of the network’s output. More recent work has shown that even networks with limited width, and any depth, closely follow the NTK dynamics (Lee et al., 2019a). For a constant NTK \( \Theta_0 \), and with a mean-squared error loss, equation (1) has the closed-form solution:

\[
f(\mathcal{X}, \theta_t) = (\mathcal{I} - e^{-\eta \Theta_0}) \mathcal{Y} + e^{-\eta \Theta_0} f(\mathcal{X}, \theta_0)
\]

Equation (2) allows us to predict the network’s output given the input-output training examples, the initial weights \( \theta_0 \), and the initial NTK \( \Theta_0 \). Further, leveraging equation (2), it has been shown that the training convergence is faster in the directions that correspond to the larger NTK eigenvalues (Arora et al., 2019). This suggests that sparse sub-networks that preserve the larger NTK eigenvalues of the original network would converge faster and with higher sampling efficiency (Wang et al., 2019).

2.2. Path kernel decomposition of NTK

More recently, an interesting decomposition of the Neural Tangent Kernel has been proposed that decouples the effects of the network architecture (and initial weights) from the data-dependent factors of the training process (Gebhart et al., 2021). We summarize this decomposition next.

Consider a neural network \( f : \mathbb{R}^D \rightarrow \mathbb{R}^K \) with ReLU activations, parametrized by \( \theta \in \mathbb{R}^m \). Let \( \mathcal{P} \) be the set of all input-output paths, indexed as \( p = 1, \ldots, P \) (we refer to a path by its index \( p \)). Let \( p_i = 1 \{ \theta_i \in p \} \) represent the presence of edge-weight \( \theta_i \) in path \( p \).

The edge-weight-product of a path is defined as the product of edge-weights present in a path, \( \pi_p(\theta) = \prod_{i=1}^m \theta_i^{p_i} \). For an input variable \( x \), the activation status of a path is, \( a_p(x) = \prod_{i \in p} \mathbb{I}\{z_i > 0\} \), where \( z_i \) is the activation of the
neuron connected to the previous layer through $\theta_i$. The $k^{th}$ output of the network can be expressed as:

$$f^k(x, \theta) = \sum_{i=1}^{D} \sum_{p \in \mathcal{P}_{i \rightarrow k}} \pi_p(\theta) \alpha_p(x) x_i,$$

where $\mathcal{P}_{i \rightarrow k}$ is the set of paths from input unit $i$ to output unit $k$. We can now decompose the NTK using the chain rule:

$$\Theta(\mathcal{X}, \mathcal{X}) = \nabla_{\theta} f(\mathcal{X}) \nabla_{\theta} \pi(\theta) \nabla_{\theta} \pi(\theta)^T \nabla_{\theta} f(\mathcal{X})^T$$

$$= J^2_{\pi}(\mathcal{X}) J^0_{\theta} (J^0_{\pi})^T J^2_{\pi}(\mathcal{X})^T$$

$$= J^2_{\pi}(\mathcal{X}) \Pi_{\theta} J^2_{\pi}(\mathcal{X})^T$$

The matrix $\Pi_{\theta}$ is referred to as the Path Kernel (Gebhart et al., 2021). The path kernel element for two paths $p$ and $p'$ is:

$$\Pi_{\theta}(p, p') = \sum_{i=1}^{m} \left( \frac{\pi_p(\theta)}{\theta_i} \right) \left( \frac{\pi_{p'}(\theta)}{\theta_i} \right) p_i p'_i$$

Note that the path kernel, $\Pi_{\theta} \in \mathbb{R}^{P \times P}$, depends only on the network architecture and the initial weights. On the other hand, the matrix $J^2_{\pi}(\mathcal{X}) \in \mathbb{R}^{NK \times P}$ captures the data-dependent activations and re-weights the paths on the basis of the training data input.

