Abstract

Fine-tuning from a collection of models pre-trained on different domains (a “model zoo”) is emerging as a technique to improve test accuracy in the low-data regime. However, model selection, i.e. how to pre-select the right model to fine-tune from a model zoo without performing any training, remains an open topic. We use a linearized framework to approximate fine-tuning, and introduce two new baselines for model selection – Label-Gradient and Label-Feature Correlation. Since all model selection algorithms in the literature have been tested on different use-cases and never compared directly, we introduce a new comprehensive benchmark for model selection comprising of:

i) A model zoo of single and multi-domain models, and

ii) Many target tasks. Our benchmark highlights accuracy gain with model zoo compared to fine-tuning Imagenet models. We show our model selection baseline can select optimal models to fine-tune in few selections and has the highest ranking correlation to fine-tuning accuracy compared to existing algorithms.

1. Introduction

A “model zoo” is a collection of pre-trained models, obtained by training different architectures on many datasets covering a variety of tasks and domains. For instance, the zoo could comprise models (or experts) trained to classify, say, trees, birds, fashion items, aerial images, etc. The typical use of a model zoo is to provide a good initialization which can then be fine-tuned for a new target task, for which we have few training data. This strategy is an alternative to the more common practice of starting from a model trained on a large dataset, say Imagenet [13], and is aimed at providing better domain coverage and a stronger inductive bias. Despite the growing usage of model zoos [10, 26, 31, 48] there is little in the way of analysis, both theoretical and empirical, to illuminate which approach is preferable under what conditions. In Fig. 1, we show that fine-tuning with a model zoo is indeed better, especially when training data is limited. Fig. 1 also shows that using a model zoo, we can outperform hyper-parameter optimization performed during fine-tuning of the Imagenet pre-trained model.

Fine-tuning with a model zoo can be done by brute-force fine-tuning of each model in the zoo, or more efficiently by using “model selection” to select the closest model (or best initialization) from which to fine-tune. The goal of model selection therefore is to find the best pre-trained model to fine-tune on the target task, without performing the actual fine-tuning. So, we seek an approximation to the fine-tuning process. In our work, we develop an analytical framework to characterize the fine-tuning process using a linearization of the model around the point of pre-training [35], drawing inspiration from the work on the Neural Tangent Kernel (NTK) [24, 29]. Our analysis of generalization bounds and training speed using linearized fine-tuning naturally suggests two criterion to select the best model to fine-tune from, which we call Label-Gradient Correlation (LGC) and Label-Feature Correlation (LFC). Given its simplicity, we consider our criteria as baselines, rather than full-fledged methods for model selection, and compare the state-of-the-art in model selection – e.g. RSA [15], LEEP [37], Domain Similarity [10], Feature Metrics [49] – against it.

Model selection being a relatively recent endeavor, there is currently no standard dataset or a common benchmark to perform such a comparison. For example, LEEP [37] performs its model selection experiments on transfer (or fine-tuning) from Imagenet pre-trained model to 200 randomly sampled tasks of CIFAR-100 [28] image classification, RSA [15] uses the Taskonomy dataset [55] to evaluate its prediction of task transfer (or model selection) performance. Due to these different experimental setups, the state-of-the-art in model selection is unclear. Therefore, in Sec. 4 we build a new benchmark comprising a large model zoo and many target tasks. For our model zoo, we use 8 large image classification datasets (from different domains) to train single-domain and multi-domain experts. We use various image classification datasets as target tasks and study fine-tuning (Sec. 4.2) and model selection (Sec. 4.3) using our model zoo. To the best of our knowledge ours is
show that, for tuning in deep learning is still debated. He et al. [20] trained model can be matched by simply training a net-

2. Related work

Fine-tuning. The exact role of pre-training and fine-
tuning in deep learning is still debated. He et al. [20] show that, for object detection, the accuracy of a pre-
trained model can be matched by simply training a net-

(a) Model zoo vs. different architectures. Fine-tuning using our model zoo is better (i.e. lower test error) than fine-tuning using different architectures with Random or Imagenet pre-trained initialization. We use fine-tuning hyper-parameters of Sec. 4.2 with \( \eta = .005 \).

(b) Model zoo vs. HPO. of Imagenet expert. Fine-tuning using our model zoo is better than fine-tuning with hyper-parameter optimization (HPO) of Imagenet pre-trained Resnet-101 model. We use fine-tuning hyper-parameters of Sec. 4.2 and perform HPO with \( \eta = .01, .005, 0.001 \).

Figure 1. Fine-tuning using our model zoo can obtain lower test error compared to: (a) using different architectures and (b) hyper-parameter optimization (HPO) of Imagenet expert. The standard fine-tuning approach entails picking a network architecture pre-trained on Imagenet to fine-tune and performing hyper-parameter optimization (HPO) during fine-tuning. We outperform this strategy by fine-tuning using our model zoo described in Sec. 4.1. We plot test error as a function of the number of per-class samples (i.e. shots) in the dataset. In (a), we compare fine-tuning with our single-domain experts in the model zoo to using different architectures (AlexNet, ResNet-18, ResNet-101, Wide ResNet-101) for fine-tuning. In (b), we show fine-tuning with our model zoo obtains lower error than performing HPO on Imagenet pre-trained Resnet-101 [19] during fine-tuning. Model zoo lowers the test error, especially in the low-data regime (5, 10, 20-shot per class samples of target task). Since we compare to Imagenet fine-tuning, we exclude Imagenet experts from our model zoo for the above plots.

the first large-scale benchmark for model selection.

By performing fine-tuning and model selection on our benchmark, we discover the following:

(a) We show (Fig. 1) that fine-tuning models in the model zoo can outperform the standard method of fine-tuning with Imagenet pre-trained architectures and HPO. We obtain better fine-tuning than Imagenet expert with, both model zoo of single-domain experts (Fig. 2) and multi-domain experts (Fig. 3). While in the high-data regime using a model zoo leads to modest gains, it sensibly improves accuracy in the low-data regime.

