

# BLOCKSWAP: FISHER-GUIDED BLOCK SUBSTITUTION FOR NETWORK COMPRESSION ON A BUDGET

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## ABSTRACT

The desire to map neural networks to varying-capacity devices has led to the development of a wealth of compression techniques, many of which involve replacing standard convolutional blocks in a large network with cheap alternative blocks. However, not all blocks are created equally; for a required compute budget there may exist a potent combination of many different cheap blocks, though exhaustively searching for such a combination is prohibitively expensive. In this work, we develop *BlockSwap*: a fast algorithm for choosing networks with interleaved block types by passing a single minibatch of training data through randomly initialised networks and gauging their *Fisher potential*. These networks can then be used as students and distilled with the original large network as a teacher. We demonstrate the effectiveness of the chosen networks across CIFAR-10 and ImageNet for classification, and COCO for detection, and provide a comprehensive ablation study of our approach. BlockSwap quickly explores possible block configurations using a simple architecture ranking system, yielding highly competitive networks in orders of magnitude less time than most architecture search techniques (e.g. 8 minutes on a single CPU for CIFAR-10).

## 1 INTRODUCTION

Convolutional Neural Networks (CNNs) are extremely popular, and demonstrate strong performance on a variety of challenging tasks. Because of this, there exists a large range of scenarios in the wild in which practitioners wish to deploy these networks e.g. pedestrian detection in a vehicle’s computer, human activity recognition with wearables (Radu et al., 2018).

CNNs are largely over-parameterised; a fact that can be exploited when specialising networks for different resource budgets. For example, there is a wealth of work demonstrating that it is possible to replace expensive convolutional blocks in a large network with cheap alternatives e.g. those using grouped convolutions (Chollet, 2017; Xie et al., 2017; Ioannou et al., 2017; Huang et al., 2018) or bottleneck structures (He et al., 2016; Sandler et al., 2018; Peng et al., 2018). This creates a smaller network which may be used as a *student*, and trained through distillation (Ba & Caruana, 2014; Hinton et al., 2015) with the original large network as a teacher to retain performance.

Typically, each network block is replaced with a single cheap alternative, producing a single-blocktype network for a given budget. By cheapening each block equally, one relies on the assumption that each block is of equal importance. We posit instead that for each budget, there exist more powerful *mixed-blocktype networks* that assign non-uniform importance to each block by cheapening them to different extents. We now have a paradigm of choice; for a given budget, it is not obvious *which* cheap alternatives to use for our student network, nor *where* to place them.

Let’s assume we have a large network that consists of  $B$  expensive convolutional blocks, and a candidate pool of  $C$  cheap blocks that we could substitute each of these out for, and we are given a limit on the number of parameters (and therefore memory) that the network can use. Which of the  $C^B$  *mixed-blocktype* networks is appropriate? It rapidly becomes infeasible to exhaustively consider these networks, even for single digit  $C$ . We could turn to neural architecture search (NAS) techniques (Zoph & Le, 2017; Pham et al., 2018; Tan et al., 2019; Wu et al., 2019; Liu et al., 2019a) but these introduce a gigantic computational overhead; even one of the fastest such approaches (Pham et al., 2018) still requires half a day of search time on a GPU. Moreover, in real-world scenarios, allocated budgets can change quickly; we would like to be able to find a good network in a matter of minutes, not hours or days.

Figure 1: The three-step BlockSwap pipeline. Beginning with a large network, we sample a list of candidate architectures by replacing its blocks with cheap alternatives. In Step (2), we rank each candidate by its Fisher potential after a single minibatch of training data. In Step (3), we select the highest ranked architecture and train it with attention transfer from the original network.

Our goal in this paper is, given a desired parameter budget, to quickly identify a suitable blocktypeversion of the original network that makes for a powerful student. We present a simple method—BlockSwap—to achieve this. First, we randomly sample a collection of candidate blocktypearchitectures that satisfy a desired parameter budget. A single minibatch is then pushed through each candidate network to calculate Fisher potential the sum of the total Fisher information for each of its blocks. Finally, the network with the highest potential is selected as a student and trained through distillation—specifically, attention transfer (Zagoruyko & Komodakis, 2017)—with the original teacher. Our method is illustrated in Figure 1.