**Convergence approximation:** The eigenstructure of the NTK depends on how the eigenvectors of $J^2_{\pi}(\mathcal{X})$ map onto the eigenvectors of the path kernel $\Pi_{\theta}$, as shown by the following result.

**Theorem 1** (Gebhart et al., 2021): Let $\lambda_i$ be the eigenvalues of $\Theta(\mathcal{X}, \mathcal{X})$, $v_i$ the eigenvalues of $J^2_{\pi}(\mathcal{X})$ and $w_i$ the eigenvalues of $\Pi_{\theta}$. Then $\lambda_i \leq v_i w_i$ and $\sum_i \lambda_i \leq \sum_i v_i w_i$.

Given the eigenvalue decomposition of $\Theta_{\theta}$, Theorem 1 provides an upper bound for the convergence in equation (2). $\Theta_{\theta}$ with eigenvalues $\lambda_i$ has the same eigenvectors as $e^{-\lambda_i t}$ with eigenvalues $e^{-\lambda_i t}$. Therefore, $\sum_i v_i w_i$ accurately captures the eigenvalues of $\Theta_{\theta}$ and it can be used to predict the convergence of the training process. Even without any training data, the convergence can be effectively approximated from the trace of the path kernel:

$$Tr(\Pi_{\theta}) = \sum_i w_i = \sum_{p=1}^{P} \Pi_{\theta}(p, p) = \sum_{p=1}^{P} \sum_{i=1}^{m} \left( \frac{\pi_p(\theta)}{\theta_i} \right)^2 p_i$$

The authors of (Gebhart et al., 2021) empirically validated the convergence predicted using the trace of the path kernel against the actual training convergence of the network.

The previous result has an important consequence for neural network pruning. Given a fully connected neural network at initialization as well as the target density for a pruned network, maximizing the path kernel trace of the pruned network preserves the largest NTK eigenvalues of the original network. Since, the directions corresponding to the larger eigenvalues of the NTK learn faster, the sub-network obtained by maximizing the path kernel trace is also expected to converge faster and learn more efficiently.

### 2.3. SynFlow-L1 and SynFlow-L2

The path kernel framework has been applied in the design of pruning algorithms that do not require any training data (Gebhart et al., 2021; Tanaka et al., 2020). SynFlow-L2 is such an iterative pruning algorithm that removes edges (parameters) based on the following saliency function:

$$S(\theta_i) = \theta_i \odot \frac{\partial R(\theta)}{\partial \theta_i} = \theta_i \odot \sum_{p=1}^{P} \left( \frac{\pi_p(\theta)}{\theta_i} \right)^2 p_i$$

The process of computing the previous saliency measure and eliminating edges with lower saliency is repeated until the required density is achieved. SynFlow-L2 maximizes the trace of the path kernel, and preserves the following data-independent loss function:

$$R(\theta) = 1^T \left( \prod_{i=1}^{L+1} \theta_i |^2 \right) 1 = \sum_{p=1}^{P} (\pi_p(\theta))^2$$

where $|\theta_i|^2$ is the matrix formed by the squares of the elements of the weight matrix at the $l^{th}$ layer and $L$ is the number of hidden layers.
We can also observe empirically in Figure 1 that SynFlow-L2 achieves the highest path kernel trace compared to other state-of-the-art pruning methods.

Another related pruning method is SynFlow-L1 (or simply “SynFlow”) – proposed in (Tanaka et al., 2020). SynFlow is based on preserving the loss function $R(\theta) = \sum_{p=1}^{P} |\pi_p(\theta)|$, which is based on the edge-weight products along each input-output path (rather than their squares).

### 3. Pruning for maximum path kernel trace

In this section we analyze the resulting architecture of a sparse network that has been pruned to maximize the path kernel trace. As discussed in the previous section, SynFlow-L2 is a pruning method that has this objective.