(b) For any given target task, we show that only a small subset of the models in the zoo lead to accuracy gain (Fig. 2). In such a scenario, brute-force fine-tuning all models to find the few that improve accuracy is wasteful. Fine-tuning with all our single-domain experts in the model zoo is 40× more compute intensive than fine-tuning an Imagenet Resnet-101 expert in Tab. 3.

(c) Our LGC model selection, and particularly its approximation LFC, can find the best models from which to fine-tune without requiring an expensive brute-force search (Tab. 3). With only 3 selections, we can select models that show gain over Imagenet expert (Fig. 4). Compared to Domain Similarity [11], RSA [15] and Feature Metrics [49], our LFC score can select the best model to fine-tune in fewer selections, and it shows the highest ranking correlation to the fine-tuning test accuracy (Fig. 6) among all model selection methods.

2. Related work

Model Selection. Empirical evidence [1, 31, 54] and theory [2] suggests that effectiveness of fine-tuning relates to a notion of distance between tasks. Taskonomy [54] defines a distance between learning tasks a-posteriori, that is, by looking at the fine-tuning accuracy during transfer learning. However, for predicting the best pre-training without performing fine-tuning, an a-priori approach is best. Achille et al. [1, 2] introduce a fixed-dimensional “task embedding” to encode distance between tasks. Cui et al. [11] propose a Domain Similarity measure, which entails using the Earth Mover Distance (EMD) between source and target features. LEEP [37, 46] looks at the conditional cross-entropy between the output of the pre-trained model and the target labels. RSA [15] compares representation dissimilarity matrices of features from pre-trained model and a small network trained on target task for model selection. As op-
posed to using the ad-hoc measure of task similarity, we rely on a linearization approximation to the fine-tuning to derive our model selection methods (Sec. 3).

**Linearization and NTK.** To analyse fine-tuning from pre-trained weights, we use a simple but effective framework inspired by the Neural Tangent Kernel (NTK) formalism [24]: We approximate the fine-tuning dynamics by looking at a linearization of the source model around the pre-trained weights \( w_0 \) (Sec. 3.1). This approximation has been suggested by [35], who also notes that while there may be doubts on whether an NTK-like approximation holds for real randomly-initialized network [16], it is more likely to hold in the case of fine-tuning, since the fine-tuned weights tend to remain close to the pre-trained weights.

**Few-shot.** Interestingly, while pre-training has a higher impact in the few-shot regime, there is only a handful of papers that experiment with it [14, 18, 47]. This could be due to over-fitting of the current literature on standard benchmarks that have a restricted scope. We hope that our proposed benchmark (Sec. 4) may foster further research.

### 3. Approach

**Notation.** We have a model zoo, \( \mathcal{F} \), of \( n \) pre-trained models or experts: \( \mathcal{F} = \{ f^1, f^2, \ldots, f^n \} \). Our aim is to classify a target dataset, \( D = \{(x_i, y_i)\}_{i=1}^N \), by fine-tuning models in the model zoo. Here, \( x_i \in \mathcal{X} \), is the \( i \)th input image and \( y_i \in \mathcal{Y} \), is the corresponding class label. For a network \( f \in \mathcal{F} \) with weights \( w \), we denote the output of the network with \( f_w(x) \). \( w_0 \) denotes the initialization (or pre-trained weights) of models in the model zoo. The goal of model selection is to predict a score \( S(f_{w_0}, D) \) that measures the fine-tuning accuracy on the test set \( D_{\text{test}} \), when \( D \) is used to fine-tune the model \( f_{w_0} \). Note, \( S \) does not have to exactly measure the fine-tuning accuracy, it needs to only predict a score that correlates to the ranking by fine-tuning accuracy. The model selection score for every pre-trained model, \( S(f^k, D) \) for \( k \in \{1, 2, \ldots, n\} \), can then be used as proxy to rank and select top-k models by their fine-tuning accuracy. Since the score \( S \) needs to estimate (a proxy for) fine-tuning accuracy without performing any fine-tuning, in Sec. 3.1 we construct a linearization approximation to fine-tuning and present several results that allow us to derive our Label-Gradient Correlation \( (S_{LG}) \) and Label-Feature Correlation \( (S_{LF}) \) (Sec. 3.2) scores for model selection from it. In Fig. 6 (b), we show our scores have higher ranking correlation to fine-tuning accuracy than existing work.

#### 3.1. Linearized framework to analyse fine-tuning

Given an initialization \( w_0 \), the weights of the pre-trained model, we can define the linearized model:

\[
f_w^{\text{lin}}(x) := f_{w_0}(x) + \nabla_w f_{w_0}(x)|_{w=w_0}(w - w_0),
\]

which approximates the output of the real model for \( w \) close to \( w_0 \). Mu et al. [35] observe that, while in general not accurate, a linear approximation can correctly describe the model throughout fine-tuning since the weights \( w \) tend to remain close to the initial value \( w_0 \). Under this linear approximation [29] shows the following proposition.

**Proposition 1** Let \( D = \{(x_i, y_i)\}_{i=1}^N \) be the target dataset. Assume the task is a binary classification problem with labels \( y_i = \pm 1 \), using the \( L_2 \) loss \( L_D(w) = \sum_{i=1}^N (y_i - f_w(x_i))^2 \). Let \( w_t \) denote the weights at time \( t \) during training. Then, the loss function evolve as:

\[
L_t = (Y - f_{w_0}(X))^T e^{-2y\Theta y'} (Y - f_{w_0}(X))
\]

where \( f_{w_0}(X) \) denotes the vector containing the output of the network on all the images in the dataset, \( Y \) denotes the vectors of all training labels, and we defined the Neural Tangent Kernel (NTK) matrix:

\[
\Theta := \nabla_w f_{w_0}(X)\nabla_w f_{w_0}(X)'^T
\]

which is the \( N \times N \) Gram matrix of all the per-sample gradients.

From Prop. 1, the behavior of the network during fine-tuning is fully characterized by the kernel matrix \( \Theta \), which depends on the pre-trained model \( f_{w_0} \), the data \( X \) and the task labels \( Y \). We then expect to be able to select the best model by looking at these quantities. To show how we can do this, we now derive several results connecting \( \Theta \) and \( Y \) to the quantities of relevance for model selection below, i.e. Training time and Generalization on the target task.

