Loss Surfaces, Mode Connectivity, and Fast Ensembling of DNNs

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Abstract

The loss functions of deep neural networks are complex and their geometric properties are not well understood. We show that the optima of these complex loss functions are in fact connected by simple curves over which training and test accuracy are nearly constant. We introduce a training procedure to discover these high-accuracy pathways between modes. Inspired by this new geometric insight, we also propose a new ensembling method entitled Fast Geometric Ensembling (FGE). Using FGE we can train high-performing ensembles in the time required to train a single model. We achieve improved performance compared to the recent state-of-the-art Snapshot Ensembles, on CIFAR-10, CIFAR-100, and ImageNet.

1 Introduction

The loss surfaces of deep neural networks (DNNs) are highly non-convex and can depend on millions of parameters. The geometric properties of these loss surfaces are not well understood. Even for simple networks, the number of local optima and saddle points is large and can grow exponentially in the number of parameters [1, 2, 3]. Moreover, the loss is high along a line segment connecting two optima [e.g., 7, 13]. These two observations suggest that the local optima are isolated.

In this paper, we provide a new training procedure which can in fact find paths of near-constant accuracy between the modes of large deep neural networks. Furthermore, we show that for a wide range of architectures we can find these paths in the form of a simple polygonal chain of two line segments. Consider, for example, Figure 1 which illustrates the ResNet-164 ℓ2-regularized cross-entropy train loss on CIFAR-100, through three different planes. We form each two dimensional plane by all affine combinations of three weight vectors. The left panel shows a plane defined by three independently trained networks. In this plane, all optima are isolated, which corresponds to the standard intuition. However, the middle and right panels show two different paths of near-constant loss between the modes in weight space, discovered by our proposed training procedure. The endpoints of these paths are the two independently trained DNNs corresponding to the two lower modes on the left panel.

∗ Equal contribution.

1Suppose we have three weight vectors \(w_1, w_2, w_3\). We set \(u = (w_2 - w_1), v = (w_3 - w_1) - (w_3 - w_1, w_2 - w_1)/\|w_2 - w_1\|^2 \cdot (w_2 - w_1)\). Then the normalized vectors \(\hat{u} = u/\|u\|, \hat{v} = v/\|v\|\) form an orthonormal basis in the plane containing \(w_1, w_2, w_3\). To visualize the loss in this plane, we define a Cartesian grid in the basis \(\hat{u}, \hat{v}\) and evaluate the networks corresponding to each of the points in the grid. A point \(P\) with coordinates \((x, y)\) in the plane would then be given by \(P = w_1 + x \cdot \hat{u} + y \cdot \hat{v}\).
Figure 1: The $\ell_2$-regularized cross-entropy train loss surface of a ResNet-164 on CIFAR-100, as a function of network weights in a two-dimensional subspace. In each panel, the horizontal axis is fixed and is attached to the optima of two independently trained networks. The vertical axis changes between panels as we change planes (defined in the main text). **Left:** Three optima for independently trained networks. **Middle** and **Right:** A quadratic Bezier curve, and a polygonal chain with one bend, connecting the lower two optima on the left panel along a path of near-constant loss. Notice that in each panel a direct linear path between each mode would incur high loss.

We believe that this geometric discovery has major implications for research into multilayer networks, including (1) improving the efficiency, reliability, and accuracy of training, (2) creating better ensembles, and (3) deriving more effective posterior approximation families in Bayesian deep learning. Indeed, in this paper we are inspired by this geometric insight to propose a new ensembling procedure that can efficiently discover multiple high-performing but diverse deep neural networks. In particular, our contributions include:

- The discovery that the local optima for modern deep neural networks are connected by very simple curves, such as a polygonal chain with only one bend.
- A new method that finds such paths between two local optima, such that the train loss and test error remain low along these paths.
- Using the proposed method we demonstrate that such mode connectivity holds for a wide range of modern deep neural networks, on key benchmarks such as CIFAR-100. We show that these paths correspond to meaningfully different representations that can be efficiently ensembled for increased accuracy.
- Inspired by these observations, we propose Fast Geometric Ensembling (FGE), which outperforms the recent state-of-the-art Snapshot Ensembles [11], on CIFAR-10 and CIFAR-100, using powerful deep neural networks such as VGG-16, Wide ResNet-28-10, and ResNet-164. On ImageNet we achieve 0.56% top-1 error-rate improvement for a pretrained ResNet-50 model by running FGE for only 5 epochs.
- We release the code for reproducing the results in this paper at [https://github.com/timgaripov/dnn-mode-connectivity](https://github.com/timgaripov/dnn-mode-connectivity)

