# Maestro: Uncovering Low-Rank Structures via Trainable Decomposition

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# **Abstract**

Deep Neural Networks (DNNs) have been a large driver for AI breakthroughs in recent years. However, these models have been getting increasingly large as they become more accurate and safe. This means that their training becomes increasingly costly and time-consuming and typically yields a single model to fit all targets. Various techniques have been proposed in the literature to mitigate this, including pruning, sparsification, or quantization of model weights and updates. While achieving high compression rates, they often incur significant computational overheads at training or lead to non-negligible accuracy penalty. Alternatively, factorization methods have been leveraged for low-rank compression of DNNs. Similarly, such techniques (e.g., SVD) frequently rely on heavy iterative decompositions of layers and are potentially sub-optimal for non-linear models, such as DNNs. We take a further step in designing efficient low-rank models and propose MAESTRO, a framework for trainable low-rank layers. Instead of iteratively applying a priori decompositions, the low-rank structure is baked into the training process through LoD, a low-rank ordered decomposition. Not only is this the first time importance ordering via sampling is applied on the decomposed DNN structure, but it also allows selecting ranks at a layer granularity. Our theoretical analysis demonstrates that in special cases LoD recovers the SVD decomposition and PCA. Applied to DNNs, MAESTRO enables the extraction of lower footprint models that preserve performance. Simultaneously, it enables the graceful trade-off between accuracy-latency for deployment to even more constrained devices without retraining.

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#### 1. Introduction

Deep Learning has been experiencing an unprecedented uptake, with models achieving a (super-)human level of performance in several tasks across modalities, giving birth to even more intelligent assistants (Radford et al., 2023) and next-gen visual perception and generative systems (Radford et al., 2021). However, the price of this performance is that models are getting significantly larger, with training and deployment becoming increasingly costly (Laskaridis et al., 2024). Therefore, techniques from Efficient ML become evermore relevant (Wan et al., 2023), and a requirement for deployment in constrained devices, such as smartphones or IoT devices (Laskaridis et al., 2022).

Typical techniques to compress the network involve i) quantization, i.e., reducing precision of the model (Wang et al., 2019) or communicated updates (Seide et al., 2014; Alistarh et al., 2017), ii) pruning the model during training, e.g., through Lottery Ticket Hypothesis (LTH) (Frankle & Carbin, 2019), iii) sparsification of the network representation and updates, i.e., dropping the subset of coordinates (Suresh et al., 2017; Alistarh et al., 2018) or iv) lowrank approximation (Wang et al., 2021; Dudziak et al., 2019), i.e. keeping the most relevant ranks of the decomposed network. Despite the benefits during deployment, that is a lower footprint model, in many cases, the overhead during training time or the accuracy degradation can be non-negligible. Moreover, many techniques can introduce multiple hyperparameters or the need to fine-tune to recover the lost accuracy.

In this work, we focus training low-rank factorization. Specifically, we pinpoint the challenges of techniques (Wang et al., 2021; 2023) when decomposing the parameters of each layer in low-rank space and the need to find the optimal ranks for each one at training time. To solve this, we propose LOD (Low-rank ordered Decomposition), a non-trivially extended version of Ordered Dropout technique from Horváth et al. (2021), applied to progressively find the optimal decomposition for each layer of a DNN while training (Fig. 1). Critical differences to prior work include *i*) the non-uniformity of the search space (i.e. we allow for different ranks per layer), *ii*) the trainable aspect of the decomposition to reflect the data distribution, and *iii*) the gains to training and deployment time without sacrificing

accuracy. Nevertheless, we also provide a latency-accuracy trade-off mechanism to deploy the model on even more constrained devices.

Our contributions can be summarized as follows:

- We propose MAESTRO<sup>1</sup>, a novel layer decomposition technique that enables learning low-rank layers in a progressive manner while training. We fuse layer factorization and ordered dropout into LoD, an extended variant of the ordered dropout, in a novel manner, by embedding ordered importance directly into the factorized weights. By decomposing layers and training on stochastically sampled low-rank models, we apply ordered importance decomposed representation of each layer. We combine this with a hierarchical group-lasso term (Yuan & Lin, 2006) in the loss function to zero out redundant ranks and progressively shrink the rank space. This way, we enable computationally efficient training achieved by the proposed decomposition without relying on inexact and potentially computationally expensive iterative decompositions such as Singular Value Decomposition (SVD).
- MAESTRO is fundamentally a theoretically motivated approach that embeds decomposition into training. First, we show that our new objective can recover *i*) the SVD of the target linear mapping for the particular case of uniform data distribution and *ii*) the Principal Component Analysis (PCA) of the data in the case of identity mapping.
- As MAESTRO's decomposition is part of the training process, it also accounts for data distribution and the target function, contrary to SVD, which operates directly on learned weights. We show that this problem *already arises* for a simple linear model and empirically generalize our results in the case of DNNs, by applying our method to different types of layers (including fully-connected, convolutional, and attention) spanning across three datasets and modalities.
- We illustrate that our technique achieves better results than SVD-based baselines at a lower cost. Indicatively, MAESTRO can achieve on par or better results than lowrank SOTA methods on vision datasets (CIFAR-10, ImageNet) with lower training overhead due to progressive shrinking, while at the same time it reaches 6% lower perplexity at a quarter of the computational cost and half of the parameters of SVD-variants in Transformer models.

## 2. Related Work

The topic of Efficient ML has received a lot of attention throughout the past decade as networks have been getting increasingly computationally expensive. We distinguish between training and deployment time, with the latter having a more significant impact and thus amortizes the potential overhead during training. Nevertheless, cost optimisation is evermore relevant for training large models, and the advent of Federated Learning (McMahan et al., 2017), efficient training becomes increasingly relevant to remain tractable.

**Efficient inference.** For efficient deployment, various techniques have been proposed that either optimize the architecture of the DNN in a hand-crafted (Howard et al., 2017) or automated manner (i.e. NAS) (Tan & Le, 2019), they remove redundant computation by means of pruning parts of the network (Han et al., 2015; Carreira-Perpinán & Idelbayev, 2018; Frankle & Carbin, 2019; Chen et al., 2021; Sreenivasan et al., 2022; Li et al., 2016; Wen et al., 2016; Hu et al., 2016; Wen et al., 2016; Zhu & Gupta, 2017; He et al., 2017; Yang et al., 2017; Liu et al., 2018; Yu & Huang, 2019b), in a structured or unstructured manner, or utilise low-precision representation (Wang et al., 2019) of the neurons and activations. However, such techniques may involve non-negligible training overheads or lack flexibility of variable footprint upon deployment. Closer to our method, there have been techniques leveraging low-rank approximation (e.g. SVD) for efficient inference (Xue et al., 2013; Sainath et al., 2013; Jaderberg et al., 2014; Wiesler et al., 2014; Dudziak et al., 2019). Last, there is a category of techniques that dynamically resize the network at runtime for compute, memory or energy efficiency, based on early-exiting (Laskaridis et al., 2021) or dynamic-width (Yu et al., 2019) and leverage the accuracy-latency tradeoff.

Efficient training. On the other hand, techniques for efficient training become very relevant nowadays when scaling DNNs sizes (Hu et al., 2021) or deploying to embedded devices (Lin et al., 2022), and oftentimes offer additional gains at deployment time. Towards this goal, there have been employed methods where part of the network is masked (Sidahmed et al., 2021) or dropped (Alam et al., 2022; Caldas et al., 2019; Wu et al., 2018) during training, with the goal of minimizing the training footprint. Similarly to early-exiting, multi-exit variants for efficient training (Kim et al., 2023; Liu et al., 2022) have been proposed, and the same applies for width-based scaling (Horváth et al., 2021; Diao et al., 2021). Last but not least, in the era of transformers and LLMs, where networks have scaled exponentially in size, PEFT-based techniques, such as adapterbased fine-tuning (Houlsby et al., 2019) (such as LoRA (Hu et al., 2021)), become increasingly important and make an important differentiator for tackling downstream tasks.

Learning ordered representation. Ordered Dropout (OD) was proposed as a mechanism for importance-based pruning for the easy extraction of sub-networks devised to allow for heterogeneous federated training (Horváth et al., 2021). Similar constructions were proposed to be applied in the representation layer of autoencoders (Rippel et al., 2014) to en-

<sup>&</sup>lt;sup>1</sup>The implementation can be found here: https://github.com/SamuelHorvath/Maestro-LoD

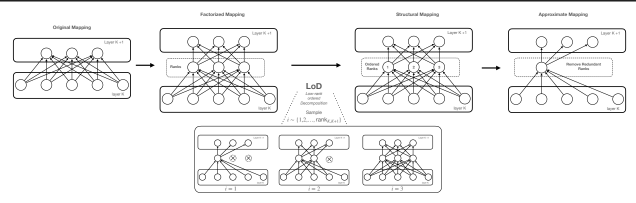


Figure 1: MAESTRO's construction. To obtain low-rank approximation, the given linear map is decomposed and trained with LoD to obtain an ordered representation that can be efficiently pruned.

force identifiability of the learned representation or the last layer of the feature extractor (Horváth et al., 2021) to learn an ordered set of features for transfer learning. Contrary to prior work, MAESTRO's LOD non-trivially extends ordered representation in three meaningful ways. First, it is the first work that is applied to the decomposed network, which gets progressively shrunk as redundant ranks converge to zero. This is achieved through hierarchical group lasso penalty, as described in Sec. 3.3. Second, LOD allows for heterogeneous (non-uniform) ranks per layer, yielding a much richer operating space. Last, through LOD we can leverage the ordered representation of ranks at inference time to further compress the model, allowing a graceful accuracy-latency tradeoff for deployment on more constrained devices, without the need to retrain.

#### 3. MAESTRO

In this work, we focus on low-rank models as a technique to reduce the neural network model's computational complexity and memory requirements. The main challenge that we face is the selection of the optimal rank or the trade-off between the efficiency and the rank for the given layer. Therefore, we devise an *importance-based* training technique, MAESTRO, which learns not only a mapping between features and responses but also *learns the decomposition* of the trained network. This is achieved by factorizing all the layers in the network.

#### 3.1. Formulation

**Low-rank approximation.** Our inspiration comes from the low-rank matrix approximation of a matrix  $A \in \mathbb{R}^{m \times n}$ . For simplicity, we assume that A has rank at most  $r = \min\{m,n\}$  with  $k \leq r$  distinct non-zero singular values  $\tilde{\sigma}_1 > \tilde{\sigma}_2 > \ldots > \tilde{\sigma}_k > 0$ , with corresponding left and right singular vectors  $\tilde{u}_1, \tilde{u}_2, \ldots, \tilde{u}_k \in R^m$  and  $\tilde{v}_1, \tilde{v}_2, \ldots, \tilde{v}_k \in R^n$ , respectively. For such a matrix, we can rewrite its best l-rank approximation as the following minimization

problem

$$\min_{U \in \mathbb{R}^{m \times l}, V \in \mathbb{R}^{n \times l}} \left\| \sum_{i=1}^{l} u_i v_i^{\top} - A \right\|_F^2 \tag{1}$$

where  $c_i$  denotes the i-th column of matrix C and  $\|\cdot\|_F$  denotes Frobenius norm. We note that Problem (1) is nonconvex and non-smooth. However, (Ye & Du, 2021) showed that the randomly initialized gradient descent algorithm solves this problem in polynomial time. In this work, we consider the best rank approximation across all the ranks. Eckart–Young–Mirsky theorem leads to the following objective

$$\min_{U \in \mathbb{R}^{m \times r}, V \in \mathbb{R}^{n \times r}} \frac{1}{r} \sum_{b=1}^{r} \left\| U_{:b} V_{:b}^{\top} - A \right\|_{F}^{2}, \quad (2)$$

where  $C_{:b}$  denotes the first b columns of matrix C. This objective, up to scaling, recovers SVD of A exactly, and for the case of distinct non-zero singular values, the solution is, up to scaling, unique (Horváth et al., 2021). This formulation, however, does not account for the data distribution, i.e., it cannot tailor the decomposition to capture specific structures that appear in the dataset.