**Training time.** In [56], it is shown that the loss \( L_t \) of the linearized model evolves with training over time \( t \) as

\[
L_t = \|\delta Y\|^2 - 2t\delta Y' \Theta \delta Y' + O(t^2).
\]

where we have defined \( \delta Y^t = Y - f_{w_0}(X) \) to be the initial projection residual. Eq. (3) suggests using the quadratic term \( \delta Y' \Theta \delta Y \) as a simple estimate of the training speed.

**Generalization.** The most important criterion for model selection is generalization performance. Unfortunately, we cannot have any close form characterization of generalization error, which depends on test data we do not have. However, in [3] the following bound on the test error is suggested:

\[
L_{\text{test}}^2 \leq \frac{1}{n} \lambda^T \Theta^{-1} \lambda = \frac{1}{n} \sum_k \frac{1}{\lambda_k} (Y \cdot v_k)^2.
\]

We see that if \( Y \) correlates more with the first principal components of variability of the per-sample gradients (so that \( Y \cdot v_k \) is larger), then we expect better generalization.

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1This is to simplify the notation, but a similar result would hold for a multi-class classification using one-hot encoding. Using the \( L_2 \) loss is necessary to have a close form expression. However, note that empirically the \( L_2 \) performs similarly to cross-entropy during fine-tuning [17, 4].
Arora et al. [3] prove that this bound holds with high-probability for a wide-enough randomly initialized 3-layer network. In practice, however, this generalization bound may be vacuous as hypotheses are not satisfied (the network is deeper, and the initialization is not Gaussian). For this reason, rather than using the above quantity as a real bound, we refer to it as an empirical “generalization score”.

Note eq. (3) and eq. (4) contain the similar terms $\delta Y^T \Theta_1 \delta Y$ and $\delta Y^T \Theta^{-1} \delta Y$. By diagonalizing $\Theta$ and applying Jensen’s inequality we have the following relation between the two:

$$\left(\frac{\delta Y^T \Theta_1 \delta Y}{\|\delta Y\|^2}\right)^{-1} \leq \frac{\delta Y^T \Theta^{-1} \delta Y}{\|\delta Y\|^2}. \tag{5}$$

Hence, good “generalization score” $\delta Y^T \Theta^{-1} \delta Y$ implies faster initial fine-tuning, that is, larger $\delta Y^T \Theta \delta Y$. In general we expect the two quantities to be correlated. Hence, selecting the fastest model to train or the one that generalizes better are correlated objectives. $\delta Y^T \Theta \delta Y$ is an approximation to $\delta Y^T \Theta_1 \delta Y$ that uses task labels $Y$ and kernel $\Theta$, and we use it to derive our model selection scores in Sec. 3.2. Large value of $\delta Y^T \Theta \delta Y$ implies better generalization and faster training and it is desirable for a model when fine-tuning.

**Should model selection use gradients or features?**

Our analysis is in terms of the matrix $\Theta$ which depends on the network’s gradients (2), not on its features. In Sec. A, we show that it suffices to use features (i.e., network activations) in (2) as an approximation to the NTK matrix. Let $\Theta(x)|_t$ denote the feature vector (or activation) extracted from layer $l$ of pre-trained network $f$ after forward pass on image, i.e. after $f(x)$. In analogy with the gradient similarity matrix $\Theta$ of (2), we define the feature similarity matrix $\Theta_F$ (which approximates $\Theta$) as follows

$$\Theta_F := [f_w(x)|_l|_t [f_w(x)|_l|_t]^T. \tag{6}$$

**3.2. Label-Feature and Label-Gradient correlation**

We now introduce our two scores for model selection, *Label-Gradient correlation* and *Label-Feature correlation*.

**Label-Gradient Correlation.** From Sec. 3.1 we know that the following score,

$$S_{LG}(f_{w_0}, D) = Y^T \Theta Y = \Theta \cdot Y^T \tag{7}$$

which we call *Label-Gradient Correlation* (LGC), can be used to estimate both the convergence time (eq. 3) and the generalization ability of a model. Here, “:” denotes the dot-product of the matrices (i.e. the sum of Hadamard product of two matrices). $Y^T \Theta$ is an $N \times N$ matrix such that $(Y^T \Theta)^{i,j} = 1$ if $x_i$ and $x_j$ have the same label and $-1$ otherwise. For this reason, we call $Y^T \Theta$ the label similarity matrix. On the other hand, $\Theta_{ij} = \nabla_x f_{w_0}(x_i) \cdot \nabla_x f_{w_0}(x_j)$ is the pair-wise similarity matrix of the gradients. Hence, eq. (7) can be interpreted as giving high LG score (i.e., the model is good for the task) if the gradients are similar whenever the labels are also similar, and are different otherwise.

**Label-Feature Correlation.** Instead of $\Theta$, we can use the approximation $\Theta_F$ from (6) and define our *Label-Feature Correlation* (LFC) score as:

$$S_{LF} = Y^T \Theta_F Y = \Theta_F \cdot Y^T Y. \tag{8}$$

Similarly to the LGC score, this score is higher if samples with the same labels have similar features extracted from the pre-trained network.

**3.3. Implementation**

Notice that the scores $S_{LG}$ and $S_{LF}$ are not normalized. Different pre-training could lead to very different scores if the gradients or the features have a different norm. Also, $\gamma^T \gamma$ used in our scores is specific to binary classification. In practice, we address this as follows: For a multi-class classification problem, let $K_N$ be the $N \times N$-matrix with $(K_N)_{i,j} = 1$ if $x_i$ and $x_j$ have the same label, and $-1$ otherwise. Let $\mu_K$ denote the mean of the entries of $K_N$, and $\mu_\Theta$ the mean of $\Theta$. We define the normalized LGC score as:

$$S_{LG} = \frac{(\Theta - \mu_\Theta) \cdot (K_N - \mu_K)}{\|\Theta - \mu_\Theta\|_2 \|K_N - \mu_K\|_2 \tag{8}.$$}

We normalize LFC similar to LGC in (8). This can also be interpreted as the Pearson’s Correlation coefficient between the entries of $\Theta$ (or $\Theta_F$) and the entries of $K_N$, justifying the name label-gradient (or label-feature) correlation.