The rest of the paper is organized as follows. Section 2 discusses existing literature on DNN loss geometry and ensembling techniques. Section 3 introduces the proposed method to find the curves with low train loss and test error between local optima, which we investigate empirically in Section 4. Section 5 then introduces our proposed ensembling technique, FGE, which we empirically compare to the alternatives in Section 6. Finally, in Section 7 we discuss connections to other fields and directions for future work.

Note that we interleave two Sections where we make methodological proposals (Sections 3, 5), with two Sections where we perform experiments (Sections 4, 6). Our key methodological proposal for ensembling, FGE, is in Section 5.

## 2 Related Work

Despite the success of deep learning across many application domains, the loss surfaces of deep neural networks are not well understood. These loss surfaces are an active area of research, which falls into two distinct categories.

The first category explores the local structure of minima found by SGD and its modifications. Researchers typically distinguish sharp and wide local minima, which are respectively found by using
large and small mini-batch sizes during training. Hochreiter and Schmidhuber [10] and Keskar et al. [13], for example, claim that flat minima lead to strong generalization, while sharp minima deliver poor results on the test dataset. However, recently Dinh et al. [4] argue that most existing notions of flatness cannot directly explain generalization. To better understand the local structure of DNN loss minima, Li et al. [16] proposed a new visualization method for the loss surface near the minima found by SGD. Applying the method for a variety of different architectures, they showed that the loss surfaces of modern residual networks are seemingly smoother than those of VGG-like models.

The other major category of research considers global loss structure. One of the main questions in this area is how neural networks are able to overcome poor local optima. Choromanska et al. [2] investigated the link between the loss function of a simple fully-connected network and the Hamiltonian of the spherical spin-glass model. Under strong simplifying assumptions they showed that the values of the investigated loss function at local optima are within a well-defined bound. In other research, Lee et al. [14] showed that under mild conditions gradient descent almost surely converges to a local minimizer and not a saddle point, starting from a random initialization.

In recent work Freeman and Bruna [6] theoretically show that local minima of a neural network with one hidden layer and ReLU activations can be connected with a curve along which the loss is upper-bounded by a constant that depends on the number of parameters of the network and the “smoothness of the data”. Their theoretical results do not readily generalize to multilayer networks. Using a dynamic programming approach they empirically construct a polygonal chain for a CNN on MNIST and an RNN on PTB next word prediction. However, in more difficult settings such as AlexNet on CIFAR-10 their approach struggles to achieve even the modest test accuracy of 80%. Moreover, they do not consider ensembling.

By contrast, we propose a much simpler training procedure that can find near-constant accuracy polygonal chains with only one bend between optima, even on a range of modern state-of-the-art architectures. Inspired by properties of the loss function discovered by our procedure, we also propose a new state-of-the-art ensembling method that can be trained in the time required to train a single DNN, with compelling performance on many key benchmarks (e.g., 96.4% accuracy on CIFAR-10).

Draxler et al. [5] simultaneously and independently discovered the existence of curves connecting local optima in DNN loss landscapes. To find these curves they used a different approach inspired by the Nudged Elastic Band method [12] from quantum chemistry.

Xie et al. [21] proposed a related ensembling approach that gathers outputs of neural networks from different epochs at the end of training to stabilize final predictions. More recently, Huang et al. [11] proposed snapshot ensembles, which use a cosine cyclical learning rate [17] to save “snapshots” of the model during training at times when the learning rate achieves its minimum. In our experiments, we compare our geometrically inspired approach to Huang et al. [11], showing improved performance.