**LoD for data-dependent low-rank approximation.** Therefore, the next step of our construction is to extend this problem formulation with data that can further improve compression, reconstruction, and generalization and incorporate domain knowledge. We assume that data comes from the distribution  $x \sim \mathcal{X}$  centered around zero, i.e.,  $\mathbf{E}_{x \sim \mathcal{X}}[x] = 0.^2$ , and the response is given by y = Ax. In this particular case, we can write the training loss as

$$\min_{U \in \mathbb{R}^{m \times r}, V \in \mathbb{R}^{n \times r}} \mathbf{E}_{x, y \sim \mathcal{X}} \left[ \sum_{b=1}^{r} \frac{1}{r} \left\| U_{:b} V_{:b}^{\top} x - y \right\|^{2} \right]. \quad (3)$$

It is important to note that the introduced problem formulation (3) for the neural network with a single hidden layer and no activations can be solved using stochastic algorithms

<sup>&</sup>lt;sup>2</sup>We make this assumption for simplicity. It can be simply overcome by adding a bias term into the model.

by sampling from the data distribution  $\mathcal{X}$  (subsampling) and rank distribution  $\mathcal{D}$ . When we apply LoD to DNNs, contrary to any prior work (Horváth et al., 2021; Rippel et al., 2014; Diao et al., 2021), our formulation is the first one to apply importance-based ordered dropout to the decomposed representation of each layer, thus preserving the dimensionality. More importantly, we decompose each layer *independently* and allow for finding the optimal rank per layer in a data-informed manner. We discuss details in the next paragraph.

**DNN low-rank approximation.** For Deep Neural Networks (DNNs), we seek to uncover the optimal ranks for a set of d linear mappings  $W^1 \in \mathbb{R}^{m_1 \times n_1}, \ldots, W^d \in \mathbb{R}^{m_d \times n_d}$ , where  $W^i$ 's are model parameters and d is model depth, e.g., weights corresponding to linear layers<sup>3</sup>, by decomposing them as  $W^i = U^i \left(V^i\right)^\top$ . We discuss how these are selected in the next section. To decompose the network, we aim to minimize the following objective

$$\mathbf{E}_{x,y\sim\mathcal{X}} \left[ \frac{1}{\sum_{i=1}^{d} r_i} \sum_{i=1}^{d} \sum_{b=1}^{r_i} l(h(U^1(V^1)^\top, \dots, U^i(V^b)^\top, \dots, U^d(V^d)^\top, W^o, x), y) \right],$$
(4)

where  $r_i = \min\{m_i, n_i\}$ , l is a loss function, h is a DNN, and  $W^o$  are the other weights that we do not decompose. We note that our formulation aims to decompose each layer, while decompositions across layers do not directly interact. The motivation for this approach is to uncover low-rank structures within each layer that are not affected by inaccuracies from other layers due to multiple low-rank approximations.

#### 3.2. Layer Factorization

The following sections discuss how we implement model factorization for different architectures.

**FC layers.** A 2-layer fully connected (FC) neural network can be expressed as  $f(x) = \sigma(\sigma(xW_1)W_2)$ , where Ws are weight matrices of each FC layer, and  $\sigma(\cdot)$  is any arbitrary activation function, e.g., ReLU. The weight matrix W can be factorized as  $UV^\top$ .

**CNN layers.** For a convolution layer with dimension,  $W \in \mathbb{R}^{m \times n \times k \times k}$  where m and n are the number of input and output channels, and k is the size of the convolution filters. Instead of directly factorizing the 4D weight of a convolution layer, we factorize the unrolled 2D matrix. Unrolling the 4D tensor W leads to a 2D matrix with shape  $W_{\text{unrolled}} \in \mathbb{R}^{mk^2 \times n}$ , where each column represents the weight of a vectorized convolution filter. Factorization can

#### Algorithm 1: MAESTRO (Training Process)

```
Input: epochs E, dataset \mathcal{D}, model h parametrized by
                     U^{1} \in \mathbb{R}^{m_{1} \times r_{1}}, 
V^{1} \in \mathbb{R}^{n_{1} \times r_{1}}, \dots, U^{d} \in \mathbb{R}^{m_{d} \times r_{d}}, V^{d} \in \mathbb{R}^{n_{d} \times r_{d}},
                     W^{o}, and hyperparameters \lambda_{al}, \varepsilon_{ps}
  1 for t \leftarrow 0 to E - 1 do // Epochs
              for (x,y) \in \mathcal{D} do // Iterate over dataset
                       Sample (i,b) \sim \{\{(i,b)\}_{b=1}^{r_i}\}_{i=1}^d; // LoD
                         l(h(U^{1}(V^{1})^{\top}, \dots, U^{i}_{:b}(V^{i}_{:b})^{\top}, \dots, U^{d}(V^{d})^{\top}, W^{o}, x),
y) + \lambda_{gl} \sum_{i=1}^{d} \sum_{b=1}^{r_{i}} (||U^{i}_{b:}|| + ||V^{i}_{b:}||) // Loss
                       L.backward() // Update weights
              end
              for i \leftarrow 1 to d do
                       for b \leftarrow 1 to r_i do
                               // rank importance thresholding
                                \begin{aligned} & \text{if } \left\| V_{b:}^{i} \right\| \left\| U_{b:}^{i} \right\| \leq \varepsilon_{ps} \text{ then} \\ & \left| \quad r_{i} = b - 1 \text{ // progressive shrinking} \right| \end{aligned} 
                                        hreak
                               end
                       end
15
              end
16 end
```

then be conducted on the unrolled 2D matrix; see (Wang et al., 2021) for details.

**Transformers.** A Transformer layer consists of a stack of encoders and decoders (Vaswani et al., 2017). The encoder and decoder contain three main building blocks: the multihead attention layer, position-wise feed-forward networks (FFN), and positional encoding. We factorize all trainable weight matrices in the multi-head attention (MHA) and the FFN layers. The FFN layer factorization can directly adopt the strategy from the FC factorization. A p-head attention layer learns p attention mechanisms on the key, value, and query (K, V, Q) of each input token:

$$MHA(Q, K, V) = Concat(head_1, ..., head_p)W^O$$
. Each head performs the computation of:

$$\begin{aligned} \text{head}_i &= \text{Attention}(QW_Q^{(i)}, KW_K^{(i)}, VW_V^{(i)}) \\ &= \text{softmax}\left(\frac{QW_Q^{(i)}W_K^{(i)\top}K^\top}{\sqrt{d/p}}\right)VW_V^{(i)}. \end{aligned}$$
 where  $d$  is the hidden dimension. The trainable weights

where d is the hidden dimension. The trainable weights  $W_Q^{(i)}, W_K^{(i)}, W_V^{(i)}, i \in \{1, 2, \dots, p\}$  can be factorized by simply decomposing all learnable weights  $W \cdot$  in an attention layer and obtaining  $U \cdot V^\top \cdot$  (Vaswani et al., 2017).

# 3.3. Training Techniques

Having defined the decomposition of typical layers found in DNNs, we move to formulate the training procedure of our method, formally described in Algorithm 1. Training the model comprises an iterative process of propagating forward on the model by *sampling a rank*  $b_i$  per decomposed layer i

<sup>&</sup>lt;sup>3</sup>We can apply our decomposition on different types of layers, such as Linear, Convolutional and Transformers as shown in Sec. 3.2.

up to maximal rank  $r_i$  (line 3). We calculate the loss, which integrates an additional *hierarchical group lasso* component (lines 4) and *backpropagate* on the sampled decomposed model (line 5). At the end of each epoch, we *progressively shrink* the network by updating the maximal rank  $r_i$ , based on an importance threshold  $\varepsilon_{ps}$  (line 11). We provide more details about each component below.

**Efficient training via sampling.** In Sec. 4, we show that for the linear case (3), the optimal solution corresponds to PCA over the linearly transformed dataset. This means that the obtained solution contains orthogonal directions. This property is beneficial because it directly implies that when we employ gradient-based optimization, not only is the gradient zero at the optimum, but the gradient with respect to each summand in Equation (3) is also zero. The same property is directly implied by overparametrization (Ma et al., 2018) or strong growth condition (Schmidt & Roux, 2013). As a consequence, this enables us to sample only one summand at a time and obtain the same quality solution. When considering (4) as an extension to (3), it is unclear whether this property still holds, which would also imply that the set of stationary points of (3) is a subset of stationary points of the original objective without decomposition. However, in the experiments, we observed that sampling is sufficient to converge to a good-quality solution. If this only holds approximately, one could leverage fine-tuning to recover the loss in performance.

#### Efficient rank extraction via hierarchical group-lasso.

By definition, (3) leads to an ordered set of ranks for each layer. This ordered structure enables efficient rank extraction and selection. To effectively eliminate unimportant ranks while retaining the important ones, thus leading to a more efficient model, we consider Hierarchical Group Lasso (HGL) (Lim & Hastie, 2015) in the form

$$\lambda_{gl} \sum_{i=1}^{d} \sum_{b=1}^{r_i} (\|U_{b:}^i\| + \|V_{b:}^i\|), \tag{5}$$

where  $C_{b:}$  denotes the matrix that contains all the columns of C except for the first b-1 columns.

**Progressive shrinking.** HGL encourages that unimportant ranks become zero and can be effectively removed from the model. To account for this, for each layer we remove  $V_{b}^i$  and  $U_{b}^i$  (i.e., set  $r_i = b-1$ ) if  $\|V_{b}^i\|\|U_{b}^i\| \le \varepsilon_{ps}$ , where  $\varepsilon_{ps}$  is a pre-selected threshold – and a hyperparameter of our method.

**Hyperparameter optimization.** We provide an algorithm for finding the optimal value for the hyperparameter  $\lambda_{gl}$  in Alg. 2. From the evaluation of Tab. 11-15, this strategy typically requires at most 2-3 times the computational effort (in terms of FLOPs) compared to a single training loop with an optimally chosen  $\lambda_{gl}$ . This is significantly easier than

tuning the per-layer maximal rank and the pretraining steps, where the full-rank model is being pretrained, as is the case in other low-rank baselines. Equivalently, the value of  $\varepsilon_{ps}$  represents the effective zero-point in our algorithm, typical value of which was 1e-7 in our experiments.