Consider a network with a single hidden-layer, $f : \mathbb{R}^D \rightarrow \mathbb{R}^D$, with $N$ hidden units and $D$ inputs and outputs. The incoming and outgoing weights $\theta$ of each unit are initialized by sampling from $\mathcal{N}(0, 1)$. Let the number of connections in the unpruned network be $M$, and let $m$ be the target number of connections in the pruned network, so that the resulting network density is $\rho = m/M$.

The optimization of the path kernel trace selects the $m$ out of $M$ parameters that maximize:

$$\sum_{p=1}^{P} \sum_{i=1}^{m} \left( \frac{\pi_p(\theta)}{\theta_i} \right)^2 p_i \quad (9)$$

In Appendix A.1, we show that this maximization results in a fully-connected network in which only $n \leq N$ of the hidden-layer units remain in the pruned network – all other units and their connections are removed. In other words, the network that maximizes the path kernel trace has the narrowest possible hidden-layer width, given a target network density.

We also show (Appendix A.2) that this network architecture maximizes the number of input-output paths $P$ : Given a target density $\rho$, the maximum number of paths results when each hidden-layer has the same number of units, and the network is fully-connected.

Intuitively, the previous results can be justified as follows, with the same weight distribution across all units of a layer, increasing the number of input-output paths $P$ results in higher path kernel trace. To maximize $P$ with a given number of edges $m$, however, forces the pruning process to only maintain the edges of the smallest possible set of units at each layer. So, the networks produced by SynFlow and SynFlow-L2 tend to have narrower layers, compared to other pruning methods that do not optimize on the basis of path kernel trace.

To examine the previous claim empirically, and in the context of convolutional networks rather than MLPs, Figure 2 compares the number of remaining units at each layer after pruning, using the VGG19 and ResNet20 architectures. The target network density in these experiments is the lowest possible such that the method of Magnitude-Pruning (that can be performed only after training) achieves within 5% of the unpruned network’s accuracy. In higher densities there is still significant redundancy, while in lower densities there is no sufficient capacity to learn the given task. For a convolutional layer, the width of a layer is the number of channels at the output of that layer. We find that both SynFlow and SynFlow-L2 result in pruned networks with very small width (“bottleneck layers”) compared to other state-of-the-art pruned networks of the same density. Further, with SynFlow and SynFlow-L2 all layers have approximately the same number of remaining units, i.e., approximately equal width. Note that for the purposes of this analysis (Figure 2), we do not include skip connections for ResNet20 – such connections complicate the definition of “layer width” and paths, but without changing the main result of Figure 2.

\[1\] In SNIP, the widest layers get pruned more aggressively as showed in (Tanaka et al., 2020). Due to this SNIP also leads to a decrease in width, but only at the widest layers.

\[2\] GraSP and PHEW are able to preserve the same width as the unpruned network for all the layers. The curves for GraSP (green) and PHEW (blue) overlap with the curve for the unpruned network in Figure 2.
Figure 3. The effect of increasing the layer width of SynFlow and SynFlow-L2 networks, while preserving the same set of parameters at each layer. The definition of the x-axis “Width Factor” appears in the main text.

3.1. Effect of network width on performance

Several empirical studies have been conducted to understand the effect of network width and over-parametrization on learning performance (Neyshabur et al., 2018; Du et al., 2018; Park et al., 2019; Lu et al., 2017). However, the previous studies do not decouple the effect of increasing width from the effect of over-parametrization. Recently, (Golubeva et al., 2021) examined the effect of network width under a constant number of parameters. That work conducted experiments with layer-wise random pruning. Starting with a fully-connected network, the width of each layer is increased while keeping the number of parameters the same. The experiments of (Golubeva et al., 2021) show that as the network width increases the performance also increases. Further, the distance between the Gaussian kernel formed by the sparse network and the infinitely wide kernel at initialization is indicative of the network’s performance. As expected, increasing the width after a certain limit without also increasing the number of parameters will inevitably cause a drop in both test and train accuracy because of very low per-unit connectivity (especially with random pruning).