### 3 Finding Paths between Modes

We describe a new method to minimize the training error along a path that connects two points in the space of DNN weights. Section 3.1 introduces this general procedure for arbitrary parametric curves, and Section 3.2 describes polygonal chains and Bezier curves as two example parametrizations of such curves. In the supplementary material, we discuss the computational complexity of the proposed approach and how to apply batch normalization at test time to points on these curves. We note that after curve finding experiments in Section 4 we make our key methodological proposal for ensembling in Section 5.

#### 3.1 Connection Procedure

Let \( \hat{w}_1 \) and \( \hat{w}_2 \) in \( \mathbb{R}^{\text{net}} \) be two sets of weights corresponding to two neural networks independently trained by minimizing any user-specified loss \( L(w) \), such as the cross-entropy loss. Here, \( \text{net} \) is the number of weights of the DNN. Moreover, let \( \phi_\theta : [0, 1] \rightarrow \mathbb{R}^{\text{net}} \) be a continuous piecewise smooth parametric curve, with parameters \( \theta \), such that \( \phi_\theta(0) = \hat{w}_1 \), \( \phi_\theta(1) = \hat{w}_2 \).
To find a path of high accuracy between \( \hat{w}_1 \) and \( \hat{w}_2 \), we propose to find the parameters \( \theta \) that minimize the expectation over a uniform distribution on the curve, \( \ell(\theta) \):

\[
\hat{\ell}(\theta) = \frac{\int \mathcal{L}(\phi_\theta) d\phi_\theta}{\int d\phi_\theta} = \frac{\int_0^1 \mathcal{L}(\phi_\theta(t)) \|\phi'_\theta(t)\| dt}{\int_0^1 \|\phi'_\theta(t)\| dt} = \int_0^1 \mathcal{L}(\phi_\theta(t)) q_\theta(t) dt = \mathbb{E}_{t \sim q_\theta} \left[ \mathcal{L}(\phi_\theta(t)) \right], \tag{1}
\]

where the distribution \( q_\theta(t) \) on \( t \in [0, 1] \) is defined as: \( q_\theta(t) = \|\phi'_\theta(t)\| \cdot \left( \int_0^1 \|\phi'_\theta(t)\| dt \right)^{-1} \). The numerator of (1) is the line integral of the loss \( \mathcal{L} \) on the curve, and the denominator \( \int_0^1 \|\phi'_\theta(t)\| dt \) is the normalizing constant of the uniform distribution on the curve defined by \( \phi_\theta(\cdot) \). Stochastic gradients of \( \hat{\ell}(\theta) \) in Eq. (1) are generally intractable since \( q_\theta(t) \) depends on \( \theta \). Therefore we also propose a more computationally tractable loss \( \ell(\theta) \):

\[
\ell(\theta) = \int_0^1 \mathcal{L}(\phi_\theta(t)) dt = \mathbb{E}_{t \sim U(0,1)} \mathcal{L}(\phi_\theta(t)), \tag{2}
\]

where \( U(0, 1) \) is the uniform distribution on \([0, 1]\). The difference between (1) and (2) is that the latter is an expectation of the loss \( \mathcal{L}(\phi_\theta(t)) \) with respect to a uniform distribution on \( t \in [0, 1] \), while (1) is an expectation with respect to a uniform distribution on the curve. The two losses coincide, for example, when \( \phi_\theta(\cdot) \) defines a polygonal chain with two line segments of equal length and the parametrization of each of the two segments is linear in \( t \).

To minimize (2), at each iteration we sample \( \hat{t} \) from the uniform distribution \( U(0, 1) \) and make a gradient step for \( \theta \) with respect to the loss \( \mathcal{L}(\phi_\theta(\hat{t})) \). This way we obtain unbiased estimates of the gradients of \( \ell(\theta) \), as

\[
\nabla_{\theta} \mathcal{L}(\phi_\theta(\hat{t})) \approx \mathbb{E}_{t \sim U(0,1)} \nabla_{\theta} \mathcal{L}(\phi_\theta(t)) = \nabla_{\theta} \mathbb{E}_{t \sim U(0,1)} \mathcal{L}(\phi_\theta(t)) = \nabla_{\theta} \ell(\theta).
\]

We repeat these updates until convergence.