## Algorithm 2: MAESTRO (Hyper-parameter optimization)

```
Input: constraints (e.g., min required accuracy, max
            #parameters), epochs E, dataset \mathcal{D}, model h,
            evaluation frequency E_{\text{every}}^{\text{eval}}, \varepsilon_{ps}, limits for HPO:
            largeValue, smallValue
 1 \lambda_{gl} = largeValue
2 old_model = NULL
   while \lambda_{al} > smallValue do
        model = RandInit(model)
        for t \leftarrow 0 to E - 1 do // Epochs
             Train(model, dataset, \lambda_{gl})
             if t \mod E_{\text{every}}^{\text{eval}} == 0 then
                  acc = CalculateAcc(model, dataset)
                  flops, params = MeasureFootprint(model, \varepsilon_{ps})
                  if acc \ge constraints[`acc'] and params \le
                    constraints['params'] then
                       return model // model satisfying constraints
                         was found
                  if params < few_params then
                      break // model too sparse
                  end
             end
        end
17
        flops, params = MeasureFootprint(model, \varepsilon_{ps})
        if params > constraints['params'] then
             /** no model satisfies constraints, return the last
              model satisfying parameters constraints**/
             return old model
        old_model = Copy(model)
        \lambda_{gl} = \lambda_{gl} / 2
25 end
```

**Initialization.** Initialization is a key component of the training procedure (He et al., 2015; Mishkin & Matas, 2015). To adopt the best practices from standard non-factorized training, we follow a similar approach to (Khodak et al., 2021; Wang et al., 2021), where we first initialize the nonfactorized model using standard initialization. For initializing factorized layers, we use the Singular Value Decomposition of the non-factorized initialization – in a full-rank form – to ensure that the resulting product matrix is the same as the original parameter decomposition. In addition, SVD is an optimal decomposition for the linear case with uniform data. However, in contrast with the adaptive baseline method (Wang et al., 2023) we only decompose once, rather than on every training iteration. As such, we only run decomposition once and progressively shrink the ranks in a data-centric manner. This is contrary to related work (Wang et al., 2021; 2023) that requires manual rank and layer selection and full-rank warmup to achieve the

desired performance, at the cost of training overhead, of course.

#### 3.4. Train-Once, Deploy-Everywhere

Up until now, we have described how our method works for training low-rank models, which yield computational, memory, network, and energy (Wu et al., 2022) bandwidth benefits during training. At deployment time, one can directly deploy the final model (rank  $r_i$  for each layer) on the device, which we acquire from performing a threshold sweep of  $\varepsilon_{ps}$ over the effective range of rank importance across layers. However, in case we want to run on even more constrained devices, such as mobile or embedded (Almeida et al., 2021) systems, the learned decomposition also gives us the flexibility to further compress the model in a straightforward manner, effectively trading off accuracy for a smaller model footprint. Inspired by (Yu & Huang, 2019a), we propose to use greedy search. We begin with the current model and compare model performance across various low-rank models, each created by removing a certain percentage of ranks from each layer. We then eliminate the ranks that cause the least decrease in performance. This process is iterated until we reach the desired size or accuracy constraint. To make this approach efficient, we estimate the loss using a single mini-batch with a large batch size (e.g., 2048). This also avoids issues with BatchNorm layers; see (Yu & Huang, 2019a) for details.

In summary, MAESTRO comprises a technique for trainable low-rank approximation during training time that progressively compresses the model, reflecting the data distribution, and a method that enables a graceful trade-off between accuracy and latency for embedded deployment, by selecting the most important parts of the network. We validate these claims in Sec. 5.2 and 5.5, respectively.

#### 4. Theoretical Guarantees

In this section, we further investigate the theoretical properties of MAESTRO for the linear mappings, i.e., the setup of the problem formulation (3).

**Theorem 4.1** (Informal). Let  $A = \tilde{U}\tilde{\Sigma}\tilde{V}^{\top}$  be a SVD decomposition of A. Then, the minimization problem (3) is equivalent to PCA applied to the transformed dataset  $x \to \tilde{\Sigma}\tilde{V}^{\top}x$ ,  $x \sim \mathcal{X}$  projected on the column space of  $\tilde{U}$ .

The formal statement can be found in Appendix C. Theorem 4.1 shows that MAESTRO can adapt to data distribution by directly operating on data  $x \sim \mathcal{X}$  and also to the target mapping by projecting data to its right singular vectors scaled by singular values. In particular, we show that in the special case, when  $\mathcal{X}$  is the uniform distribution on the unit ball, (3), i.e., MAESTRO, exactly recovers truncated SVD of A, which is consistent with the prior results (Horváth et al., 2021). In the case A is the identity, it is straightforward to see that MAESTRO is equivalent to PCA. We can see

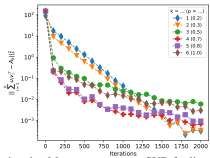
that MAESTRO can efficiently extract low-rank solutions by filtering out directions corresponding to the null space of the target mapping A and directions with no data. We also numerically verify both of the special cases–PCA and SVD, by minimizing (3) using stochastic gradient descent (SGD) with  $\mathcal{D}$  being the uniform distribution. These experiments are provided in Fig. 2a and 2b. We provide further evidence on the adaptivity of MAESTRO in Appendix E.1 and E.2.

We showed that MAESTRO could recover SVD in a particular case of the linear model and the uniform data distribution on the unit ball. We note that in this case, SVD is optimal, and we cannot acquire better decomposition. Therefore, it is desired that MAESTRO is equivalent to SVD in this scenario. More generally, we argue that MAESTRO decomposition should be preferable to SVD due to the following reasons:

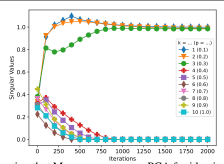
- MAESTRO formulation is *directly built into the training* and tailored to obtain the best low-rank decomposition, while SVD relies on linearity assumption.
- SVD does not account for data, and even in the linear NN case, the learned singular vectors might exhibit wrong ordering. We demonstrate this issue using a simple example where we take matrix A with rank 3. We construct the dataset  $\mathcal{X}$  in such a way that the third singular vector is the most important, the second one is the second, and the first is the third most important direction. Clearly, SVD does not look at data. Therefore, it cannot capture this phenomenon. We showcase that MAESTRO learns the correct order; see Fig. 6 of the Appendix.
- Pre-factorizing models allow us to apply *hierarchical* group-lasso penalty (Yuan & Lin, 2006) for decomposed weights to directly regularize the rank of different layers.
- SVD is computationally expensive and can only run rarely, while MAESTRO is directly built into the training and, therefore, does not require extra computations. In addition, MAESTRO supports rank sampling so training can be made computationally efficient.

# 5. Experiments

We start this section by describing the setup of our experiments, including the models, datasets and baselines with which we compare MAESTRO. We then compare MAESTRO against the baselines on accuracy and training Multiply-Accumulate operations (MACs) and discuss the results. Subsequently, we analyze the behaviour of our system in-depth and provide additional insights on the performance of our technique, along with an ablation study and sensitivity analysis to specific hyperparameters. Finally, we showcase the performance of models upon deployment and how we can derive a smaller footprint model with some accuracy tradeoff, without the need to fine-tune.



(a) Verification that MAESTRO recovers SVD for linear mapping with uniform data. We display the L2 distance between the best rank k and MAESTRO's approximation of mapping A. The target matrix was randomly generated  $9\times 6$  matrix with rank 3. p and k represent relative and actual rank.



(b) Verification that MAESTRO recovers PCA for identity mapping. The plot displays the estimates of singular values. The data distribution has only 3 directions. It is expected that the top 3 ranks will converge to value one and the rest to zero. p and k stand for relative and actual rank, respectively.

Figure 2: Empirical showcase of theoretical properties of the MAESTRO's formulation.

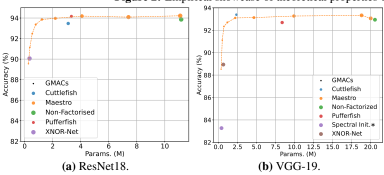


Figure 3: Maestro vs. baselines on CIFAR10. Spectral-Init results is taken from the original work; For XNOR-Net each weight is quantized from 32 to 1-bit. Thus, we report a compression rate of 3.125%; Detailed results are presented in table form in the Appendix E.5.

# **Table 2:** Datasets and models for evaluation. The footprints depict the vanilla models.

Dataset	Model	# GMACs	# Params (M)	Task
MNIST	LeNet	$2e^{-4}$	0.04	Image classification
CIFAR10	ResNet-18	0.56	11.18	Image classification
CIFAR10	VGG-19	0.40	20.00	Image classification
ImageNet	ResNet-50	4.12	25.56	Image classification
Multi30k	6-layer Transformer	1.37	48.98	Translation (en-ge)

Table 3: Maestro vs. baselines on Multi30k.

Variant		Perplexity	GMACs	Params. $(M)$
Non-factorized	Transformer	$9.85_{\pm 0.10}$	1.370	53.90
Pufferfish*	Transformer	$7.34 \pm 0.12$	0.996	26.70
$MAESTRO^{\dagger}$	Transformer	$\textbf{6.90} \scriptstyle{\pm 0.07}$	0.248	13.80

#### 5.1. Experimental Setup

**Models & datasets.** The datasets and models considered in our experiments span across four datasets, concisely presented along with the associated models on Tab. 2. We have implemented our solution in PyTorch (Paszke et al., 2017)(v1.13.0) trained our models on NVidia A100 (40G) GPUs. Details for the learning tasks and hyperparameters used are presented in Appendix D.

**Baselines.** We have selected various baselines from the literature that we believe are closest to aspects of our system. On the *pruning* front, we compare with the IMP (Paul et al., 2023) and RareGems (Sreenivasan et al., 2022) techniques. On the *quantization* front, we compare with XNOR-Net (Rastegari et al., 2016). With respect to *low-rank* methods, we compare with Spectral Initialisation (Khodak et al., 2021), Pufferfish (Wang et al., 2021) and Cuttlefish (Wang et al., 2023).

#### 5.2. Performance Comparison

We start off by comparing MAESTRO with the mentioned baselines from the literature across the datasets and models

of Tab. 2<sup>4</sup>. Results are depicted in Fig. 3 and Tab. 3, while additional performance points of MAESTRO for different model footprints are presented in the Appendix E.3 and E.4.

Comparisons with low-rank methods. The low-rank methods we are comparing against are Pufferfish (Wang et al., 2021) and Cuttlefish (Wang et al., 2023). These methods try to reduce training and inference runtime while preserving model accuracy by leveraging low-rank approximations. For ResNet-18, we achieve  $94.19\pm0.07\%$  for 4.08M parameters and 93.97±0.25% for 2.19M parameters compared to the 94.17% of Pufferfish at 3.3M parameters. For VGG-19, we achieve +0.41pp (percentage points) higher accuracy compared to Pufferfish and -0.29pp to Cuttlefish at 44.8% and 93.2% of the sizes, respectively. Finally, comparing with the spectral initialization (Khodak et al., 2021) for VGG-19, we achieve +5.26pp higher accuracy for 87.5% of parameter size. Detailed results are shown in Tab. 16. This performance benefits also apply in the case of Transformers (Tab. 3), where MAESTRO performs 6% better in terms of perplexity at 25% of the cost (MACs) and 51.7%

<sup>&</sup>lt;sup>4</sup>The operating points we select for MAESTRO are the closest lower to the respective baseline in terms of footprint. Where the result is not present in the Fig. 3, we provide the  $\lambda_{gp}$  value so that it can be referenced from the Appendix, Tab. 12, 13.