We present similar experiments for SynFlow and SynFlow-L2 in Figure 3. For a given network density, we first obtain the layer-wise density and number of active units that result from the previous two pruning algorithms. We then gradually increase the number of active units by randomly shuffling the masks of each layer (so that the number of weights at each layer is preserved). The increase in layer width can be expressed as the fraction $x = (w' - w) / (W - w)$, where $W$ is the layer width of the unpruned network, $w$ is the layer width in the Synflow (or Synflow-L2) pruned network, and $w' \geq w$ is the layer width that results through the shuffling method described above. The maximum value $x = 1$ results when $w' = W$.

Figure 3 shows the results of these models on CIFAR-10/100 tasks using ResNet20 and VGG19. We can see that as the width increases so does the performance of the sparse network, even though the layer-wise number of edges is the same. Similar results appear in the ablation studies of (Frankle et al., 2021) using SynFlow. That study redistributes the edges of a layer, creating a uniform distribution across all units in the layer – doing so increases the performance of the network (see Appendix C).

Summary: Let us summarize the observations of this section regarding the maximization of the path kernel trace – and the resulting decrease in network width. Even without any training data, pruned networks that result by maximizing the path kernel trace are expected to converge faster and learn more efficiently. As we showed however, for a given density, such methods tend to maximize the number of input-output paths, resulting in pruned networks with very narrow layers. Narrow networks, however, attain lower performance as compared to wider networks of the same layer-wise density. In the next section, we present a method that aims to achieve the best of both worlds: high path kernel trace for fast convergence, and large layer-wise width for better generalization and learning.

4. The PHEW network sparsification method

Given a weight-initialized architecture, and a target number of learnable parameters, we select a set of input-output paths that are conserved in the network – and prune every connection that does not appear in those paths. The selection of conserved paths is based strictly on their initial weights – not on any training data. The proposed method is called “Paths with Higher Edge Weights” (PHEW) because it has a bias in favor of higher weight connections. Further, the path selection is probabilistic, through biased random walks from input units to output units. Specifically, the next-hop of each path, from unit $i$ to $j$ at the next layer, is taken with a probability that is proportional to the weight magnitude of the connection from $i$ to $j$. We show that conserving paths with higher edge weight product results in higher path kernel trace. The probabilistic nature of PHEW avoids the creation of “bottleneck layers” and leads to larger network width than methods with similar path kernel trace. Additionally, the procedure of selecting and conserving input-output paths completely avoids layer collapse.

In more detail, let us initially consider a fully-connected
MLP network with $L$ hidden layers and $N_l$ units at each layer (we consider convolutional networks later in this section). Suppose that the weights are initialized according to Kaiming’s method (He et al., 2015), i.e., they are sampled from a Normal distribution in which the variance is inversely proportional to the width of each layer: $\theta_{i,j} \sim \mathcal{N}(0, \sigma_l^2)$, where $\sigma_l^2 = 2/N_l$.

First, let us consider two input-output paths $u$ and $b$: $u$ has been selected via a uniform random-walk in which the probability $Q(j,i)$ that the walk moves from unit $i$ to unit $j$ at the next layer is the same for all $j$; $b$ has been selected via the following weight-biased random-walk process:

$$Q(j,i) = \frac{[\theta(j,i)]}{\sum_j[\theta(j,i)]}$$

where $\theta(j,i)$ is the weight of the connection from $i$ to $j$.

In Appendix A.4 we show that the biased-walk path $b$ contributes more in the path kernel trace than path $u$:

$$\mathbb{E}[\Pi_\theta(b,b)] = 2^L \times \mathbb{E}[\Pi_\theta(u,u)]$$

As the number of hidden layers $L$ increases the ratio between the two terms becomes exponentially higher. On the other hand, as the layer’s width increases the ratio of two values remains the same. The reason that PHEW paths result in higher path kernel trace, compared to the same number of uniformly chosen paths, is that the former tend to have higher edge weights, and thus higher $\pi_p(\theta)$ values (see Equation 6). Empirically, Figure 1 shows that PHEW achieves a path kernel trace greater than or equal to SNIP and GraSP, and close to the upper bound of SynFlow-L2.