In Section 3 we describe the block substitutions used in BlockSwap, distillation via attention transfer, and Fisher information. We describe our method in Section 4 as well as providing a comprehensive ablation study. Finally, we experimentally verify the potency of BlockSwap on CIFAR-10 (Section 5) as well as ImageNet (Section 6) and COCO (Section 7). Our contributions are as follows:

1. We introduce BlockSwap, an algorithm for reducing large neural networks by performing block-wise substitution. We show that this outperforms other top-down approaches such as depth/width scaling, parameter pruning, and random substitution.
2. We outline a simple method for quickly evaluating candidate models via Fisher information, which matches the performance of bottom-up approaches while reducing search time from days to minutes.
3. We conduct ablation studies to validate our methodology, highlighting the benefits of block mixing, and confirming that our ranking metric is highly correlated to the final error.

## 2 RELATED WORK

Neural networks tend to be overparameterised: Denil et al. (2013) accurately predict most of the weights in a network from a small subset; Frankle & Carbin (2019) hypothesise that within a large network, there exists a fortuitously initialised subnetwork that drives its performance. However, it remains difficult to exploit this overparameterisation without taking a hit in performance.

One means to combat this is to use a large network to regularise the training of a small student network; a process known as distillation. The small network is trained from scratch, but is also forced to match the outputs (Ba & Caruana, 2014; Hinton et al., 2015) or activation statistics (Romero et al., 2015; Zagoruyko & Komodakis, 2017; Kim et al., 2018) of the teacher using an additional loss term. When utilising distillation one must decide how to create a student network. A simple approach would be to reduce the depth of the original large network, although this can prove detrimental (Urban et al., 2017). An effective strategy is to create a student by replacing all the teacher’s convolutions with grouped alternatives (Crowley et al., 2018).

Grouped convolutions are a popular replacement for standard convolutions as they drastically cut the number of parameters used by splitting the input along the channel dimension and applying a much cheaper convolution on each split. They were originally used in AlexNet (Krizhevsky et al., 2012) due to GPU memory limitations, and have appeared in several subsequent architectures (Ioffe & Szegedy, 2015; Chollet, 2017; Xie et al., 2017; Ioannou et al., 2017; Huang et al., 2018). However,

as the number of groups increases, fewer channels are mixed, which hinders representational capacity. MobileNet (Howard et al., 2017) compensates for this by following its heavily-grouped depthwise convolutions by a pointwise (1) convolution to allow for channel mixing.

The increasing complexity of neural network designs has encouraged the development of methods for automating neural architecture search (NAS). Zoph & Le (2017) use an RNN to generate network descriptions and iter the options using reinforcement learning. Storing such a large quantity of possible networks is expensive, and their evaluation strategy uses 450 GPUs over the course of 3 days. To address this, Pham et al. (2018) propose giving all models access to a shared set of weights, achieving similar performance to Zoph & Le (2017) with a single GPU in less than 24 hours. Subsequent works have made extensive use of this technique (Liu et al., 2019a; Luo et al., 2018; Chen et al., 2018). However, it has been observed that under the constrained architecture search space of the above methods, random architecture search provides a competitive baseline (Li & Talwalkar, 2019; Sciuto et al., 2019). In particular, Sciuto et al. (2019) show that weight sharing hampers the ability of candidate networks to learn and causes many NAS techniques to find suboptimal architectures.

NAS techniques predominantly take a bottom-up approach; they find a powerful building block, and form neural networks using stacks of these blocks. Other works have taken a top-down approach by searching for architectures using pruning (Lee et al., 2019; Liu et al., 2019b; Frankle & Carbin, 2019). SNIP (Lee et al., 2019) is a fast and effective means of doing this: one randomly initialises a large network and quantifies the sensitivity of each connection using gradient magnitudes. The lowest sensitivity connections are removed to produce a sparse architecture which is then trained as normal.