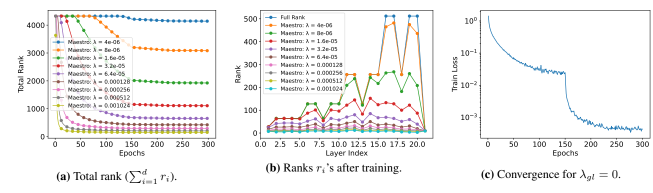


Figure 4: Training dynamics of MAESTRO for ResNet18 on CIFAR10. Results for other datasets can be found in Appendix E.3.

Table 4: Maestro vs. baselines on ImageNet-1k.

Variant	Model	Acc. (%)	Params. $(M)$	GMACs			
No decompositi	on						
Non-factorized	ResNet-50	75.32	25.26	4.12			
Not decomposing first four blocks and last layer							
Pufferfish <sup>†</sup>	ResNet-50	75.99	15.2	3.6			
Cuttlefish <sup>†</sup>	ResNet-50	76.00	14.9	3.6			
MAESTRO *	ResNet-50	76.04	14.0	3.4			
Decomposing al	ll layers						
Pufferfish <sup>†</sup>	ResNet-50	71.03	9.4	2.1			
MAESTRO *	ResNet-50	71.54	9.2	2.0			

<sup>\*</sup> $\lambda_{gl}$  chosen such that the final number of parameters and accuracy is similar to the baseline models; † without label smoothing (same as our setup for Maestro)

of the size (parameters) compared to Pufferfish. It is worth noting that both Pufferfish and Cuttlefish, by default, do not decompose all layers and have warm-up full-training rounds, both of which cause training and hyperparameter optimization overheads. In contrast, our technique only introduces two hyperarameters, namely  $\lambda_{gl}$  and  $\varepsilon_{ps}$ , which govern the whole training process. We have scaled up our experiments to ImageNet-1k levels (Tab. 4) and for the same setup of full decomposition, we achieve slightly higher accuracy (+0.51pp) at 97.8% of the size of Pufferfish. For partial decomposition, MAESTRO performs on par with Pufferfish and Cuttlefish at a lower training and inference cost.

Comparisons with pruning methods. The next family of baselines is related to the LTH (Frankle & Carbin, 2019). Specifically, we compare against IMP (Paul et al., 2023) and witness that MAESTRO can achieve +1.25pp ( $\lambda_{gp}=128e^{-6}$ ) and +0.24pp ( $\lambda_{gp}=32e^{-6}$ ) higher accuracy for ResNet-18 and VGG-19 respectively. The detailed results are shown in Tab. 16 of the Appendix. Although we cannot scale to the size that RareGems (Sreenivasan et al., 2022) for ResNet-18, the sparsity that they achieve is unstructured, which most modern hardware cannot take advantage of. In contrast, our technique performs ordered structured sparsity compatibly with most computation targets. On the other hand, for VGG-19, we achieve +6.82pp higher accuracy at 43.6% of the footprint.

**Table 5:** Ablation study for ResNet18 on CIFAR10

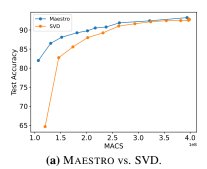
Variant	Acc. (%) Rel	l. GMACs (Trai	n.) Params. $(M)$
MAESTRO	94.19±0.39	1.00×	4.08±0.020
w/out GL	$94.04_{\pm0.10}$	1.33×	$11.2 \pm 0.000$
w/out PS	$94.12 \pm 0.36$	1.33×	$4.09{\scriptstyle\pm0.027}$
w/ full-trainir	<b>ng</b> 94.05±0.32	1.97×	$4.09{\scriptstyle\pm0.032}$

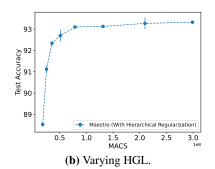
Comparisons with quantized models. We also compare against XNOR-Net (Rastegari et al., 2016), which binarizes the network to achieve efficient inference. Training continues to happen in full precision, and inference performance is dependent on the operation implementation of the target hardware. Nonetheless, assuming a compression rate of 3.125%, for the same model size, we achieve +1.08pp ( $\lambda_{gp}=512e^{-6}$ ) and +2.18pp ( $\lambda_{gp}=256e^{-6}$ ) higher accuracy on ResNet-18 and VGG-19.

#### **5.3. Training Behaviour of MAESTRO**

Having shown the relative performance of our framework to selected baselines, we now move to investigate how our method behaves with respect to its convergence and lowrank approximations.

Model and rank convergence. In Fig. 4, we present the training dynamics for MAESTRO. Fig. 4a illustrates the evolution of total rank throughout the training steps. We observe that the ranks are pruned incrementally. This aligns with the observations made during Pufferfish (Wang et al., 2021) training, where the authors suggest warm-start training with full precision to enhance the final model performance. In our situation, we do not need to integrate this heuristic because MAESTRO automatically prunes rank. Fig. 4b reveals the ranks across layers after training. We notice an intriguing phenomenon: the ranks are nested for increasing  $\lambda_{al}$ . This could imply apart from a natural order of ranks within each layer, a global order. We briefly examine this captivating occurrence in the following section, and we plan to investigate it more thoroughly in future work, as we believe this might contribute to a superior rank selection and sampling process. Lastly, Fig. 4c depicts the progression of training loss. We find that our hypothesis that sampling does not adversely impact training is also supported empirically.





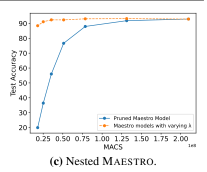


Figure 5: MAESTRO accuracy-latency trade-off under different settings for VGG19 on CIFAR10. Additional results in Appendix E.4.

#### 5.4. Ablation Study

In this section, we examine the impact of each component on the performance of MAESTRO. Specifically, we run variants of our method i) without the hierarchical group lasso regularization (HGL), ii) without progressive shrinking (PS). Additionally, we integrate iii) an extra full low-rank pass ( $b=r_i$ ) into the training at each step to assess whether extra sampling would be beneficial. The results are displayed in Tab. 5. As anticipated, our findings confirm that neither the inclusion of hierarchical group lasso with a tuned  $\lambda_{gl}$  nor progressive shrinking impair the final performance, but they do significantly enhance the efficiency of MAESTRO. Moreover, sampling more ranks at each training step does not improve the final performance, and, in fact, it hampers training efficiency, making it approximately twice as computationally demanding.

# 5.5. Accuracy-Latency Trade-Off at Training and Deployment Time

In Fig. 5, we illustrate various approaches to balance latency (proxied through MACs operations) and accuracy in model training and deployment. Fig. 5a demonstrates how MAE-STRO ( $\lambda_{ql} = 0$ ) can be pruned effectively for deployment using the greedy search method discussed in Section 3.4. We contrast this with the greedy pruning of a non-factorized model that has been factorized using SVD. We reveal that this straightforward baseline does not measure up to the learned decomposition of MAESTRO and results in a significant performance decrease. Next, Fig. 5b portrays the final accuracy and the number of model parameters for varying hierarchical group lasso penalties. This leads to the optimal latency-accuracy balance for both training and inference. However, it is crucial to point out that each model was trained individually, while greedy pruning only necessitates a single training cycle. Lastly, we delve into the observation of nested ranks across increasing  $\lambda_{gl}$ . Fig. 5c displays the performance of MAESTRO ( $\lambda_{gl}=0$ ) across different ranks selected by smaller models MAESTRO ( $\lambda_{ql} > 0$ ). Intriguingly, we observe that MAESTRO ( $\lambda_{al} = 0$ ) performs very well—for instance, we can decrease its operations in half (and parameters by  $10\times$ ) and still maintain an accuracy of

87.7% without fine-tuning, just by reusing rank structure from independent runs. As aforementioned, we intend to further explore this in the future.

#### 6. Conclusion and Future Work

In this work, we have presented MAESTRO, a method for trainable low-rank approximation of DNNs that leverages progressive shrinking by applying a generalized variant of Ordered Dropout to the factorized weights. We have shown the theoretical guarantees of our work in the case of linear models and empirically demonstrated its performance across different types of models, datasets, and modalities. Our evaluation has demonstrated that MAESTRO outperforms competitive compression methods at a lower cost. In the future, we plan to expand our technique to encompass more advanced sampling techniques and apply it to different distributed learning scenarios, such as Federated Learning, where data are natively non-independent or identically distributed (non-IID).

#### **Impact Statement**

The goal of our work is to make the training and deployment of DNNs more efficient, affecting the total computation, memory and bandwidth of systems, as well as the energy they require to run the respective tasks. DNN model training requires significant amounts of energy, whether in a data center or at the edge (Wu et al., 2022; Patterson et al., 2022). However, such techniques should not be used in lieu of making data centers less green, but as a complementary measure to further reduce the carbon footprint of Deep Learning.

Additionally, as our technique involves a training-aware methodology for progressively selecting ranks, it depends on the quality of data used in training. Deploying the model in the wild for various downstream tasks may result in behavior different from the intended outcomes. Therefore, it should be thoroughly tested before deployment to ensure it adheres to the required Service Level Objectives (SLOs), especially in performance-critical use cases, such as self-driving vehicles or UAV navigation.

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# **Appendix**

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#### A. Limitations

In this work, we have proposed a method for trainable low-rank approximation of DNNs that provides performance benefits for both training and inference times. While we suggest that this could have repercussions on the energy consumption of these tasks, we have not yet evaluated this hypothesis experimentally across different devices, be they data center-grade or at the edge.

Additionally, we have applied our technique to CNN and Transformer models spanning across vision and NLP tasks. While we anticipate generalization to any type of network, it remains to be seen whether our techniques can also be applied to alternative types of layers, such as recurrent ones, and the benefits they may bring.

Although we have provided a thorough investigation of the behaviour of our proposed system, the only way we can control the end footprint of the model during training is via the  $\lambda_{gl}$  and  $\varepsilon_{ps}$  hyperparameters. However, there is no guarantee about the final footprint of the model. If we are willing to sacrifise accuracy, then the technique illustrated in Sec. 3.4 and evaluated in Sec. 5.5 is a start. More robust ways of globally ranking per-layer importances are left as future work.

Lastly, our sampling method during training is uniform up to the maximum rank during progressive shrinking. Although this method has proven effective, alternative sampling methods could potentially accelerate rank exploration, thereby hastening the shrinking and convergence of the network during training.

# **B. Extended Background**

**Ordered Dropout.** Ordered Dropout is a technique of importance-based, nested and ordered pruning that works along the indices of a layer's parameters (neurons, filters, etc.) Introduced by (Horváth et al., 2021), the authors describe a training technique where a layer's width is discretised in |P| values, where  $P = \{s_1, s_2, \ldots, s_{|P|}\}$ , and at each training step, they sample  $p \sim U_P$  to get a specific subnetwork, extracted by selecting the first  $\lceil p*K_l-1 \rceil$  neurons per layer and dropping the rest. In contrast to our work, sampling is happening directly on model parameters (rather than ranks) and is uniform across layers (i.e. a single p-value is set). Nested-ness refers to the fact that larger p-value models include the parameters of lower p-values and importance-based pruning means that via stochastic sampling, the right-most (in terms of index) parameters train on progressively less data due to the probability of sampling and nestedness (i.e. all data pass from the parameters of minimal subnetwork, less pass the higher the p-value).

# C. Theoretical Properties of Low-Rank Layers

In this section, we show that for the case of linear mappings, i.e., the problem formulation discussed in (3), MAESTRO acts as PCA applied to the original dataset  $\mathcal{X}$  projected onto the space weighted by the corresponding singular values. Before proceeding with the theorem, we first recall the assumptions and notations introduced in the main paper.