If the PHEW paths were chosen deterministically (say in a greedy manner, always taking the next hop with the highest weight) the path kernel trace would be slightly higher but the resulting network would have “bottlenecks” at the few units that have the highest incoming weights. PHEW avoids this problem by introducing randomness in the path selection process. Specifically, in Appendix A.3 we show that the expected number of random walks through each unit of a layer $l$ is $W/N_l$, where $W$ is the required number of walks to achieve the target network density. Thus, as long as $W > N_l$, every unit is expected to be traversed by at least one walk – and thus every unit of that layer is expected to be present in the sparsified network.

This is very different than the behavior of SynFlow or SynFlow-L2, in which the width of several layers in the pruned network is significantly reduced. Empirically, Figure 2 confirms that PHEW achieves the larger per-layer width, compared to SynFlow and SynFlow-L2. Additionally, the per-layer width remains the same as the width of the original unpruned network.

Layer-Collapse: Layer collapse is defined as a network state in which all edges of a specific layer are eliminated, while there are still connections in other layers (Tanaka et al., 2020). Layer collapse causes a disruption of information flow through the sparse network making the network untrainable. SynFlow and SynFlow-L2 have been shown to avoid layer collapse by iteratively computing gradient based importance scores and pruning (Tanaka et al., 2020). PHEW also avoids layer collapse due to its path-based selection and conservation process. Even a single input-output path has one connection selected at each layer, and so it is impossible for PHEW networks to undergo layer collapse.

4.1. Additional PHEW details

Balanced, bidirectional walks: Without any information about the task or the data, the only reasonable prior is to assume that every input unit is equally significant – and the same for every output unit. For this reason, PHEW attempts to start the same number of walks from each input. And to terminate the same number of walks at each output.

To do so, we create paths in both directions with the same probability: forward paths from input units, and reverse paths from output units. The selection of the starting unit in each case is such that the number of walks that start (or terminate) at each input (or output) unit is approximately the same. The creation of random-walks continues until we have reached the given, target number of parameters.

PHEW in convolutional neural networks: A convolutional layer takes as input a 3D-vector with $n_i$ channels and transforms it into another 3D-vector of $n_{i+1}$ channels. Each of the $n_{i+1}$ units in a layer produces a single 2D-channel corresponding to the $n_{i+1}$ channels. A 2D channel is produced applying convolution on the input vector with $n_i$ channels, using a 3D-filter of depth $n_i$. Therefore each input from a unit at the previous layer has a corresponding 2D-kernel as one of the channels in the filter. So, even though MLPs have an individual weight per edge, convolutional networks have a 2D-kernel per edge.

A random-walk can traverse an edge of a convolutional network in two ways: either traversing a single weight in the corresponding 2D kernel – or traversing the entire kernel with all its weights. Traversing a single weight from a kernel conserves that edge and produces a non-zero output channel. This creates sparse kernels and allows for the processing of multiple input channels at the same unit and with fewer parameters. On the other hand, traversing the entire 2D-kernel that corresponds to an edge means that several other kernels will be eliminated. Earlier work in pruning has shown empirically the higher performance of creating sparse kernels instead of pruning entire kernels (Blalock et al., 2020; Liu et al., 2019). Therefore, in PHEW we choose to conserve individual parameters during a random-walk rather than conserving entire kernels.
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Figure 4. Comparison of the Top-1 accuracy for sparse networks obtained using PHEW and other state-of-the-art data-independent baselines. The mean is shown as a solid line while the standard deviation is shown as shaded area.

In summary, PHEW follows a two-step process in convolutional networks: first an edge (i.e., 2D-kernel) is selected using equation (10). Then a single weight is chosen from that kernel, randomly, with a probability that is proportional to the weight of the sampled parameter. We have also experimented with the approach of conserving the entire kernel, and we also present results for that case in the next section.