**Which features and gradients to use?** For LFC, we extract features from the layer before the fully-connected classification layer (for both Resnet-101 [19] and DenseNet-169 [22] models in our model zoo of Sec. 4.1). We use these features to construct our $\Theta_F$ and compute the normalized LFC. For LGC, following [35], we use gradients corresponding to the last convolutional layer in the pre-trained network. For a large gradient vector, to perform fast computation of LGC, we take a random projection to $10K$ dimensions and compute the normalized LGC score. This results in a trade-off between accuracy and computation for LGC.

**Sampling of target task.** Model selection is supposed to be an inexpensive pre-processing step before actual fine-tuning. To reduce its computation, following previous work of RSA [15], we sample the training set of target dataset $D$ and pick at most 25 images per class to compute our model selection scores. Note, test set is hidden from model selection. Our results show, this still allows us to select models that obtain accuracy gain over Imagenet expert (Fig. 4), and we need few selections (< 7 for model zoo size 30) to select the optimal models (Fig. 6) to fine-tune. We include additional implementation details of our model selection methods and other baselines: RSA [15], Domain Similarity [11], LEEP [37], Feature Metrics [49] in Sec. C.
Table 1. Model zoo of single-domain experts. We train 30 models, Resnet-101 and Densenet-169, on 8 source datasets and measure the top-1 test accuracy. We train our models starting with (√) and without (×) Imagenet pre-training. For all datasets we have higher test accuracy with Resnet-101 (√) than what is reported in the literature (last row), except for iNaturalist [21] by -1.03%. We order datasets from left to right by increasing dataset size. Nwpu-resisc45 [7] has 25K training images while Places-365 [57] has 1.8M. We chose datasets that are publicly available and cover different domains.

Table 2. Multi-domain expert. The top-1 test accuracy of multi-domain model – Multi-BN, Adapter – is comparable to single domain expert for small datasets (Nwpu-Resisc45, Food-101, Logo 2k), while the accuracy is lower on other large datasets. Multi-BN performs better than Shared, Adapter on all datasets and we use this as our multi-domain expert for fine-tuning and model selection.

4. Experiments

Having established the problem of model selection for fine-tuning (Sec. 3), we now put our techniques to test. Sec. 4.1 describes our construction of model zoos with single-domain and multi-domain experts. In Sec. 4.2, we then verify the advantage of fine-tuning using our model zoo with various target tasks. In Sec. 4.3, we compare our LFC, LGC model selection (Sec. 3.2) to previous work, and show that our method can select the optimal models to fine-tune from our model zoo (without performing the actual fine-tuning).

4.1. Model Zoo

We evaluate model selection and fine-tuning with both, a model zoo of single-domain experts (i.e. models trained on single dataset) and a model zoo of multi-domain experts described below.

Source Datasets. Tab. 1 and Tab. 4 lists the source datasets, i.e. the datasets used for training our model zoo. We include publicly available large source datasets (from 25K to 1.8M training images) from different domains, e.g. Nwpu-resisc45 [7] consists of aerial imagery, Food-101 [6] and iNaturalist 2019 [21] consist of food, plant images, Places-365 [57] and Google Landmark v2 [39] contain scene images. This allows us to maximize the coverage of our model zoo to different domains and enables more effective transfer when fine-tuning on different target tasks.

Model zoo of single-domain experts. We build a model zoo of a total of 30 models (Resnet-101 [19] and Densenet-169 [22]) trained on 8 large image classification datasets (i.e. source datasets). Since each model is trained on a single classification dataset (i.e. domain), we refer to these models as single-domain experts. This results in a model zoo, \( \mathcal{F} = \{f_k^i\}_{k=1}^{30} \), to evaluate our model selection.

On each source dataset of Tab. 1, we train Resnet-101 and Densenet-169 models for 90 epochs, with the following hyper-parameters: initial learning rate of 0.1, with decay by 0.1× every 30 epochs, SGD with momentum of 0.9, weight decay of 10^{-4} and a batch size 512. We use the training script\(^2\) from PyTorch [42] library and ensure that our models are well-trained.

In Tab. 1, we show slightly higher top-1 test accuracy for our models trained on ImageNet [13] when compared to the PyTorch [42] model zoo\(^3\). Our Resnet-101 model trained on ImageNet has +.17% top-1 test accuracy and our Densenet-169 model has +.4% top-1 test accuracy vs. PyTorch. On source datasets other than ImageNet, we train our models with (√) and without (×) Imagenet pre-training. This allows us to study the effect of pre-training on a larger dataset when we fine-tune and perform model selection. Note that our Resnet-101 models with (√) ImageNet pre-training have higher accuracy compared to that reported in the literature for all source datasets, except iNaturalist [21] by -1.03%.

Model zoo of multi-domain expert. We also train a Resnet-101 based multi-dataset (or multi-domain) [45] model on the combination of all the 8 source datasets. Our multi-domain Resnet-101 expert, \( f_{w_s,\{w_d\}_D=1} \), uses shared weights (or layers), i.e. \( w_s \), across different domains (or datasets), and in addition it has some domain-specific parameters, i.e. \( \{w_d\}_{D=1} \), for each domain. We have 8 source datasets or domains, so \( D = 8 \) in our benchmark. Note, for fine-tuning we can choose any one of the \( D \) domain-specific parameters to fine-tune. For a given multi-domain expert, this results in a model zoo of \( D \) models (one per domain) that we can fine-tune, \( \mathcal{F} = \{f_{w_s,\{w_d\}_D=1}, f_{w_1}, f_{w_2}, \ldots, f_{w_D}\} \).