We denote  $C_{:b}$  as the first b columns of matrix C,  $C_{:a,:b}$  denotes the first a rows, and b columns of a matrix C, a+1: denotes the all the columns/rows from index a+1,: denotes the all the columns/rows, and for vectors, we use a single subscript. As discussed in the main paper, we reformulate the original least squares problems to the following decomposition problem

$$\min_{U \in \mathbb{R}^{m \times r}} \mathbf{E}_{x, y \sim \mathcal{X}} \left[ \mathbf{E}_{b \sim \mathcal{D}} \left[ \left\| U_{:b} V_{:b}^{\top} x - y \right\|^{2} \right] \right], \tag{6}$$

where  $\mathcal{D}$  is a distribution that samples  $b \in \{1, 2, \dots, r\}$  with probability  $p_b > 0$  and we assume that y is linked with x through linear map A, i.e., y = Ax.

**Theorem C.1.** Let  $A = \tilde{U}\tilde{\Sigma}\tilde{V}^{\top}$  be a SVD decomposition of A. Then, the minimization problem (6) is equivalent to PCA applied to the transformed dataset  $x \to \tilde{\Sigma}\tilde{V}^{\top}x$ ,  $x \sim \mathcal{X}$  projected on the column space of  $\tilde{U}$ . Concretely, we can first solve

$$\min_{U \in \mathbb{R}^{m \times r}, V \in \mathbb{R}^{n \times r}} \mathbf{E}_{z \sim \mathcal{X}} \left[ \mathbf{E}_{b \sim \mathcal{D}} \left[ \left\| \left( U_{:b} V_{:b}^{\top} - I \right) \tilde{\Sigma} \tilde{V}^{\top} x \right\|^{2} \right] \right], \tag{7}$$

and then we can obtain the solutions of (6) using  $U^* = \tilde{U}^T \bar{U}, V^* = \tilde{V}^T \bar{V}$ , where  $\bar{U}, \bar{V}$  belong to the set of optimal solutions of problem (7).zx

In the particular case, where  $\mathcal{X}$  is a uniform distribution on the unit ball, (6) recovers the best rank approximation of A across all ranks, i.e., up to the scale of U and V recovers truncated SVD. In the case, A is identity, (6) leads to standard PCA decomposition.

*Proof.* From the assumptions that y = Ax and  $A = \tilde{U}\tilde{\Sigma}\tilde{V}^{\top}$ , we can rewrite (6) as

$$\min_{U \in \mathbb{R}^{m \times r}, V \in \mathbb{R}^{n \times r}} \mathbf{E}_{x \sim \mathcal{X}} \left[ \mathbf{E}_{b \sim \mathcal{D}} \left[ \left\| \left( U_{:b} V_{:b}^{\top} - \tilde{U} \tilde{\Sigma} \tilde{V}^{\top} \right) x \right\|^{2} \right] \right].$$

Since  $\tilde{U}$  is orthogonal, we have  $||z|| = ||\tilde{U}^{\top}z||$ . Therefore, the above problem is equivalent to

$$\min_{U \in \mathbb{R}^{m \times r}, V \in \mathbb{R}^{n \times r}} \mathbf{E}_{x \sim \mathcal{X}} \left[ \mathbf{E}_{b \sim \mathcal{D}} \left[ \left\| \left( \tilde{U}^{\top} U_{:b} V_{:b}^{\top} - \tilde{\Sigma} \tilde{V}^{\top} \right) x \right\|^{2} \right] \right],$$

which is also equivalent to

$$\min_{U \in \mathbb{R}^{m \times r}, V \in \mathbb{R}^{n \times r}} \mathbf{E}_{x \sim \mathcal{X}} \left[ \mathbf{E}_{b \sim \mathcal{D}} \left[ \left\| \left( U_{:b} V_{:b}^{\top} - \tilde{\Sigma} \tilde{V}^{\top} \right) x \right\|^{2} \right] \right]$$

after reparametrization. The next step involves injecting identity in the form  $\tilde{V}\tilde{V}^{\top}$  as that leads to the equivalent reformulation

$$\min_{U \in \mathbb{R}^{m \times r}, V \in \mathbb{R}^{n \times r}} \mathbf{E}_{x \sim \mathcal{X}} \left[ \mathbf{E}_{b \sim \mathcal{D}} \left[ \left\| \left( U_{:b} V_{:b}^{\top} \tilde{V} - \tilde{\Sigma} \right) \tilde{V}^{\top} x \right\|^{2} \right] \right].$$

As for the previous case, we can reparametrise the problem to obtain

$$\min_{U \in \mathbb{R}^{m \times r}, V \in \mathbb{R}^{n \times r}} \mathbf{E}_{x \sim \mathcal{X}} \left[ \mathbf{E}_{b \sim \mathcal{D}} \left[ \left\| \left( U_{:b} V_{:b}^{\top} - \tilde{\Sigma} \right) \tilde{V}^{\top} x \right\|^{2} \right] \right].$$

Let  $k = \operatorname{rank}(\tilde{\Sigma}) = \operatorname{rank}(A) \leq r$  and  $z = \tilde{V}^\top x$ . Furthermore, let  $g = \tilde{\Sigma} z$  for any  $z \in \mathbb{R}^n$ , then  $g_{k+1} = \vec{0}$ . This, combined with the nested structure of the optimization problem, implies that the optimal solution for U has to be of the form  $u_{i,k+1} = \vec{0}$  for all interesting (non-zero mapping) directions, i.e., there exists  $x \in \mathcal{X}$  such that  $v_i^\top \tilde{V}^\top x \neq 0$ . These are the only interesting solutions since the case where for all  $x \in \mathcal{X} : v_i^\top \tilde{V}^\top x = 0$  yields zero mapping on  $\mathcal{X}$ , which is not

of interest and could be dropped, e.g., using group lasso penalty discussed in the main part. Therefore, to solve the original problem, we could first solve the following problem

$$\min_{U \in \mathbb{R}^{k \times r}, V \in \mathbb{R}^{n \times r}} \mathbf{E}_{z \sim \mathcal{X}} \left[ \mathbf{E}_{b \sim \mathcal{D}} \left[ \left\| \left( U_{:k,:b} V_{:b}^{\top} - \tilde{\Sigma}_{:k,:} \right) z \right\|^{2} \right] \right]$$

and then reconstruct the corresponding solution of the original problem by appending zeros to the resulting matrix U. By a similar argument, we can argue that for all non-zero mapping directions, it has to be the case that  $v_{i,k+1} = \vec{0}$ . Therefore, solving the original minimization reduces to

$$\min_{U \in \mathbb{R}^{k \times r}, V \in \mathbb{R}^{k \times r}} \mathbf{E}_{z \sim \mathcal{X}} \left[ \mathbf{E}_{b \sim \mathcal{D}} \left[ \left\| \left( U_{:b} V_{:b}^{\top} - \tilde{\Sigma}_{:k,:k} \right) z_{:k} \right\|^{2} \right] \right].$$

This can be further simplified using reparametrization  $V^{\top} \to V^{\top} \tilde{\Sigma}^{-1}_{:k::k}$ , which leads to

$$\min_{U \in \mathbb{R}^{k \times r}, V \in \mathbb{R}^{k \times r}} \mathbf{E}_{z \sim \mathcal{X}} \left[ \mathbf{E}_{b \sim \mathcal{D}} \left[ \left\| \left( U_{:b} V_{:b}^{\top} - I_{k} \right) \tilde{\Sigma}_{:k,:k} z_{:k} \right\|^{2} \right] \right], \tag{8}$$

where  $I_k$  is  $k \times k$  identity. If  $\mathcal{X}$  is centred around zero, then  $\tilde{\Sigma}_{:k,:k}z_{:k}$  is also centred around zero, and the above problem is up to scaling equivalent to PCA of  $\tilde{\Sigma}_{:k,:k}z_{:k}$  as shown by Rippel et al., (Rippel et al., 2014). Since  $\tilde{\Sigma}$  is a diagonal matrix with only  $k \times k$  non-zero upper left sub-matrix, therefore, PCA on  $\tilde{\Sigma}_{:k,:k}z_{:k}$  is equivalent to PCA on  $\tilde{\Sigma}z$  by appending zeros to the obtained principal component vectors. Thus, we can write an equivalent formulation

$$\min_{U \in \mathbb{R}^{m \times r}, V \in \mathbb{R}^{n \times r}} \mathbf{E}_{z \sim \mathcal{X}} \left[ \mathbf{E}_{b \sim \mathcal{D}} \left[ \left\| \left( U_{:b} V_{:b}^{\top} - I \right) \tilde{\Sigma} \tilde{V}^{\top} x \right\|^{2} \right] \right].$$

Furthermore, let  $\bar{U}, \bar{V}$  belong to the set of optimal solutions of problem (7). Then  $U^* = \tilde{U}^\top \bar{U}, V^* = \tilde{V}^\top \bar{V}$  belong to the set of optimal solutions of problem (6). This can be proved by reversing our construction and ignoring scaling since (7) is scaling invariant.

For the case  $\mathcal{X}$  is a uniform distribution on the unit ball, we have  $\tilde{\Sigma}_{:k,:k}z_{:k}$  is a k-dimensional ellipsoid with principal axes being standard basis vectors  $\{e_i\}_{i=1}^k$ , where the length of the axes is given by ordered singular values, i.e., the first basis vector corresponds to the largest singular vector. Therefore, its principal component vectors correspond to the basis vectors. Following our construction, one can see that the solution to the original problems leads to truncated SVD up to the scaling factor.

For the case A is an identity, we have  $k=r=m=m, \tilde{\Sigma}$  is an identity, and  $\tilde{U}=\tilde{V}$ . Under this setting, the principal component vectors obtained from (8) corresponds to principal component vectors of  $\mathcal{X}$  in basis given by columns of  $\tilde{U}$ . Similarly, as in the previous case, reversing the transformations to return back to the original problem, we conclude that the optimal solution of the original problem corresponds to principal component vectors of  $\mathcal{X}$  since we reverse the transformation by  $\tilde{U}^{\top}$ .

## D. Experimental setup

#### D.1. Datasets

**MNIST.** The MNIST dataset (LeCun et al., 2010) is a database of  $28 \times 28$  greyscale handwritten digits, with a training set of 60k examples and a test set of 10k samples.

CIFAR-10. The CIFAR10 dataset (Krizhevsky et al., 2009) is a computer vision dataset that consists of  $32 \times 32$  RGB images classified into 10 labels. It is split into 50k training images and 10k test images which are balanced across labels.

**ImageNet-1k.** The ImageNet dataset (ILSVRC) (Deng et al., 2009) is an image classification challenge. The task comprises to classify an 300×300 RGB image among 1000 classes. In total there are 1.2M training samples and 50k test images.

**WMT16.** The WMT dataset from statmt is machine translation dataset, spanning news commentaries and parliament proceedings, that aims to investigate the applicability of machine translation techniques when translating between language pairs. Specifically, we focus on the task of German-English language translation of image descriptions, commonly referred to as **Multi30k** (Elliott et al., 2016). We only utilise the text modality for the translation task. Data is taken straight from torchtext.