5. Experimental results

In this section we present several experiments conducted to compare the performance of PHEW against state-of-the-art pruning methods and other baselines. We present results both for standard image classification tasks using convolutional networks as well as an image-transformation task using MLPs (that task is described in more detail in Appendix B). We also conduct a wide variety of ablation studies to test the efficacy of PHEW.

5.1. Classification comparisons

We compare PHEW against two baselines, Random pruning and Initial (Weight) Magnitude pruning, along with four state-of-the-art algorithms: SNIP (Lee et al., 2019b), GraSP (Wang et al., 2019), SynFlow (Tanaka et al., 2020) and SynFlow-L2 (Gebhart et al., 2021). We also present results for the original Unpruned network as well as for Gradual Magnitude Pruning (Zhu & Gupta, 2017) to upper bound the network’s performance for a given density. Because PHEW is data-independent, the most relevant comparison is with SynFlow and SynFlow-L2, which also do not require any training data.

We present results for three networks: ResNet20, VGG19 and ResNet18 and three datasets: CIFAR-10, CIFAR-100 and Tiny ImageNet. The network density range is chosen such that the magnitude pruning after training method (our performance upper bound) maintains a comparable accuracy to the unpruned network. We claim that it is not important in practice to consider densities in which any pruned network would perform poorly. The hyper-parameters used were tuned only for the unpruned network (see Appendix D). We run each experiment three times for different seed values and present the mean and standard deviation of the test accuracy.

Data-independent methods: We first compare PHEW against data-independent pruning baselines random pruning, initial magnitude pruning, and methods SynFlow and SynFlow-L2 in Figure 4. At the density levels considered, PHEW performs better than both SynFlow and SynFlow-L2. We attribute this superior performance to the large per-layer width of the sparse networks obtained through PHEW. As we showed earlier, differences in layer-wise width of sparse networks accounts for most of the performance differences, under the same number of parameters. Further, the performance gap increases for datasets with more classes, such as Tiny-ImageNet as compared to CIFAR-10 and CIFAR-100. We believe that this is due to the increased complexity of the dataset and the need for larger width to learn finer and
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Figure 5. Comparison of the Top-1 accuracy for sparse networks obtained using PHEW and other state-of-the-art data-dependent pruning methods and baselines. The mean is shown as a solid line while the standard deviation is shown as shaded area.

Disconnected decision regions (Nguyen et al., 2018).

Data-dependent methods: Figure 5 compares the test accuracy of PHEW networks against various data-dependent pruning algorithms. It is interesting that PHEW outperforms in many cases the data-dependent pruning algorithms GraSP and SNIP even though it is agnostic to the data or task.

At higher network densities, PHEW is competitive with SNIP. Note that SNIP also utilizes paths with higher weight magnitudes (Gebhart et al., 2021). At lower network densities, SNIP’s accuracy falls quickly while the accuracy of PHEW falls more gradually creating a significant gap. This is because SNIP eliminates units at a higher rate at the largest layer (Figure 2), which eventually leads to layer collapse (Tanaka et al., 2020).

GraSP on the other hand is competitive with PHEW at lower density levels and falls short of PHEW at moderate and higher network densities. This is because GraSP maximizes gradient flow after pruning, and the obstruction to gradient flow is prominent only at lower density levels. Further, it has been shown that GraSP performs better than SNIP only at lower density levels (Wang et al., 2019), which is consistent with our observations.

PHEW is able to perform relatively well across all density levels. At lower density levels, the obstruction to the flow of gradients is avoided by conserving input-output paths. Further, by selecting higher weight magnitudes, PHEW avoids the problem of vanishing gradients.

5.2. Ablation Studies

In this section we conduct a wide variety of ablation studies to test PHEW’s efficacy as well as to understand fully the cause behind its generalization performance.