We experiment with a few different variants of domain-specific parameters – i) Shared: The domain-specific parameters are also shared, therefore we simply train a Resnet-101 on all datasets, ii) Multi-BN: We replace each batch norm in Resnet-101 architecture with a domain-specific batch norm. Note, for a batch norm layer we replace

\(^2\)https://bit.ly/38NMvyu
\(^3\)https://bit.ly/35V2pFE
**Fine-tuning with model zoo of single-domain experts.** We plot top-1 test error (vertical axis) for fine-tuning with different single domain models in our model zoo. For every target task (on horizontal axis), we have 4 columns of markers from left to right: 1) Imagenet experts in red, 2) Densenet-169 experts with pre-train (✓) and without pre-train (×), 3) Resnet-101 experts with pre-train (✓) and without pre-train (×), 4) We use “black ←” to highlight models that perform better than imagenet expert (i.e. lower error than first column of Imagenet expert per task). Our observations are the following: i) For full target task, we observe better accuracy than Imagenet expert for Magnetic Tile Defects, UC Merced Land Use and iCassava (see black ←). For 20 and 5-shot per class sampling of target task, with the model zoo we outperform Imagenet expert on more datasets, see Oxford Flowers 102, European Flood Depth, Belga Logos and Cub200. Our empirical result, on the importance of different pre-trainings of our model zoo experts when training data is limited, adds to the growing body of similar results in existing literature [20, 31, 58], and ii) The accuracy gain over Imagenet expert is only obtained for fine-tuning with select few models for a given target task, e.g. only one expert for UC Merced Land Use target task in Full, 20-shot setting above. Therefore, brute-force fine-tuning with model zoo leads to wasteful computation. Model selection (Sec. 3) picks the best models to fine-tune and avoids brute-force fine-tuning. Figure is best viewed in high-resolution.

4.2. Fine-tuning on Target Tasks

**Target Tasks.** We use various target tasks (Tab. 4) to study transfer learning from our model zoo of Sec. 4.1: Cucumber [12], Describable Textures [9], Magnetic Tile Defects [23], iCassava [36], Oxford Flowers 102 [38], Oxford-IIIT Pets [41], European Flood Depth [5], UC Merced Land Use [53]. For few-shot, due to lesser compute needed, we use additional target tasks: CUB-200 [52], Stanford Cars [27] and Belga Logos [25]. Note, while some target tasks have domain overlap with our source datasets, e.g. aerial images of UC Merced Land Use [53], other tasks do not have this overlap, e.g. defect images in Magnetic Tile Defects [23], texture images in Describable Textures [9].

**Fine-tuning with single-domain experts in model zoo.** For fine-tuning, Imagenet pre-training is a standard technique. Note, most deep learning frameworks, e.g. PyTorch3, MxNet/Gluon4 etc., just have the Imagenet pre-trained models for different architectures in their model zoo. Fig. 2 shows the top-1 test error obtained by fine-tuning single-domain experts in our model zoo vs. Imagenet expert.

Our fine-tuning hyper-parameters are: 30 epochs, weight decay of 10^{-4}, SGD with Nesterov momentum 0.9, batch size of 32 and learning rate decay by 0.1× at 15 and 25 epochs. We observe that the most important hyper-parameter for test accuracy is the initial learning rate η, so for each fine-tuning we try η = 0.01, 0.005, 0.001 and report the best top-1 test accuracy.

**Does fine-tuning with model zoo perform better than fine-tuning a Imagenet expert?** While fine-tuning an Imagenet pre-trained model is standard and works well on most target tasks, we show that by fine-tuning models of a large model-zoo we can indeed obtain a lower test error on some target tasks (see models highlighted by black ← in Fig. 2). The reduction in error is more pronounced in the low-data regime. Therefore, we establish that maintaining a model zoo of models trained on different datasets is helpful to transfer to a diverse set of target tasks with different amounts of training data.

We demonstrate gains in the low-data regime by training on a smaller subset of the target task, with only 20, 5 samples per class in Fig. 2 (i.e., we train in a 20-shot and 5-shot setting). In few-shot cases we still test on the full test set.

**Fine-tuning with multi-domain expert.** In Sec. 4.1, we show that fine-tuning can be done by choosing different

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4https://gluon-cv.mxnet.io/api/model_zoo.html
domain-specific parameters within the multi-domain expert for fine-tuning. In Fig. 3, we fine-tune the multi-domain expert, i.e., Multi-BN of Tab. 2, on our target tasks by choosing different domain-specific parameters to fine-tune. Similar to Fig. 2, we show the accuracy gain obtained by fine-tuning multi-domain expert with respect to fine-tuning the standard Resnet-101 pre-trained on ImageNet. We observe that selecting the correct domain to fine-tune, i.e., the correct \( w_d \), where \( d \in \{1, 2, \ldots, D\} \) from multi-domain model zoo \( \mathcal{F} = \{ f_{w_1}, w_d \}_{d=1}^D \), is important to obtain high fine-tuning test accuracy on the target task. In Sec. 4.3, we show that model selection algorithms help in selecting the optimal domain-specific parameters for fine-tuning our multi-domain model zoo.

We also observe that fine-tuning with our multi-domain expert improves over the fine-tuning of single-domain model zoo for some tasks, e.g., iCassava: +1.4\% accuracy gain with multi-domain expert compared to +.72\% accuracy gain with single domain model expert over ImageNet expert. However, the comparison between single do-

Figure 3. Fine-tuning with the multi-domain expert for the full target task. We use the same notation as Fig. 2. For every target task (horizontal axis), we have 4 columns corresponding to fine-tuning different models from left to right: 1) ImageNet single and multi-domain expert in red, 2) Fine-tuning with different domains of multi-domain expert in green and 3) Single-domain Resnet-101 experts in blue, 4) We highlight multi-domain experts that obtain lower error than ImageNet single domain with black circle. Note, since our multi-domain expert is Resnet-101 based, we only use all our Resnet-101 experts for for fair comparison. Our observations are: i) We see gains over ImageNet expert (both single and multi-domain) by fine-tuning some (not all) domains of the multi-domain expert, for Magnetic Tile Defects, Oxford Flowers 102, Cucumber and iCassava target tasks. Therefore, it is important to pick the correct domain from the multi-domain expert for fine-tuning. ii) We observe the variance in error is smaller for fine-tuning with different domains of multi-domain experts, possibly due to shared parameters across domains, iii) Finally in some cases, e.g., Oxford Flowers 102 and iCassava, our multi-domain experts outperform both, all single domain and ImageNet experts. Figure is best viewed in high-resolution.