#### D.2. Models

**LeNet.** LeNet is a simple convolutional network, introduced by LeCun at al. for recognizing handwritten digits (LeCun et al., 2010). It consists of a sequence of two convolutional layers, followed by three fully-connected layers. However, we are using a ReLU instead of the initially proposed sigmoid activation. The detailed architecture of the network is depicted in Tab. 6

**ResNet.** ResNet (He et al., 2016) is a deep neural network whose prominent feature is the existence of skip (or residual) connections, that is connections that perform identity mappings merged with the target layer it joins with through summation. Multiple residual blocks are stacked to form the network. The result is an easier to optimise network that offers enhanced accuracy. We use ResNet-18 in our experiments, the architecture of which is depicted in Tab. 7, except for ImageNet, where we use ResNet-50.

**Table 6:** Detailed architecture of the LeNet-5 architecture used in our experiments. Each convolution and linear layer is followed by a ReLU activation that is ommitted from the table. The shapes for convolution layers follows (m, n, k, k).

Parameter	Shape	Layer hyper-parameter
layer1.conv1.weight	$1\times 6\times 5\times 5$	stride:1;padding:1
pooling.max	N/A	kernel size:2;stride:1;dilation:1
layer2.conv2.weight	$6\times16\times5\times5$	stride:1;padding:0;dilation:1
pooling.max	N/A	kernel size:2;stride:2
layer3.fc1.weight	$256\times120$	N/A
layer4.fc2.weight	$120 \times 84$	N/A
layer5.fc3.weight	$84 \times 10$	N/A

Layer Name ResNet-18 ResNet-50 3×3, 64, stride 1, padding 1 7×7, 64, stride 2, padding 1 conv1 3×3 maxpool, stride 2  $1 \times 1,64$ conv2\_x  $3 \times 3,64$  $3 \times 3,64$  $3 \times 3,64$  $1 \times 1, 256$  $1 \times 1, 128$  $3 \times 3, 128 \\ 3 \times 3, 128$ conv3\_x  $3 \times 3, 128$   $1 \times 1, 512$  $1 \times 1,256 \\ 3 \times 3,256$  $3 \times 3, 256 \\ 3 \times 3, 256$ conv4 x  $\times 6$  $1 \times 1, 1024$  $1 \times 1,512$  $3 \times 3,512$  $1 \times 1,2048$ conv5 x  $\times 3$ 

Avg Pool, 20-dim FC, SoftMax

Avg Pool, 10-dim FC, SoftMax

Table 7: The hybrid ResNet architecture for the CIFAR-10 and

ImageNet datasets used in the experiments.

**VGG.** VGG (Simonyan & Zisserman, 2015) is a also a convolutional network that leverages smaller 3×3 convolutions that enables deeper architecture than before. For our experiments we are using VGG-19, the architecture of which is depicted in Tab. 8.

**Table 8:** Detailed architecture of the VGG-19 architecture used in our experiments. There is a BatchNorm layer followed by a ReLU activation (omitted in the table) after each convolution layer. The shapes for convolution layers follows (m, n, k, k).

Parameter	Shape	Layer hyper-parameter
layer1.conv1.weight	$3 \times 64 \times 3 \times 3$	stride:1;padding:1
layer2.conv2.weight	$64 \times 64 \times 3 \times 3$	stride:1;padding:1
pooling.max	N/A	kernel size:2;stride:2
layer3.conv3.weight	$64 \times 128 \times 3 \times 3$	stride:1;padding:1
layer4.conv4.weight	$128\times128\times3\times3$	stride:1;padding:1
pooling.max	N/A	kernel size:2;stride:2
layer5.conv5.weight	$128\times256\times3\times3$	stride:1;padding:1
layer6.conv6.weight	$256\times256\times3\times3$	stride:1;padding:1
layer7.conv7.weight	$256\times256\times3\times3$	stride:1;padding:1
layer8.conv8.weight	$256\times256\times3\times3$	stride:1;padding:1
pooling.max	N/A	kernel size:2;stride:2
layer9.conv9.weight	$256\times512\times3\times3$	stride:1;padding:1
layer10.conv10.weight	$512\times512\times3\times3$	stride:1;padding:1
layer11.conv11.weight	$512\times512\times3\times3$	stride:1;padding:1
layer12.conv12.weight	$512\times512\times3\times3$	stride:1;padding:1
pooling.max	N/A	kernel size:2;stride:2
layer13.conv13.weight	$512\times512\times3\times3$	stride:1;padding:1
layer14.conv14.weight	$512\times512\times3\times3$	stride:1;padding:1
layer15.conv15.weight	$512\times512\times3\times3$	stride:1;padding:1
layer16.conv16.weight	$512\times512\times3\times3$	stride:1;padding:1
pooling.avg	N/A	kernel size:2
classifier.weight	$512 \times 10$	N/A
classifier.bias	10	N/A

**Transformers.** The transformer architecture (Vaswani et al., 2017) has been lately revolutionising deep learning. Based on the notion of self-attention, for each input token, it produces a weighted combination of other relevant tokens weighed by the attention weight. Each attention unit has three weight matrices, namely  $W_Q$ ,  $W_K$ ,  $W_V$ , for query, key and value weights respectively producing the equivalent vectors. Attention is defined as the scaled dot product between key and query. For our translation task, we use the architecture depicted in Tab. 10.

**Table 10:** Detailed information of the decoder layer in the Transformer architecture in our experiment

**Table 9:** Detailed information of the encoder layer in the Transformer architecture in our experiment

Parameter	Shape	Hyper-param.
embedding	$9521 \times 512$	padding index: 1
positional encoding	N/A	N/A
dropout	N/A	p = 0.1
${\bf encoder.self\text{-}attention.wq}(W^Q)$	$512\times512$	N/A
${\bf encoder.self\text{-}attention.wk}(W^K)$	$512\times512$	N/A
${\bf encoder.self\text{-}attention.wv}(\boldsymbol{W}^{V})$	$512\times512$	N/A
${\bf encoder.self\text{-}attention.wo}(W^O)$	$512\times512$	N/A
encoder.self-attention.dropout	N/A	p = 0.1
encoder.self-attention.layernorm	512	$\varepsilon = 10^{-6}$
encoder.ffn.layer1	$512\times2048$	N/A
encoder.ffn.layer2	$2048\times512$	N/A
encoder.layernorm	512	$\varepsilon = 10^{-6}$
dropout	N/A	p = 0.1

Parameter	Shape	Hyper-param.
embedding	$9521 \times 512$	padding index: 1
positional encoding	N/A	N/A
dropout	N/A	p = 0.1
${\bf decoder.self\text{-}attention.wq}(W^Q)$	$512\times512$	N/A
${\bf decoder.self\text{-}attention.wk}(W^K)$	$512\times512$	N/A
${\bf decoder.self\text{-}attention.wv}(W^V)$	$512\times512$	N/A
${\bf decoder.self\text{-}attention.wo}(W^O)$	$512\times512$	N/A
decoder.self-attention.dropout	N/A	p = 0.1
decoder.self-attention.layernorm	512	$\varepsilon=10^{-6}$
${\bf decoder.enc\text{-}attention.wq}(W^Q)$	$512\times512$	N/A
${\bf decoder.enc\text{-}attention.wk}(\boldsymbol{W}^K)$	$512\times512$	N/A
${\bf decoder.enc\text{-}attention.wv}(W^V)$	$512\times512$	N/A
${\bf decoder.enc\text{-}attention.wo}(W^O)$	$512\times512$	N/A
decoder.enc-attention.dropout	N/A	p = 0.1
decoder.enc-attention.layernorm	512	$\varepsilon = 10^{-6}$
decoder.ffn.layer1	$512\times2048$	N/A
decoder.ffn.layer2	$2048\times512$	N/A
encoder.layernorm	512	$\varepsilon = 10^{-6}$
dropout	N/A	p = 0.1

# **D.3. Hyperparameter Selection**

**LeNet.** We use a standard configuration that is commonly employed for training LeNet models — a step size of 0.01, a momentum of 0.9, and no weight decay. We train for a total of 20 epochs.

**VGG and ResNet-18.** Similarly, we use a standard configuration that is commonly employed for training VGG and ResNet-18 models — a step size of 0.01, a momentum of 0.9, weight decay of  $1e^{-4}$ , and a learning schedule with step size reductions by a factor of 10 at epochs 150 and 250. We train for a total of 300 epochs.

**ResNet-50.** Similarly, we use a standard configuration that is commonly employed for training ResNet-50 models — a step size of 0.01, a momentum of 0.9, weight decay of  $1e^{-4}$ , and a learning schedule with step size reductions by a factor of 10 at epochs 30 and 60. We train for a total of 90 epochs.

**Transformers.** For the Transformer model, we use the Adam optimizer with an initial learning rate at 0.001,  $\beta s = (0.9, 0.98)$ ,  $\varepsilon = 10^{-8}$  batch size at 256. We also conduct gradient norm clipping with norm bound at 0.25. The entire training takes 400 epochs. For the vanilla warm-up training, we use warm-up epoch  $E_{wu} = 10$ . We enable label smoothing, weight sharing for the source and target word embedding, and weight sharing between target word embedding and the last dense layer. The learning rate schedule follows directly from the one proposed (Vaswani et al., 2017).

#### **D.4. Deciding Against Decomposition**

During inference, if the rank of a given layer is so large that keeping it as a non-decomposed layer is more efficient, we opt not to decompose that particular layer.

# E. Extended evaluation

#### E.1. MAESTRO Recovers Correct Ordering

In the main text, we pointed out that SVD fails to consider data. Indeed, even in the case of linear NN, the acquired singular vectors may exhibit incorrect ordering. To illustrate this problem, we provide a simple example in which we use a matrix A

with a rank of 3. We organize the dataset  $\mathcal{X}$  such that the third singular vector has the highest importance, followed by the second and then the first singular vector in decreasing order of significance. It is clear that SVD doesn't consider the data, and as a result, it cannot comprehend this behavior. Below (in Fig. 6), we demonstrate how MAESTRO is able to correctly discern the order.

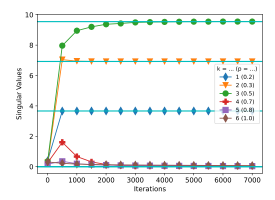
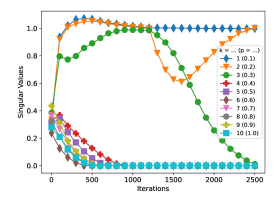


Figure 6: Verification that MAESTRO recovers correct order of importance. Target mapping is of rank 3, and the dataset is constructed in such a way that the singular vectors have reversed the order of importance. p and k stand for relative and actual rank, respectively.

#### E.2. Rank Adaptivity of MAESTRO to Data Complexity

So far, we have found that different models can have different ranks on different datasets. However, we did not reach the conclusion that more complex tasks lead to higher ranks because the model architecture is not invariant, i.e., we cannot compare ranks between layers of different dimensionality.



**Figure 7:** MAESTRO adaptivity when PCA dimensionality drops during training. The plot displays the estimates of singular values. The data distribution has initially 3 directions. It is expected that the top 3 ranks will converge to value one and the rest to zero. After removing one direction, ranks drop to 2, as the data complexity changes. *p* and *k* stand for relative and actual rank, respectively.