### 3.2 Example Parametrizations

**Polygonal chain** The simplest parametric curve we consider is the polygonal chain (see Figure 1 right). The trained networks \( \hat{w}_1 \) and \( \hat{w}_2 \) serve as the endpoints of the chain and the bends of the chain are the parameters \( \theta \) of the curve parametrization. Consider the simplest case of a chain with one bend \( \theta \). Then

\[
\phi_\theta(t) = \begin{cases} 
2(t \theta + (0.5 - t) \hat{w}_1), & 0 \leq t \leq 0.5 \\
2((t - 0.5) \hat{w}_2 + (1 - t) \theta), & 0.5 \leq t \leq 1.
\end{cases}
\]

**Bezier curve** A Bezier curve (see Figure 1 middle) provides a convenient parametrization of smooth paths with given endpoints. A quadratic Bezier curve \( \phi_\theta(t) \) with endpoints \( \hat{w}_1 \) and \( \hat{w}_2 \) is given by

\[
\phi_\theta(t) = (1 - t)^2 \hat{w}_1 + 2t(1 - t)\theta + t^2 \hat{w}_2, \quad 0 \leq t \leq 1.
\]

These formulas naturally generalize for \( n \) bends \( \theta = \{w_1, w_2, \ldots, w_n\} \) (see supplement).

## 4 Curve Finding Experiments

We show that the proposed training procedure in Section 3 does indeed find high accuracy paths connecting different modes, across a range of architectures and datasets. Moreover, we further investigate the properties of these curves, showing that they correspond to meaningfully different representations that can be ensembled for improved accuracy. We use these insights to propose an improved ensembling procedure in Section 5 which we empirically validate in Section 6.

In particular, we test VGG-16 [19], a 28-layer Wide ResNet with widening factor 10 [22] and a 155-layer ResNet [9] on CIFAR-10, and VGG-16, 164-layer ResNet-bottleneck [9] on CIFAR-100. For CIFAR-10 and CIFAR-100 we use the same standard data augmentation as Huang et al. [11]. We
provide additional results, including detailed experiments for fully connected and recurrent networks, in the supplement.

For each model and dataset we train two networks with different random initializations to find two modes. Then we use the proposed algorithm of Section 3 to find a path connecting these two modes in the weight space with a quadratic Bezier curve and a polygonal chain with one bend. We also connect the two modes with a line segment for comparison. In all experiments we optimize the loss (2), as for Bezier curves the gradient of loss (1) is intractable, and for polygonal chains we found loss (2) to be more stable.

Figures 1 and 2 show the results of the proposed mode connecting procedure for ResNet-164 on CIFAR-100. Here loss refers to $\ell_2$-regularized cross-entropy loss. For both the Bezier curve and polygonal chain, train loss (Figure 2, left) and test error (Figure 2, middle) are indeed nearly constant. In addition, we provide plots of train error and test loss in the supplementary material. In the supplement, we also include a comprehensive table summarizing all path finding experiments on CIFAR-10 and CIFAR-100 for VGGs, ResNets and Wide ResNets, as well as fully connected networks and recurrent neural networks, which follow the same general trends. In the supplementary material we also show that the connecting curves can be found consistently as we vary the number of parameters in the network, although the ratio of the arc length for the curves to the length of the line segment connecting the same endpoints decreases with increasing parametrization. In the supplement, we also measure the losses (1) and (2) for all the curves we constructed, and find that the values of the two losses are very close, suggesting that the loss (2) is a good practical approximation to the loss (1).

The constant-error curves connecting two given networks discovered by the proposed method are not unique. We trained two different polygonal chains with the same endpoints and different random seeds using VGG-16 on CIFAR-10. We then measured the Euclidean distance between the turning points of these curves. For VGG-16 on CIFAR-10 this distance is equal to 29.6 and the distance between the endpoints is 50, showing that the curves are not unique. In this instance, we expect the distance between turning points to be less than the distance between endpoints, since the locations of the turning points were initialized to the same value (the center of the line segment connecting the endpoints).