### 3 PRELIMINARIES

#### 3.1 SUBSTITUTE BLOCKS

Here, we will briefly elaborate on the block substitutions used in BlockSwap. A tabular comparison is given in Appendix A. The block choices are deliberately simplistic. We can therefore demonstrate that it is the combination of blocks that is important rather than the representational capacity of a specific highly-engineered block e.g. one from the NAS literature.

**Standard Block – S** The blocks considered are variations of the standard block used in residual networks. In the majority of these blocks, the input has the same number of channels as the output, so we describe their parameter cost assuming this is the case. This standard block contains two convolutional layers, each using lots of  $N \times k \times k$  filters where  $k$  is the kernel size. Assuming the costs of batch-norm (BN) layers and shortcut convolutions (where applicable) are negligible, the block uses a total of  $2N^2k^2$  parameters.

**Grouped+Pointwise Block – G(g)** In a grouped convolution, the channel input is split along the channel dimension into groups, each of which has  $g$  channels. Each group goes through its own convolution which outputs  $g$  channels, and all the outputs are concatenated along the channel dimension. This uses  $(N/g)^2k^2 = (N^2k^2)/g$  parameters. To compensate for reduced channel mixing, each convolution is further followed by a pointwise (1) convolution, incurring an extra cost of  $N^2$ . For this block, each full convolution has been replaced with a grouped+pointwise convolution, and so the block uses  $2((N^2k^2)/g + N^2)$  parameters.

**Bottleneck Block – B(b)** In this block, a pointwise convolution is used to reduce the number of channels of the input by a factor  $b$  before a standard convolution is applied. Then, another pointwise convolution brings the channel size back up. This uses  $(N/b)^2k^2 + 2N^2 = b$  parameters.

**Bottleneck Grouped+Pointwise Block – BG(b, g)** This is identical to the bottleneck block, except the standard convolution is further split into groups, and so uses  $(N/bg)^2k^2 + 2N^2 = b$  parameters.

#### 3.2 DISTILLATION VIA ATTENTION TRANSFER

Attention transfer (Zagoruyko & Komodakis, 2017) is a distillation technique whereby a student network is trained such that its attention maps several distinct attention points are made to be similar to those produced by a large teacher network. A formal definition follows: Consider a choice of layers  $i = 1; 2; \dots; N_L$  in a teacher network, and the corresponding layers in the student network. At each chosen layer of the teacher network, collect the spatial map of the activations for channel  $j$  into the vector  $a_{ij}^t$ . Let  $A_i^t$  collect  $a_{ij}^t$  for all  $j$ . Likewise for the student network we correspondingly collect into  $a_{ij}^s$  and  $A_i^s$ . Now given some choice of mapping  $f(A_i)$  that maps each collection of the

form  $A_i$  into a vector, attention transfer involves learning the student network by minimising

$$L_{AT} = L_{CE} + \sum_{i=1}^L \frac{f(A_i^t)}{\|f(A_i^t)\|_2} \cdot \frac{f(A_i^s)}{\|f(A_i^s)\|_2}; \tag{1}$$

where  $\lambda$  is a hyperparameter, and  $L_{CE}$  is the standard cross-entropy loss. In Zagoruyko & Komodakis (2017) the authors use  $f(A_i) = (1 - \frac{1}{N_{A_i}}) \sum_{j=1}^{N_{A_i}} a_{ij}^2$ , where  $N_{A_i}$  is the number of channels at layer

### 3.3 FISHER INFORMATION

Theis et al. (2018) derive a second order approximation of the change in loss that would occur on the removal of a particular channel activation in a neural network. They use this signal to identify the least important activation channels, and remove their corresponding weights while performing channel pruning. Formally, let us consider a single channel of an activation in a network due to some input minibatch of  $N$  examples. Let us denote the values for this channel as an  $N \times W \times H$  tensor where  $W$  and  $H$  are the channel's spatial width and height. Let us refer to the entry corresponding to example  $n$  in the mini-batch at location  $(i, j)$  as  $A_{nij}$ . If the network has a loss function  $L$ , then we can back-propagate to get the gradient of the loss with respect to this activation channel. Let us denote this gradient as  $g_{nij}$ .  $c$  can then be computed by

$$c = \frac{1}{2N} \sum_n \sum_i \sum_j A_{nij}^2 g_{nij}^2 \tag{2}$$

Using an approximation to the Taylor expansion of the loss around an activation originated in LeCun et al. (1989) and has inspired many works in pruning (Hassibi & Stork, 1993; Molchanov et al., 2017; Guo et al., 2016; Srinivas & Babu, 2015) and quantisation (Choi et al., 2017; Hou et al., 2017).