Kernel-conserved PHEW variant: We study a PHEW variant for convolutional neural networks where instead of conserving a single weight of a kernel each time a random walk traverses that kernel, we conserve the entire kernel. This approach reduces the FLOP count immensely by eliminating the operations performed on several 2D feature maps in specific units. We present the comparison for CIFAR10 and CIFAR100 in Figure 6. Note that at moderate network densities, this kernel-conserved variant performs as well as the other methods we consider – therefore this variant can
be utilized when decreasing the FLOP count is a priority.

**Different weight initializations:** PHEW depends on initial weights, and so it is important to examine the robustness of PHEW’s performance across the major weight initialization methods used in practice: Kaiming (He et al., 2015), Normal $\mathcal{N}(0, 0.1)$, and Xavier uniform ((Glorot & Bengio, 2010)). Figure 6 shows results with such initializations for VGG19 and ResNet20 on CIFAR10 and CIFAR100. Note that PHEW’s performance is quite robust across all these weight initializations. It is interesting that PHEW’s performance is not altered by initializing all layers with the same distribution. This may be due to the random walk procedure, where the probability distribution at each hop is only dependent on the initial weights of that layer.

**Unbiased and “inverse-weight biased” random walks:** We also present two variants of PHEW: a) unbiased (uniform) walks, and b) random walks that are biased in favor of lower weight-magnitudes. The former selects the next-hop of the walk randomly, ignoring the weights, while the latter gives higher probability to lower weight-connections.

The two variants perform similar with PHEW in terms of accuracy, as shown in Figure 6, because they create networks that have the same per-layer width with PHEW.

Their difference with PHEW is in terms of convergence speed. The reason is that they are not biased in favor of higher-weight connections, and so the resulting path kernel trace is lower than that in PHEW. Indeed, we have also confirmed empirically that the training error drops faster initially (say between epochs 1 to 20) in PHEW than in these two variants.

**Re-initializing PHEW Networks:** We have also conducted experiments with reinitialized sparse networks selected through PHEW. Here, the architecture of the sparse network remains the same but the individual weights are re-sampled. This causes the path-kernel trace to be lower while the architecture produced by PHEW is maintained.

We present results for three reinitialization schemes, a) dense reinitialization, b) layer-wise sparse reinitialization (Liu et al., 2019) and c) neuron-wise sparse reinitialization (Evci et al., 2020).

We observe that reinitialized PHEW networks also achieve the same performance as the original PHEW network. This further confirms that the sparse architecture produced through the random walk process is sufficient to obtain PHEW’s generalization performance.

In Figure 7 we also compare with reinitialized randomly pruned networks, as those networks have been shown to improve performance. Although we see some improvements with sparse reinitialization in randomly pruned networks, the performance still falls short of PHEW and its variants.

6. Conclusion

We proposed a probabilistic approach called PHEW to construct sparse networks without any training data. Sparse networks that result by maximizing the path kernel trace are expected to converge faster. We showed that, for a given density, methods that maximize the path kernel trace result in very narrow layers and lower performance compared to wider networks of the same layer-wise density. On the other hand, conserving paths with higher edge-weight magnitudes leads to sparse networks with higher path kernel trace. Further, introducing randomness in the path selection process preserves the layer-wise width of the unpruned network. Empirically, we showed that PHEW achieves significant improvements over current data-independent state-of-the-art pruning at initialization methods.

Some open questions for future research are: 1) A comparison between PHEW networks and “winning tickets” (Frankle & Carbin, 2018), given the same number of parameters, both in terms of their convergence speed and structural properties. 2) Development of path-based network sparsification methods that can utilize a limited amount of training data to get even higher performance than PHEW. 3) How to identify path-based sparse networks that can perform as well as PHEW networks but without any training? 4) How to dynamically determine the optimal number of parameters in a sparse network at the early stages of training – instead of starting with a pre-determined target number of parameters?

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References


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