Although high accuracy connecting curves can often be very simple, such as a polygonal chain with only one bend, we note that line segments directly connecting two modes generally incur high error. For VGG-16 on CIFAR-10 the test error goes up to 90% in the center of the segment. For ResNet-158 and Wide ResNet-28-10 the worst errors along direct line segments are still high, but relatively less, at 80% and 66%, respectively. This finding suggests that the loss surfaces of state-of-the-art residual networks are indeed more regular than those of classical models like VGG, in accordance with the observations in Li et al. [16].

In this paper we focus on connecting pairs of networks trained using the same hyper-parameters, but from different random initializations. Building upon our work, Gotmare et al. [8] have recently shown that our mode connectivity approach applies to pairs of networks trained with different batch sizes, optimizers, data augmentation strategies, weight decays and learning rate schemes.

To motivate the ensembling procedure proposed in the next section, we now examine how far we need to move along a connecting curve to find a point that produces substantially different, but still
useful, predictions. Let $\hat{w}_1$ and $\hat{w}_2$ be two distinct sets of weights corresponding to optima obtained by independently training a DNN two times. We have shown that there exists a path connecting $\hat{w}_1$ and $\hat{w}_2$ with high test accuracy. Let $\phi_\theta(t), t \in [0, 1]$ parametrize this path with $\phi_\theta(0) = \hat{w}_1$, $\phi_\theta(1) = \hat{w}_2$. We investigate the performance of an ensemble of two networks: the endpoint $\phi_\theta(0)$ of the curve and a point $\phi_\theta(t)$ on the curve corresponding to $t \in [0, 1]$. Figure 2 (right) shows the test error of this ensemble as a function of $t$, for a ResNet-164 on CIFAR-100. The test error starts decreasing at $t \approx 0.1$ and for $t \geq 0.4$ the error of an ensemble is already as low as the error of an ensemble of the two independently trained networks used as the endpoints of the curve. Thus even by moving away from the endpoint by a relatively small distance along the curve we can find a network that produces meaningfully different predictions from the network at the endpoint. This result also demonstrates that these curves do not exist only due to degenerate parametrizations of the network (such as rescaling on either side of a ReLU); instead, points along the curve correspond to meaningfully different representations of the data that can be ensembled for improved performance. In the supplementary material we show how to create trivially connecting curves that do not have this property.

5 Fast Geometric Ensembling

In this section, we introduce a practical ensembling procedure, Fast Geometric Ensembling (FGE), motivated by our observations about mode connectivity.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure3.png}
\caption{Left: Plot of the learning rate (Top), test error (Middle) and distance from the initial value $\hat{w}$ (Bottom) as a function of iteration for FGE with Preactivation-ResNet-164 on CIFAR-100. Circles indicate the times when we save models for ensembling. Right: Ensemble performance of FGE and SSE (Snapshot Ensembles) as a function of training time, using ResNet-164 on CIFAR-100 ($B = 150$ epochs). Crosses represent the performance of separate “snapshot” models, and diamonds show the performance of the ensembles constructed of all models available by the given time.}
\end{figure}

In the previous section, we considered ensembling along mode connecting curves. Suppose now we instead only have one set of weights $\hat{w}$ corresponding to a mode of the loss. We cannot explicitly construct a path $\phi_\theta(\cdot)$ as before, but we know that multiple paths passing through $\hat{w}$ exist, and it is thus possible to move away from $\hat{w}$ in the weight space without increasing the loss. Further, we know that we can find diverse networks providing meaningfully different predictions by making relatively small steps in the weight space (see Figure 2 right).

Inspired by these observations, we propose the Fast Geometric Ensembling (FGE) method that aims to find diverse networks with relatively small steps in the weight space, without leaving a region that corresponds to low test error. While inspired by mode connectivity, FGE does not rely on explicitly finding a connecting curve, and thus does not require pre-trained endpoints, and so can be trained in the time required to train a single network.