## 4 METHOD

Let us denote a large teacher network composed of  $B$  blocks each of type  $S$  as  $T = [S_1; S_2; \dots; S_B]$ . Each of these may be replaced by a cheap block chosen from a list of candidates  $C_1; C_2; \dots; C_N$  of various representational capacities. We wish to construct a smaller model  $M = [C_{r1}; C_{r2}; \dots; C_{rB}]$  that is powerful, and within a given parameter budget. But, the space of possible block configurations is very large ( $\sim 10^{23}$  in Section 5). Even when using a cheap network evaluation strategy it is not possible to exhaustively search through the available network architectures. What we require is a method to quickly propose and score possible configurations with as little training as possible.

BlockSwap therefore relies on randomly sampling block configurations that satisfy a parameter budget, and ranking the samples by Fisher potential to decide which architectures to train. We obtain a score for each network as follows: we place a probe after the last convolution in each block. When a single minibatch of training data is passed through the network and the resulting cross-entropy loss is backpropagated, the probe measures the Fisher potential of the block by summing (Equation 2) for each channel in the layer it is placed after. For this step, minibatch size is set equal to the size used during training. We then sum the Fisher potential of each block to give us a score for the network.

The intuition for this metric is as follows: the Fisher information measures the sensitivity of the loss to each parameter at initialisation. Where parameters are insensitive at initialisation, they commonly remain insensitive as the other parameters are adjusted to adapt to the signal. Hence the accumulated Fisher potential measures the parameter wastage: how much of the parameter space is likely to be redundant for learning. We wish to choose architectures which are efficient (low parameter numbers) and have low wastage (use the parameters they have well).

Once we have scored each candidate architecture using the Fisher potential, we select the best one and train it using attention transfer. The training hyperparameters can be mirrored from the use of the following blocks (defined in Section 3.1) as candidate blocks:

- B(b) for  $b \in \{2, 4\}$
- G(g) for  $g \in \{2, 4, 8, 16\}$ ;  $N \in \{16, 8, 4, 2\}$ ;  $N_g$
- BG(2; g) for  $g \in \{2, 4, 8, 16\}$ ;  $M \in \{16, 8, 4, 2\}$ ;  $M_g$

where  $N$  is the number of channels in a block, and  $m$  is the number of channels after a bottleneck—this is  $N=2$  when  $b=2$ . We also use the standard block as a candidate choice, so as to not force blocks to reduce capacity where it is imperative.

#### 4.1 WHY MIX BLOCK TYPES?

A reasonable question to ask is whether mixed-blocktype architectures will perform better than the use of a single blocktype. As a simple check for this, we chose parameter budgets of 200K, 400K, and 800K and at each budget, compared single-blocktype models to one hundred randomly assembled, mixed-blocktype models. The models are WideResNets (Zagoruyko & Komodakis, 2016) with substituted blocks trained on CIFAR-10.

Figure 2: For parameter budgets of (a) 200K, (b) 400K and (c) 800K we train a single blocktype network and 100 random mixed blocktype networks. This lets us examine the distribution of mixed blocktype architectures to verify the existence of performant ones. The dotted red line on each plot represents models composed of a single block type, and the shaded blue density represents randomly constructed, mixed-blocktype networks. For example, (a) shows that a WRN-40-2 with every block substituted for  $BG(2,4)$  trains to 5.0% error, while the mean error of randomly constructed networks at the same parameter budget lies at around 5.15% error. This implies that random architecture search will not work here, and that a better strategy is needed.