To test this hypothesis in silo, we designed a simplified experiment on a linear autoencoder example with the same setup as considered in Fig. 2b). To showcase the adaptivity of MAESTRO to changing data, we start by training with data that have intrinsic dimension 3. In the middle of the training (iteration 1250/2500), we removed one dimension using projection, thus simplifying the data. In the resulting graph (Fig. 7), we see that while the initial rank had converged to 3, it now drops to 2 as the data complexity changes. For completeness, this adaptivity is further showcased in Appendix E.1, where we have illustrated how SVD fails to consider data-centric factors, whereas Maestro recovers the correct order of importance.

#### E.3. Training Behaviour of MAESTRO

For completeness, we also include an extended version of Fig. 4 from the main paper, where we presented the training dynamics for MAESTRO. Fig. 8, 9 and 10 present similar plots, but across both MNIST and CIFAR-10. Specifically, Fig. 8 illustrates the evolution of total rank throughout the training steps. We observe that the ranks are pruned incrementally. This

aligns with the observations made during Pufferfish (Wang et al., 2021) training, where the authors suggested warm-start training with full precision to enhance the final model performance. In our case, the necessity to implement this heuristic is avoided, as MAESTRO prunes rank automatically. Fig. 9 demonstrates the ranks across layers post-training. An intriguing trend is observed: the ranks are nested for increasing  $\lambda_{gl}$ , suggesting a potential inherent ordering of ranks not only within each layer but also possibly a global one. We provide a preliminary exploration of this fascinating pattern in the subsequent section and intend to probe it more deeply in future studies. We believe this may enhance the rank selection and sampling process. Finally, Fig. 10 portrays the evolution of the training loss. Our premise that sampling does not negatively affect training is validated by empirical performance.

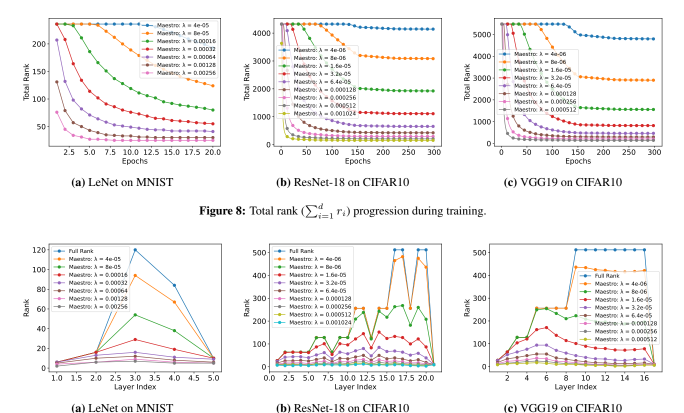


Figure 9: Ranks  $r_i$ 's across different layers after training.

#### E.4. Model Size-Accuracy Trade-Off at Training and Deployment Time

In addition to the original illustrations, we present an extended interpretation of Fig. 5, where we depict diverse strategies to maintain a balance between model size and accuracy in the process of model training and deployment. In Fig. 11, we demonstrate the effective pruning of MAESTRO ( $\lambda_{gl}=0$ ) for deployment, utilizing the greedy search methodology discussed in Section 3.4. This is juxtaposed with the greedy pruning of a model not originally factorized but later factorized through SVD. Our findings reveal that this straightforward baseline does not match the performance of MAESTRO's learned decomposition, leading to a considerable performance drop.

Subsequently, Fig. 12 displays the end accuracy and the count of model parameters corresponding to various hierarchical group lasso penalties. This results in an optimal compromise between latency and accuracy for both the training and inference stages. It's worth noting, though, that each model was trained separately, in contrast to greedy pruning, which demands just a single training round. Additionally, we scrutinize the training expense for each model illustrated in Fig. 12, the results of which are exhibited in Tables 11, 12, 13, 14 and 15, where we display and the final accuracy of the model, MACs and the number of parameters for inference, and relative total training cost in terms of the number of model parameters and MACs compared to the non-factorized model. Interestingly, smaller models are not only advantageous in terms of inference efficiency, but they can also be trained at a small portion of the cost required by full-rank models. On the downside, the smallest models cause a non-negligible reduction in performance.

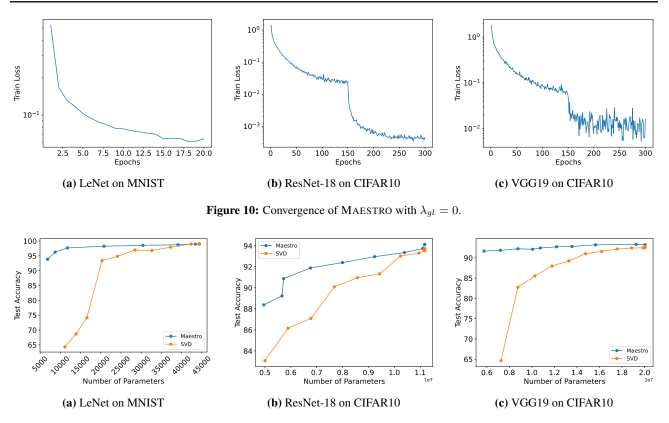


Figure 11: Accuracy-latency trade-off comparing MAESTRO ( $\lambda_{gl=0}$ ) and SVD.

Lastly, we delve deeper into the observation of nested ranks with increasing  $\lambda_{gl}$ . Fig. 13 outlines the performance of MAESTRO ( $\lambda_{gl}=0$ ) across various ranks chosen by smaller models MAESTRO ( $\lambda_{gl}>0$ ). We observe that MAESTRO ( $\lambda_{gl}=0$ ) delivers impressive results—for example, we can reduce its parameters by 10x for VGG while preserving an accuracy of 87.7% without any fine-tuning simply by leveraging rank structure from separate runs. For LeNet, a reduction in model size by a factor of three is achievable without sacrificing accuracy. Last, for ResNet-18 the reduction is  $1.7\times$ . As highlighted earlier, we aim to delve deeper into this subject in future studies.

#### E.5. Detailed Comparison with Baselines

Tab. 16 presents the details of MAESTRO's performance compared to the selected baselines leveraging pruning, quantization and low-rank techniques presented in Sec. 5.2 for CIFAR-10. These numbers along with the operating points from Tab. 12 and 13 are illustrated in Fig. 3.

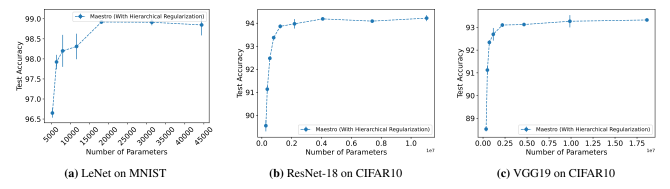


Figure 12: Impact of hierarchical group lasso on the accuracy-memory trade-off. Exact values are provided in Tables 11, 12 and 13, respectively.

**Table 11:** LeNet performance on MNIST for different regularization parameters. The last column in the table displays the relative total training cost in terms of the number of Multiply-Accumulate operations (MACs) and model parameters, compared to the non-factorized model

Variant	Acc. (%)	MACs (Inf.)	Params. (Inf.)	Rel. MACs / Params. (Train.)
Non-Factorized	98.99 <sub>±0.09</sub>	281640±0 (1.00×)	44426±0 (1.00×)	1.00× / 1.00×
Maestro ( $\lambda_{gp} = 0$ .)	$99.06 \pm 0.09$	$281640 \pm 0 \ (1.00 \times)$	$44426\pm0(1.00\times)$	$1.14\times/1.49\times$
MAESTRO ( $\lambda_{qp} = 8e^{-5}$ )	$98.91{\scriptstyle\pm0.09}$	$268577 \pm 389 (0.95 \times)$	$31363\pm0 (0.71\times)$	$1.08 \times / 1.14 \times$
MAESTRO ( $\lambda_{gp} = 16e^{-5}$ )	$98.92 \scriptstyle{\pm 0.05}$	255369±217 (0.91×)	44426±217 (0.41×)	$1.06 \times /~0.80 \times$
Maestro ( $\lambda_{gp} = 32e^{-5}$ )	$98.31 \pm 0.39$	$237084 \pm 6268 \ (0.84 \times)$	$18155\pm271~(0.26\times)$	$0.93\times$ / $0.53\times$
Maestro ( $\lambda_{gp} = 64e^{-5}$ )	$98.20_{\pm 0.49}$	$178165 \pm 19098 \ (0.63 \times)$	$7996_{\pm 662} (0.18 \times)$	$0.77 \times / 0.33 \times$
MAESTRO ( $\lambda_{gp} = 128e^{-5}$ )	$97.92{\scriptstyle\pm0.22}$	$131789 \pm 8965 (0.47 \times)$	$6375_{\pm 77} (0.14 \times)$	$0.54\times$ / $0.21\times$
Maestro ( $\lambda_{gp} = 256e^{-5}$ )	$96.65{\scriptstyle\pm0.14}$	$99969 {\scriptstyle \pm 6252}  (0.35 \times)$	$5293\pm_{214}(0.12\times)$	$0.39\times/0.14\times$

**Table 12:** ResNet-18 performance on CIFAR10 for different regularization parameters. The last column in the table displays the relative total training cost in terms of the number of Multiply-Accumulate operations (MACs) and model parameters, compared to the non-factorized model.

Variant	Acc. (%)	GMACs (Inf.)	Params. (M) (Inf.)	Rel. MACs / Params. (Train.)
Non-Factorized	93.86±0.20	$0.56_{\pm 0} (1.00 \times)$	$11.2_{\pm 0} (1.00 \times)$	1.00× / 1.00×
Maestro ( $\lambda_{gp} = 0.$ )	$94.04 \pm 0.10$	$0.56_{\pm 0} (1.00 \times)$	$11.2_{\pm 0} (1.00 \times)$	$1.10 \times / 1.13 \times$
MAESTRO ( $\lambda_{gp} = 4e^{-6}$ )	$94.22{\scriptstyle\pm0.16}$	$0.55 \pm 0.0047 \ (1.00 \times)$	$11.1 \pm 0.030 \ (0.99 \times)$	$1.09 \times$ / $1.10 \times$
MAESTRO ( $\lambda_{qp} = 8e^{-6}$ )	$94.09_{\pm 0.01}$	$0.49_{\pm 0.0002} (0.89 \times)$	$7.41_{\pm 0.004} (0.66 \times)$	$1.00 \times$ / $0.85 \times$
MAESTRO ( $\lambda_{gp} = 16e^{-6}$ )	$94.19{\scriptstyle\pm0.07}$	$0.39 \pm 0.0008 \ (0.70 \times)$	$4.08 \pm 0.020 \ (0.37 \times)$	$0.83 \times$ / $0.58 \times$
MAESTRO ( $\lambda_{qp} = 32e^{-6}$ )	$93.97 \pm 0.25$	$0.25 \pm 0.0013 \ (0.45 \times)$	$2.19\pm0.007$ (0.20×)	$0.60 \times$ / $0.36 \times$
MAESTRO ( $\lambda_{gp} = 64e^{-6}$ )	$93.86 \pm 0.11$	$0.15\pm0.0006$ $(0.27\times)$	$1.23\pm0.004~(0.11\times)$	$0.39 \times$ / $0.22 \times$
MAESTRO ( $\lambda_{qp} = 128e^{-6}$ )	$93.37 \pm 0.07$	$0.094 \pm 0.0006 \ (0.17 \times)$	$0.79 \pm 0.009 (0.07 \times)$	$0.25 \times$ / $0.13 \times$
MAESTRO ( $\lambda_{gp} = 256e^{-6}$ )	$92.48{\scriptstyle\pm0.04}$	$0.064 \pm 0.0002 \ (0.12 \times)$	$0.54 \pm 0.006 \ (0.05 \times)$	$0.16 \times$ / $0.08 \times$
Maestro ( $\lambda_{gp} = 512e^{-6}$ )	$91.14{\scriptstyle\pm0.16}$	$0.044_{\pm 0.0004}  (0.08 \times)$	$0.37_{\pm 0.007} (0.03 \times)$	$0.11 \times$ / $0.05 \times$
Maestro ( $\lambda_{gp}^{37} = 1024e^{-6}$ )	$89.55{\scriptstyle\pm0.30}$	$0.032 \pm 0.0002 \ (0.06 \times)$	$0.27_{\pm 0.007} (0.02 \times)$	0.07  imes / $0.03  imes$

**Table 13:** VGG19 performance on CIFAR10 for different regularization parameters. The last column in the table displays the relative total training cost in terms of the number of Multiply-Accumulate operations (MACs) and model parameters, compared to the non-factorized model.