Let us describe Fast Geometric Ensembling. First, we initialize a copy of the network with weights $\hat{w}$ set equal to the weights of the trained network $\hat{w}$. Now, to force $w$ to move away from $\hat{w}$ without substantially decreasing the prediction accuracy we adopt a cyclical learning rate schedule $\alpha(\cdot)$ (see Figure 3 left), with the learning rate at iteration $i = 1, 2, \ldots$ defined as

$$
\alpha(i) = \begin{cases} 
(1 - 2t(i))\alpha_1 + 2t(i)\alpha_2 & 0 < t(i) \leq \frac{1}{2} \\
(2 - 2t(i))\alpha_2 + (2t(i) - 1)\alpha_1 & \frac{1}{2} < t(i) \leq 1
\end{cases}
$$
where \( t(i) = \frac{1}{2}(\text{mod}(i - 1, c) + 1) \), the learning rates are \( \alpha_1 > \alpha_2 \), and the number of iterations in one cycle is given by even number \( c \). Here by iteration we mean processing one mini-batch of data. We can train the network \( w \) using the standard \( \ell_2 \)-regularized cross-entropy loss function (or any other loss that can be used for DNN training) with the proposed learning rate schedule for \( n \) iterations. In the middle of each learning rate cycle when the learning rate reaches its minimum value \( \alpha(i) = \alpha_2 \) (which corresponds to \( \text{mod}(i - 1, c) + 1 = c/2 \), \( t(i) = \frac{c}{2} \)) we collect the checkpoints of weights \( w \). When the training is finished we ensemble the collected models. An outline of the algorithm is provided in the supplement.

Figure 4 (left) illustrates the adopted learning rate schedule. During the periods when the learning rate is large (close to \( \alpha_1 \)), \( w \) is exploring the weight space doing larger steps but sacrificing the test error. When the learning rate is small (close to \( \alpha_2 \)), \( w \) is in the exploitation phase in which the steps become smaller and the test error goes down. The cycle length is usually about 2 to 4 epochs, so that the method efficiently balances exploration and exploitation with relatively-small steps in the weight space that are still sufficient to gather diverse and meaningful networks for the ensemble.

To find a good initialization \( \hat{w} \) for the proposed procedure, we first train the network with the standard learning rate schedule (the schedule used to train single DNN models) for about 80% of the time required to train a single model. After this pre-training is finished we initialize FGE with \( \hat{w} \) and run the proposed fast ensembling algorithm for the remaining computational budget. In order to get more diverse samples, one can run the algorithm described above several times for a smaller number of iterations initializing from different checkpoints saved during training of \( \hat{w} \), and then ensemble all of the models gathered across these runs.

Cyclical learning rates have also recently been considered in Smith and Topin [20] and Huang et al. [11]. Our proposed method is perhaps most closely related to Snapshot Ensembles [11], but has several distinctive features, inspired by our geometric insights. In particular, Snapshot Ensembles adopt cyclical learning rates with cycle length on the scale of 20 to 40 epochs from the beginning of the training as they are trying to do large steps in the weight space. However, according to our analysis of the curves it is sufficient to do relatively small steps in the weight space to get diverse networks, so we only employ cyclical learning rates with a small cycle length on the scale of 2 to 4 epochs in the last stage of the training. As illustrated in Figure 5 (left), the step sizes made by FGE between saving two models (that is the euclidean distance between sets of weights of corresponding models in the weight space) are on the scale of 7 for Preactivation-ResNet-164 on CIFAR-100. For Snapshot Ensembles for the same model the distance between two snapshots is on the scale of 40. We also use a piecewise linear cyclical learning rate schedule following Smith and Topin [20] as opposed to the cosine schedule in Snapshot Ensembles.