Figure 4. Model selection among single-domain experts. The heatmap shows the accuracy gain over Resnet-101 ImageNet expert obtained by fine-tuning the top-3 selected models for different model selection methods (column) on our target tasks (row). Higher values of gain are better. Note, for every method we fine-tune all the top-3 selected models (with same hyper-parameters as Sec. 4.2) and pick the one with the highest accuracy. Model selection performs better than “Worst Gain” and random selection. On average, LFC, LGC and LEEP [37] outperform Domain Similarity [11], RSA [15]. Feature Metrics [49] performs better than LFC, LEEP in high-data regime, but under-performs in the low-data regime.

Figure 5. Model Selection with multi-domain expert. The heatmap shows accuracy gain obtained by fine-tuning selected domain over fine-tuning ImageNet domain from the multi-domain expert. We show results for top-1 and top-3 selections. LFC, LEEP [37] are close to the best gain and they outperform Feature Metrics [49] and Random. main and multi-domain experts and their transfer properties is not the focus of our research and we refer the reader to [32, 44, 45].

4.3. Model Selection

In Sec. 4.2, using our benchmark we find that fine-tuning with a model zoo, both single-domain and multi-domain domain, improves the test accuracy on the target tasks. Now, we demonstrate that using a model selection algorithm we can select the best model or domain-specific parameters from our model zoos with only a few selections or trials.

Model Selection Algorithms. We use the following scores, \( S \), for our model selection methods: LFC (see \( S_{LF} \) defined in Sec. 3.3), LGC (see \( S_{LG} \) defined in (8)), which we introduce in Sec. 3.2. We compare against alternative measures of model selection and/or task similarity pro-
Fine-tuning top-3 models

| Shots      | Brute-force | Fine-tuning top-3 models | Model selection from single-domain model zoo |
|------------|-------------|--------------------------|---------------------------------------------|
|            | LFC         | LGC          | LEEP     | Feat. Met. | Dom. Sim. | LFC    | LGC    | LEEP    | Feat. Met. | Dom. Sim. |
| Full       | 48.17 ×     | 5.15 ×       | 3.89 ×   | 5.01 ×    | 6.02 ×    | 4.87 × | .41 ×  | 8.65 ×  | .02 ×      | .00 ×      | .40 ×    |
| 20-shot    | 41.67 ×     | 4.35 ×       | 3.40 ×   | 3.85 ×    | 4.86 ×    | 4.11 × | 1.09 × | 15.26 × | 0.03 ×     | 0.00 ×     | 1.31 ×   |

Table 3. Computation cost of model selection and fine-tuning the selected models from single-domain model zoo. We measure the average run-time for all our target tasks (of Fig. 2) of: Brute-force fine-tuning and Fine-tuning with 3 models chosen by model selection (Fig. 4). We divide the run-time by the run-time of fine-tuning a Resnet-101 Imagenet expert. For the single domain model zoo, brute-force fine-tuning of all 30 experts requires 40× more computation than fine-tuning Imagenet Resnet-101 expert. Note, Densest-169 models in our model zoo need more computation to fine-tune than Resnet-101, therefore the gain is > 30× for our model zoo size of 30. With model selection, we can fine-tune with selected models in only 3 – 6× the computation. LFC and LEEP compute model selection scores for 30 models in our zoo with < 1× the computation of fine-tuning Imagenet Resnet-101 expert. LGC model selection is expensive due to backward passes and large dimension of the gradient vector. However, our LFC approximation to LGC is good at selecting models (Fig. 4) and fast.

Figure 6. In (a), we measure the number of trials to select the best model, i.e., highest accuracy, from the model zoo. LFC, LGC and LEEP [37] require fewer trials than Domain Similarity [11], RSA [15] and Random selection baselines. In (b), we show that model selection scores of LFC obtain the highest Spearman’s ranking correlation to the actual fine-tuning accuracy compared to other model selection methods. Model selection scores are proxy for fine-tuning accuracy, therefore high correlation is desirable.

Is fine-tuning with model selection faster than brute-force fine-tuning? In Tab. 3, we show that brute-force fine-tuning is expensive. We can save computation by performing model selection using LFC and LEEP and fine-tuning only the selected top-3 models.

How many trials to select the model with best fine-tuning accuracy? In Fig. 6, we measure the average of selections or trials, across all target tasks, required to select the best model for fine-tuning from the model zoo. The best model corresponds to the highest fine-tuning test accuracy on target task. Our label correlation and LEEP [37] methods can select the best model in < 7 trials for our single domain model zoo of 30 experts and in < 3 trials for the multi-domain model zoo with 8 domain experts.

Are model selection scores a good proxy for fine-tuning accuracy? In Fig. 6, we show our LFC scores have the highest Spearman’s ranking correlation to the actual fine-tuning accuracy for different experts. Note, we average the correlation for all our target tasks. Our LFC score is a good proxy for ranking by fine-tuning accuracy and it can allow us to select (or reject) models for fine-tuning.

5. Conclusions

Fine-tuning using model zoo is a simple method to boost accuracy. We show that while a model zoo may have modest gains in the high-data regime, it outperforms Imagenet experts networks in the low-data regime. We show that simple baseline methods derived from a linear approximation of fine-tuning – Label-Gradient Correlation (LGC) and Label-Feature Correlation (LFC) – can select good models (single-domain) or parameters (multi-domain) to fine-tune, and match or outperform relevant model selection methods in the literature. Our model selection saves the cost of brute-force fine-tuning and makes model zoos viable.
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Proof of Proposition 1. The proof follows easily from [29]. We summarize the steps to make the section self-contained. Assuming, as we do, that the network is trained with a gradient flow (which is the continuous limit of gradient descent for small learning rate), then the weights and activations of the linearized model satisfies the differential equation:

$$
\dot{w}_t = -\eta \nabla_w f_w(\mathbf{x})^T \nabla f_w(\mathbf{x}) \mathcal{L}
$$

$$
\dot{f}_t(\mathbf{x}) = -\eta \nabla_w f_w(\mathbf{x})^T \nabla f_w(\mathbf{x}) \mathcal{L} = -\eta \Theta \nabla f_w(\mathbf{x}) \mathcal{L}
$$

For the MSE loss $$\mathcal{L} := \sum_{i=1}^N (y_i - f_t^{lin}(x_i))^2$$, the second differential equations become a first order linear differential equation, which we can easily solve in close form. The solution is

$$f_t^{lin}(\mathbf{x}) = (I - e^{-\eta\Theta t}) \mathcal{L} + e^{-\eta\Theta t} f_0(\mathbf{x}).$$

Putting this result in the expression for the loss at time $$t$$ gives

$$
\mathcal{L}_t = \sum_{i=1}^N (y_i - f_t^{lin}(x_i))^2
= (\mathcal{L} - f_t^{lin}(\mathbf{x}))^T (\mathcal{L} - f_t^{lin}(\mathbf{x}))
= (\mathcal{L} - f_{w_0}(\mathbf{x}))^T e^{-\eta\Theta t} (\mathcal{L} - f_{w_0}(\mathbf{x})),
$$

as we wanted.