Variant	Acc. (%)	GMACs (Inf.)	Params. (M) (Inf.)	Rel. MACs / Params. (Train.)
Non-Factorized	$92.94_{\pm 0.17}$	$0.40_{\pm 0} (1.00 \times)$	$20_{\pm 0} (1.00 \times)$	1.00× / 1.00×
Maestro ( $\lambda_{qp} = 0$ .)	$93.06{\scriptstyle\pm0.17}$	$0.40_{\pm 0}  (1.00 \times)$	$20_{\pm 0} (1.00 \times)$	$1.10 \times$ / $1.12 \times$
MAESTRO ( $\lambda_{gp} = 4e^{-6}$ )	$93.33{\scriptstyle\pm0.08}$	$0.39 \pm 0.0017 \ (0.97 \times)$	$18.8_{\pm0} (0.94 \times)$	$1.06 \times$ / $1.04 \times$
MAESTRO ( $\lambda_{gp} = 8e^{-6}$ )	$93.27{\scriptstyle\pm0.33}$	$0.30 \pm 0.0017 \ (0.76 \times)$	$9.91_{\pm 0.008} (0.49 \times)$	$0.90 \times$ / $0.73 \times$
MAESTRO ( $\lambda_{gp} = 16e^{-6}$ )	$93.13{\scriptstyle\pm0.07}$	$0.21 \pm 0.0014 \ (0.53 \times)$	$4.66 \pm 0.052 \ (0.23 \times)$	$0.69 \times$ / $0.46 \times$
Maestro ( $\lambda_{gp} = 32e^{-6}$ )	$93.10{\scriptstyle\pm0.10}$	$0.13\pm_{0.0009} (0.33\times)$	$2.20_{\pm 0.025} (0.11 \times)$	$0.47 \times$ / $0.27 \times$
MAESTRO ( $\lambda_{gp} = 64e^{-6}$ )	$92.70{\scriptstyle\pm0.34}$	$0.08 \pm 0.0005 \ (0.20 \times)$	$1.17 \pm 0.010 \ (0.06 \times)$	$0.30 \times$ / $0.16 \times$
MAESTRO ( $\lambda_{gp} = 128e^{-6}$ )	$92.34{\scriptstyle\pm0.12}$	$0.05 \pm 0.0005 (0.13 \times)$	$0.72 \pm 0.002 \ (0.04 \times)$	$0.19 \times$ / $0.09 \times$
MAESTRO ( $\lambda_{qp} = 256e^{-6}$ )	$91.12{\scriptstyle\pm0.19}$	$0.04 \pm 0.0007 \ (0.09 \times)$	$0.50 \pm 0.023 \ (0.02 \times)$	$0.12 \times$ / $0.05 \times$
Maestro ( $\lambda_{gp} = 512e^{-6}$ )	$88.53{\scriptstyle\pm0.13}$	$0.03_{\pm 0.0003} \ (0.06 \times)$	$0.35 \pm 0.003 \ (0.02 \times)$	$0.08 \times$ / $0.03 \times$

**Table 14:** Transformer performance on Multi30k for different regularization parameters. The last column in the table displays the relative total training cost in terms of the number of Multiply-Accumulate operations (MACs) and model parameters, compared to the non-factorized model.

Variant	Acc. (%)	Ppl.	GMACs (Inf.)	Params. (M) (Inf.)	Rel. MACs / Params. (Train.)
Non-Factorized	65.33±1.13	$9.85_{\pm 0.10}$	1.370±0.0000 (1.00×)	53.9±0.000 (1.00×)	1.00× / 1.00×
Maestro ( $\lambda_{qp} = 0.32$ )	$61.30 \pm 0.26$	$12.99 \pm 0.31$	$1.125 \pm 0.0030 \ (0.82 \times)$	$45.1_{\pm 0.101} (0.84 \times)$	$1.03 \times$ / $1.14 \times$
MAESTRO ( $\lambda_{qp} = 0.64$ )	$63.78 \scriptstyle{\pm 0.14}$	$9.37{\scriptstyle\pm0.32}$	$0.957_{\pm 0.0112}  (0.70 \times)$	$39.1_{\pm 0.413} (0.73 \times)$	$0.95 \times$ / $1.05 \times$
MAESTRO ( $\lambda_{gp} = 1.28$ )	$66.14{\scriptstyle\pm0.08}$	$7.02{\scriptstyle\pm0.17}$	$0.570_{\pm 0.0088} (0.42 \times)$	$25.3_{\pm 0.315} (0.47 \times)$	$0.75 \times$ / $0.86 \times$
MAESTRO ( $\lambda_{gp} = 2.56$ )	$66.08 \scriptstyle{\pm 0.09}$	$6.90{\scriptstyle \pm 0.07}$	$0.248 \pm 0.0032 \ (0.18 \times)$	$13.8 \pm 0.113 \ (0.26 \times)$	$0.47 \times$ / $0.58 \times$
Maestro ( $\lambda_{gp} = 5.12$ )	$57.70{\scriptstyle\pm0.13}$	$13.97{\scriptstyle\pm0.43}$	$0.123_{\pm 0.0002} (0.9 \times)$	$9.3_{\pm 0.001} (0.17 \times)$	0.28  imes / $0.39  imes$

**Table 15:** ResNet50 performance on ImageNet-1k for different regularization parameters. The last column in the table displays the relative total training cost in terms of the number of Multiply-Accumulate operations (MACs) and model parameters, compared to the non-factorized model.

Variant	Acc. (%)	GMACs (Inf.)	Params. (M) (Inf.)	Rel. MACs / Params. (Train)			
No decomposition							
Non-Factorized	76.00	4.12 (1.00×)	25.56 (1.00×)	1.00  imes / $1.00  imes$			
Not decomposing first four blocks and last layer							
MAESTRO ( $\lambda_{gp} = 2e^{-6}$ )	76.04	$3.43 (0.83 \times)$	$14.02 (0.55 \times)$	$0.87 \times$ / $0.64 \times$			
MAESTRO ( $\lambda_{gp} = 4e^{-6}$ )	75.74	$3.39(0.82\times)$	$13.11 (0.51 \times)$	$0.85 \times$ / $0.59 \times$			
Maestro ( $\lambda_{gp} = 8e^{-6}$ )	75.15	3.21 (0.78×)	11.46 (0.45×)	$0.83 \times$ / $0.55 \times$			
Decomposing all layers							
Maestro ( $\lambda_{gp} = 0$ .)	72.82	$4.12(1.00\times)$	$25.56 (1.00 \times)$	$1.22 \times$ / $1.24 \times$			
MAESTRO ( $\lambda_{gp} = 1e^{-6}$ )	72.81	$3.62(0.88\times)$	$18.77 (0.73 \times)$	$1.00 \times$ / $0.87 \times$			
MAESTRO ( $\lambda_{gp} = 2e^{-6}$ )	72.07	$2.66 (0.65 \times)$	$11.54 (0.45 \times)$	$0.76 \times$ / $0.59 \times$			
MAESTRO ( $\lambda_{gp} = 4e^{-6}$ )	71.54	$2.01~(0.49\times)$	9.21 (0.36×)	$0.57 \times$ / $0.57 \times$			
MAESTRO ( $\lambda_{gp} = 8e^{-6}$ )	71.02	1.69 (0.41×)	7.21 (0.28×)	0.50  imes / $0.39  imes$			

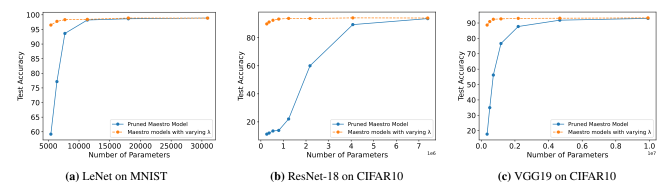


Figure 13: MAESTRO with progressive pruning to showcase nested rank importance structure. The original model corresponds to an evaluation in Fig. 12, and pruned models are based on MAESTRO with  $\lambda_{gl}=0$ , and they are pruned using the same ranks as selected by MAESTRO with  $\lambda_{gl}>0$ .

Table 16: Maestro vs. baselines on CIFAR10.

Variant	Model	Acc. (%)	GMACs	Params. $(M)$
Non-factorized	ResNet-18	$93.86 \pm 0.20$	0.56	11.17
Pufferfish	ResNet-18	94.17	0.22	3.336
Cuttlefish	ResNet-18	93.47	0.3	3.108
IMP	ResNet-18	92.12	-	0.154
RareGems	ResNet-18	92.83	-	0.076
XNOR-Net	ResNet-18	90.06	-	$0.349^{\dagger}$
$MAESTRO^{\dagger}$ $(\lambda_{gp} = 16e^{-6})$	ResNet-18	$94.19 \scriptstyle{\pm 0.07}$	$0.39{\scriptstyle\pm0.00}$	$4.08{\scriptstyle\pm0.02}$
$\begin{array}{l}\text{MAESTRO}^{\dagger}\\ (\lambda_{gp} = 64e^{-6})\end{array}$	ResNet-18	93.86±0.11	$0.15{\scriptstyle\pm0.00}$	$1.23{\scriptstyle \pm 0.00}$
Non-factorized	VGG-19	$92.94_{\pm0.17}$	0.40	20.56
Pufferfish	VGG-19	92.69	0.29	8.37
Cuttlefish	VGG-19	93.39	0.15	2.36
RareGems	VGG-19	86.28	-	5.04
IMP	VGG-19	92.86	-	5.04
XNOR-Net	VGG-19	88.94	-	$0.64^{\dagger}$
Spectral Init.*	VGG-19	83.27	-	$\approx 0.4$
$MAESTRO^{\dagger}$ $(\lambda_{gp} = 32e^{-6})$	VGG-19	$93.10{\scriptstyle\pm0.10}$	$0.13{\scriptstyle \pm 0.00}$	$2.20{\scriptstyle \pm 0.03}$
$MAESTRO^{\dagger}$ $(\lambda_{gp} = 512e^{-6})$	VGG-19	$88.53{\scriptstyle\pm0.13}$	$\textbf{0.03} {\pm 0.00}$	$0.35{\scriptstyle\pm0.00}$

<sup>\*</sup>Results from original work; †: XNOR-Net employs binary weights and activations; although the overall #trainable parameters remain the same as the vanilla network, each model weight is quantized from 32-bit to 1-bit. Therefore, we report a compression rate of 3.125%(1/32).