### 6 Fast Geometric Ensembling Experiments

<table>
<thead>
<tr>
<th>DNN (Budget)</th>
<th>method</th>
<th>CIFAR-100</th>
<th>CIFAR-10</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>1B</td>
<td>2B</td>
</tr>
<tr>
<td>VGG-16 (200)</td>
<td>Ind</td>
<td>27.4 ± 0.1 25.28 24.45</td>
<td>6.75 ± 0.16 5.89 5.9</td>
</tr>
<tr>
<td></td>
<td>SSE</td>
<td>26.4 ± 0.1 25.16 24.69</td>
<td>6.57 ± 0.12 6.19 5.95</td>
</tr>
<tr>
<td></td>
<td>FGE</td>
<td>25.7 ± 0.1 24.11 23.54</td>
<td><strong>6.48 ± 0.09</strong> 5.82 5.66</td>
</tr>
<tr>
<td>ResNet-164 (150)</td>
<td>Ind</td>
<td>21.5 ± 0.4 19.04 18.59</td>
<td>4.72 ± 0.1 4.1 <strong>3.77</strong></td>
</tr>
<tr>
<td></td>
<td>SSE</td>
<td>20.9 ± 0.2 19.28 18.91</td>
<td>4.66 ± 0.02 4.37 4.3</td>
</tr>
<tr>
<td></td>
<td>FGE</td>
<td><strong>20.2 ± 0.1</strong> 18.67 18.21</td>
<td><strong>4.54 ± 0.05</strong> 4.21 3.98</td>
</tr>
<tr>
<td>WRN-28-10 (200)</td>
<td>Ind</td>
<td>19.2 ± 0.2 17.48 17.01</td>
<td>3.82 ± 0.1 3.4 <strong>3.31</strong></td>
</tr>
<tr>
<td></td>
<td>SSE</td>
<td>17.9 ± 0.2 17.3 16.97</td>
<td>3.73 ± 0.04 3.54 3.55</td>
</tr>
<tr>
<td></td>
<td>FGE</td>
<td><strong>17.7 ± 0.2</strong> 16.95 16.88</td>
<td><strong>3.65 ± 0.1</strong> 3.38 3.52</td>
</tr>
</tbody>
</table>

In this section we compare the proposed Fast Geometric Ensembling (FGE) technique against ensembles of independently trained networks (Ind), and SnapShot Ensembles (SSE) [11], a recent state-of-the-art fast ensembling approach.
For the ensembling experiments we use a 164-layer Preactivation-ResNet in addition to the VGG-16 and Wide ResNet-28-10 models. Links for implementations to these models can be found in the supplement.

We compare the accuracy of each method as a function of computational budget. For each network architecture and dataset we denote the number of epochs required to train a single model as $B$. For a $kB$ budget, we run each of Ind, FGE and SSE $k$ times from random initializations and ensemble the models gathered from the $k$ runs. In our experiments we set $B = 200$ for VGG-16 and Wide ResNet-28-10 (WRN-28-10) models, and $B = 150$ for ResNet-164, since 150 epochs is typically sufficient to train this model. We note the runtime per epoch for FGE, SSE, and Ind is the same, and so the total computation associated with $kB$ budgets is the same for all ensembling approaches.

For Ind, we use an initial learning rate of $0.1$ for ResNet and Wide ResNet, and $0.05$ for VGG. For FGE, with VGG we use cycle length $c = 2$ epochs, and a total of $22$ models in the final ensemble. With ResNet and Wide ResNet we use $c = 4$ epochs, and the total number of models in the final ensemble is 12 for Wide ResNets and 6 for ResNets. For VGG we set the learning rates to $\alpha_1 = 10^{-2}$, $\alpha_2 = 5 \cdot 10^{-4}$; for ResNet and Wide ResNet models we set $\alpha_1 = 5 \cdot 10^{-2}$, $\alpha_2 = 5 \cdot 10^{-4}$. For SSE, we followed Huang et al. [11] and varied the initial learning rate $\alpha_0$ and number of snapshots per run $M$. We report the best results we achieved, which corresponded to $\alpha_0 = 0.1, M = 4$ for ResNet, $\alpha_0 = 0.1, M = 5$ for Wide ResNet, and $\alpha_0 = 0.05, M = 5$ for VGG. The total number of models in the FGE ensemble is constrained by network choice and computational budget. Further experimental details are in the supplement.

Table 1 summarizes the results of the experiments. In all conducted experiments FGE outperforms SSE, particularly as we increase the computational budget. The performance improvement against Ind is most noticeable for CIFAR-100. With a large number of classes, any two models are less likely to make the same predictions. Moreover, there will be greater uncertainty over which representation one should use on CIFAR-100, since the number of classes is increased tenfold from CIFAR-10, but the number of training examples is held constant. Thus smart ensembling strategies will be especially important on this dataset. Indeed in all experiments on CIFAR-100, FGE outperformed all other methods. On CIFAR-10, FGE consistently improved upon SSE for all budgets and architectures. FGE also improved against Ind for all training budgets with VGG, but is more similar in performance to Ind on CIFAR-10 when using ResNets.