Proof of using feature approximation in kernel. Using the notation $$\mathbb{E}_{i,j}[a_{ij}] := \frac{1}{N} \sum_{i,j=1}^N a_{ij}$$ we have

$$
\mathcal{Y}^T \mathcal{Y} = N^2 \mathbb{E}_{i,j}[y_i y_j \Theta_{ij}]
= N^2 \mathbb{E}_{i,j}[y_i y_j \nabla_w f_w(x_i) \cdot \nabla_w f_w(x_j)]
$$

Now let’s consider an $$f_w$$ in the form of a DNN, that is $$f_w(x) = W_L \phi(W_{L-1} \ldots \phi(W_0 x)).$$ By the chain rule, the gradient of the weights at layer $$l$$ is given by:

$$\nabla_{W_l} f_w(x) = J_{l+1}(x) \otimes f'_l(x)$$

where $$J_{l+1}$$ is the gradient of the output pre-activations coming from the upper layer and $$f'_l(x)$$ are the input activations at layer $$l$$ and “$$\otimes$$” denotes the Kronecker’s product or, equivalently since both are vectors, the outer product of the two vectors. Recall that $$\|A \otimes B\|_2 = \|A\|_2 \|B\|_2$$, which will be useful later. Using this, we can rewrite $$\mathcal{Y}^T \mathcal{Y}$$ as follows:

$$
\mathcal{Y}^T \mathcal{Y} = N^2 \mathbb{E}_{i,j}[y_i y_j \nabla_w f_w(x_i) \cdot \nabla_w f_w(x_j)]
= N^2 \mathbb{E}_{i,j}[\nabla_w f_w(x_i)] \cdot \mathbb{E}_{j} [\nabla_w f_w(x_j)]
= N^2 \sum_{l=1}^L \mathbb{E}_{i,j}[y_i J_{l+1}(x_i) \otimes f'_l(x_i)] \cdot \mathbb{E}_{j} [y_j J_{l+1}(x_j) \otimes f'_l(x_j)]
$$

We now introduce a further approximation and assume that $$J_{l+1}$$ is uncorrelated from $$f'_l(x_l).$$ The same assumption is used by [34] (see Section 3.1) who also provide theoretical and empirical justifications. Using this assumption, we have:

$$
\mathcal{Y}^T \mathcal{Y} = N^2 \sum_{l=1}^L \left\| \mathbb{E}_{i,j}[y_i J_{l+1}(x_i) \otimes f'_l(x_i)] \right\|^2
= N^2 \sum_{l=1}^L \left\| \mathbb{E}_{i,j}[J_{l+1}(x_i)] \otimes \mathbb{E}_{j} [f'_l(x_j)] \right\|^2
= N^2 \sum_{l=1}^L \left\| \mathbb{E}_{i,j}[J_{l+1}(x_i)] \right\|^2 \left\| \mathbb{E}_{j} [f'_l(x_j)] \right\|^2
$$

The term $$\mathbb{E}_{i,j}[y_j f'_l(x_j)]$$ measures the correlation between each individual feature and the label. If features are correlated with labels, then $$\mathcal{Y}^T \mathcal{Y}$$ is larger, and hence initial convergence is faster. Note that we need not consider only the last layer, convergence speed is determined by the correlation at all layers. Note however that the contribution of earlier layers is discounted by a factor of $$\left\| \mathbb{E}_{i,j}[J_{l+1}(x_i)] \right\|^2.$$ As we progress further down the network, the average of the gradients may become increasingly smaller, decreasing the term $$\left\| \mathbb{E}_{i,j}[J_{l+1}(x_i)] \right\|^2$$ and hence diminishing the contribution of earlier layer clustering to convergence speed.