Figure 3 (right) illustrates the results for Preactivation-ResNet-164 on CIFAR-100 for one and two training budgets. The training budget $B$ is 150 epochs. Snapshot Ensembles use a cyclical learning rate from the beginning of the training and they gather the models for the ensemble throughout training. To find a good initialization we run standard independent training for the first 125 epochs before applying FGE. In this case, the whole ensemble is gathered over the following 22 epochs (126-147) to fit in the budget of each of the two runs. During these 22 epochs FGE is able to gather diverse enough networks to outperform Snapshot Ensembles both for $1B$ and $2B$ budgets.

Diversity of predictions of the individual networks is crucial for the ensembling performance [e.g., 15]. We note that the diversity of the networks averaged by FGE is lower than that of completely independently trained networks. Specifically, two independently trained ResNet-164 on CIFAR-100 make different predictions on 19.97% of test objects, while two networks from the same FGE run make different predictions on 14.57% of test objects. Further, performance of individual networks averaged by FGE is slightly lower than that of fully trained networks (e.g. 78.0% against 78.5% on CIFAR100 for ResNet-164). However, for a given computational budget FGE can propose many more high-performing networks than independent training, leading to better ensembling performance (see Table 1).

6.1 ImageNet

ImageNet ILSVRC-2012 [18] is a large-scale dataset containing 1.2 million training images and 50000 validation images divided into 1000 classes.

CIFAR-100 is the primary focus of our ensemble experiments. However, we also include ImageNet results for the proposed FGE procedure, using a ResNet-50 architecture. We used a pretrained model with top-1 test error of 23.87 to initialize the FGE procedure. We then ran FGE for 5 epochs with a cycle length of 2 epochs and with learning rates $\alpha_1 = 10^{-3}, \alpha_2 = 10^{-5}$. The top-1 test error-rate of the final ensemble was 23.31. Thus, in just 5 epochs we could improve the accuracy of the model by
0.56 using FGE. The final ensemble contains 4 models (including the pretrained one). Despite the harder setting of only 5 epochs to construct an ensemble, FGE performs comparably to the best result reported by Huang et al. [11] on ImageNet, 23.33 error, which was also achieved using a ResNet-50.

7 Discussion and Future Work

We have shown that the optima of deep neural networks are connected by simple pathways, such as a polygonal chain with a single bend, with near constant accuracy. We introduced a training procedure to find these pathways, with a user-specific curve of choice. We were inspired by these insights to propose a practical new ensembling approach, Fast Geometric Ensembling, which achieves state-of-the-art results on CIFAR-10, CIFAR-100, and ImageNet.

There are so many exciting future directions for this research. At a high level we have shown that even though the loss surfaces of deep neural networks are very complex, there is relatively simple structure connecting different optima. Indeed, we can now move towards thinking about valleys of low loss, rather than isolated modes.

These valleys could inspire new directions for approximate Bayesian inference, such as stochastic MCMC approaches which could now jump along these bridges between modes, rather than getting stuck exploring a single mode. One could similarly derive new proposal distributions for variational inference, exploiting the flatness of these pathways. These geometric insights could also be used to accelerate the convergence, stability and accuracy of optimization procedures like SGD, by helping us understand the trajectories along which the optimizer moves, and making it possible to develop procedures which can now search in more structured spaces of high accuracy. One could also use these paths to construct methods which are more robust to adversarial attacks, by using an arbitrary collection of diverse models described by a high accuracy curve, returning the predictions of a different model for each query from an adversary. We can also use this new property to create better visualizations of DNN loss surfaces. Indeed, using the proposed training procedure, we were able to produce new types of visualizations showing the connectivity of modes, which are normally depicted as isolated. We also could continue to build on the new training procedure we proposed here, to find curves with particularly desirable properties, such as diversity of networks. Indeed, we could start to use entirely new loss functions, such as line and surface integrals of cross-entropy across structured regions of weight space.

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