Appendix B. Datasets

We choose our source and target datasets such that they cover different domains, and are publicly available for download. Detailed data statistics are in the respective citations for the datasets, and we include a few statistics e.g., training images, testing images, number of classes in Tab. 4. For all the datasets, if available we use the standard train and test split of the dataset, else we split the dataset randomly into 80% train and 20% test images. If images are indexed by URLs in the dataset, we download all accessible URLs with a python script.
| Dataset                        | Training Images | Testing Images | # Classes | URL                                                                 |
|-------------------------------|-----------------|----------------|-----------|----------------------------------------------------------------------|
| NWPU-RESISC45 [7]             | 25,200          | 63,000         | 43        | http://www.tensorflow.org/datasets/catalog/resisc45                   |
| Food-101 [6]                  | 75,750          | 25,250         | 101       | https://www.tensorflow.org/datasets/catalog/food101                   |
| Logo 2k [59]                  | 134,907         | 32,233         | 2341      | https://github.com/umfoundation/google-landmark                      |
| Goog. Landmark [39]           | 200,000         | 15,601         | 256       | https://github.com/min99595/Logos-2k-plus-Dataset                    |
| iNaturalist [21]              | 265,213         | 3030           | 1010      | https://github.com/vision/visualizations/landmark                    |
| iMaterialist [33]             | 965,782         | 9639           | 2019      | https://github.com/malongtech/imaterialist-product-2019              |
| Imagenet [13]                 | 1,281,167       | 50,000         | 1000      | http://image-net.org/download                                        |
| Places-365 [57]               | 1,803,460       | 36,500         | 365       | http://www-sop.inria.fr/members/Alexis.Joly/BelgaLogos/BelgaLogos.html |
| Magnetic Tile Defects [23]     | 1008            | 336            | 6         | http://www.vision.caltech.edu/visipedia/CUB-200-2011.html            |
| UC Merced Land Use [53]       | 1575            | 525            | 21        | https://www.tensorflow.org/datasets/catalog/ucmerced-landuse          |
| Oxford Flowers 102 [38]       | 2040            | 6149           | 102       | https://www.tensorflow.org/datasets/catalog/oxford-flower102          |
| Cucumber [12]                 | 2326            | 597            | 30        | https://github.com/workpikes/CUCUMBER-9                              |
| European Flood Depth [5]      | 3153            | 557            | 2         | https://github.com/vision/eu-flood-dataset                            |
| Oxford-IIT Pets [41]          | 3680            | 3669           | 37        | https://www.tensorflow.org/datasets/catalog/oxford-iit-pets           |
| Describable Textures [9]      | 4230            | 1410           | 47        | https://github.com/vision/eu-flood-dataset                            |
| iCassava [36]                 | 5367            | 280            | 5         | https://sites.google.com/view/fgvc6/competitions/icassava-2019        |
| CUB-200 [52]                  | 5994            | 5793           | 200       | http://www.vision.caltech.edu/visipedia/CUB-200-2011.html            |
| Belga Logos [25]              | 7100            | 2300           | 27        | http://www.vision.caltech.edu/visipedia/CUB-200-2011.html            |
| Stanford Cars [27]            | 3444            | 8041           | 196       | http://www.tensorflow.org/datasets/catalog/cars                       |

Table 4. The number of training images, testing images and classes as well as the URL to download the dataset are listed above. The top part contains our source datasets used to train the model zoo and the bottom part lists our target tasks used for fine-tuning and model selection with our model zoo.

Appendix C. Details of model selection methods

Domain Similarity [11]. As per [11], we extract avg. features for every class for source and target datasets using pre-trained model. We compute an earth movers distance between these average class vectors and convert them to domain similarity score. We use the code provided by the authors at https://github.com/richardaecn/cvpr18-inaturalist-transfer. We exclude classes with less than 5 training images for Earth-Movers Distance computation.

RSA [15]. Following the procedure outlined in [15], we extract features before the classification layer (e.g. 2048 dim features of Resnet-101 after average pool) for images in the target dataset. We denote this set of features as \( f(x) \), \( \forall (x, y) \in D \). We build a representation dissimilarity matrix (RDM) as follows:

\[
\text{rdm}_{f}(i, j) = 1 - \text{correlation}(f(x_i), f(x_j)) \tag{9}
\]

We train a small neural network \( f_{small} \) on target dataset. Note, this is much cheaper to train than fine-tuning the model zoo. Features are extracted from \( f_{small} \) and we build another rdm:

\[
\text{rdm}_{f_{small}}(i, j) = 1 - \text{correlation}(f_{small}(x_i), f_{small}(x_j)) \tag{10}
\]

If rdm’s of trained small network \( f_{small} \) and our pre-trained model \( f \) are similar, then the pre-trained model is a good candidate for fine-tuning with target dataset. The final RSA model selection score is:

\[
S_{RSA}(f, D) = \text{spearmanr}(\text{rdm}_f, \text{rdm}_{f_{small}}) \tag{11}
\]

Since the method requires training a small neural network on target task, we train a Resnet-18 as the small neural network with the same fine-tuning configuration used in Section 4.1 of the paper with initial learning rate = .005.

Feature Metrics [49]. Features are extracted for all images of target dataset from pre-trained model, i.e. \( f(x), \forall x \in D \). We use same features as RSA, our LFC/LGC and compute variance, sparsity metrics of [49]. We use the sparsity metrics as model selection score, \( S_{\text{Feat. Metrics}}(f, D) = \text{sparsity}(f(x), \forall x \in D) \). Note, we use the optimal linear combination of the two sparsity metrics proposed in the paper. For feature metrics, the hypothesis is that if the pre-trained model generates more sparse representations, they are can generalize with fine-tuning to the target task.

LEEP [37]. LEEP builds an empirical classifier from source dataset label space to target dataset label space using base model \( f \). The likelihood of target dataset \( D \) under this empirical classifier is the model selection score for the pre-trained model and target dataset. See [37] for a detailed explanation.

Appendix D. Different dataset size for model selection

In Fig. 7, we perform an ablation study on different sampling size of the target task used for model selection. We find that, our choice of 25 samples per class for model selection, suffices to select good models to fine-tune in top-3 selections at low-computational cost.
Figure 7. **Ablation study of dataset size for model selection.** Above we use 25, 50-samples per class and full target task to perform model selection with different methods. We plot accuracy gain vs. Imagenet expert for top-3 selected models for every method (similar to Fig. 4 of the paper). The accuracy gain increases for LFC, LEEP and RSA with more samples of the target task. However, we see that even as small as 25 samples suffice to obtain good accuracy gain with low computational cost.

**Appendix E. Visualization of $\Theta_F$ with fine-tuning**

In Fig. 8, we plot the feature correlation matrix for different pre-trained models across different epochs of fine-tuning (i.e. $0^{th}$, $15^{th}$, $30^{th}$ epoch) for the UC Merced Land Use [53] target task. We see that the pre-trained model on NWPU-RESISC45 [7], exhibits the ideal correlation wherein features of the images with the same class are correlated and features of images with different classes are uncorrelated. This NWPU-RESISC45 [7] also has the highest LFC score.
Figure 8. We plot the feature correlation matrix, $\Theta_F$, for different pre-trainings (row) and different epochs (columns) of fine-tuning. Above, we fine-tune on the UC Merced Land Use [53] dataset comprising of aerial images. Images with same class label, 25 images per class, are grouped along the vertical/horizontal axis. Since, features of the same class should be correlated and features of different classes should be uncorrelated, the matrix is expected to have higher values along block diagonal and zero elsewhere. We observe that the matrix exhibits this ideal behaviour for pre-training on semantically related domain (aerial images) of NWPU-RESISC45 [7] (top row) and has highest LFC score for this pre-training.