In Figure 2 we plot the final test error of the random architectures (blue densities) against their single-blocktype counterparts (the red dotted lines). We observe that for each of the given parameter budgets there exist combinations of mixed blocks that are more powerful than their single blocktype equivalents. However, single-blocktype networks typically sit below the mean of the density of the random networks. This suggests that, though random search has been shown to work well in architecture search settings (Sciuto et al., 2019), it will yield sub-optimal structures in this case.

#### 4.2 WHY USE A SINGLE MINIBATCH? HOW DOES YOUR RANKING METRIC COMPARE?

Though there are no common metrics for architecture ranking, neural architecture search methods often use information from early learning stages to evaluate possible candidates. In order to assess the fastest, most accurate indicator of final error, we examine our 100 random mixed-blocktype networks for parameter budgets of 200K, 400K, and 800K and record how final error correlates with (i) train accuracy, (ii) total  $L_2$  norms, (iii) summed absolute gradient norms (Golub et al., 2019), and (iv) Fisher potential after 1, 10, and 100 minibatches of training in Table 1. We found that Fisher potential was by far the most robust metric, and that there were diminishing returns for adding more than a single minibatch.

#### 4.3 HOW MANY SAMPLES ARE NEEDED?

Figure 2 suggests that sampling 100 random architectures will reliably yield at least one mixed-blocktype network that outperforms a similarly parameterised single blocktype network. However, accepting that there is some noise to our ranking metric, we assume that we will need to take more than 100 samples in order to reliably detect powerful architectures. As an illustration, at the budget of 400K parameters a single blocktype alternative has a test error of 4.5%, whereas BlockSwap finds networks with final test errors of 4.85%, 4.54%, and 4.21% after 10, 100, and 1000 samples respectively. We empirically found that 1000 samples was enough to be robust to various parameter budgets on the tasks we considered.

Table 1: Spearman Rank correlation scores for several ranking metrics when ranking 100 random architectures at three parameter budgets. Negative correlation implies that as the metric score goes up, val error goes down. Summing gradient norms follows a similar pattern to summing Fisher potential but appears to be less robust and moderately less accurate. Gradients are extremely volatile and largely uninformative. Crucially, adding more than 1 minibatch has diminishing returns.

	200K Params.			400K Params.			800K Params		
Minibatches	1	10	100	1	10	100	1	10	100
Accuracy	-0.004	-0.022	-0.495	-0.068	-0.282	-0.386	0.042	-0.211	-0.257
$\ell_2$ norms	-0.083	-0.185	-0.289	0.093	-0.213	-0.251	0.368	0.226	0.140
Grad Norms	-0.541	-0.490	0.326	-0.612	-0.444	0.402	-0.608	0.340	0.558
Fisher	-0.602	-0.621	-0.439	-0.685	-0.667	-0.508	-0.635	-0.638	-0.277

#### 4.4 WHAT DO “GOOD NETWORKS” LOOK LIKE?

For each of the three parameter budgets (200K, 400K, 800K) we inspected the most common block choices for “good” and “bad” networks, where good and bad are networks with val error greater two standard deviations below or above the mean respectively. We found that overwhelmingly, the types of blocks used in both good and bad networks was very similar, implying that the placement of the blocks instead of the types of the blocks that matters. We examine this further in Appendix B.

## 5 CIFAR EXPERIMENTS

Here, we evaluate student networks obtained using BlockSwap on the CIFAR-10 image classification dataset (Krizhevsky, 2009). We benchmark these against competing student networks for a range of parameter budgets. To recapitulate, the BlockSwap networks are found by taking 1000 random samples from the space of possible block combinations that satisfy our constraint (in this case, parameter budget). These points are ranked by Fisher potential after a single minibatch of training data, and the network with the highest potential is chosen.

The structure we use is that of a WideResNet (Zagoruyko & Komodakis, 2017), since we can use them to construct compact, high performance networks that are broadly representative of the most commonly used neural architectures. A WideResNet with depth 40, and width multiplier 2—WRN-40-2—is trained and used as a teacher. It consists of 18 blocks and has 2.2 million parameters. A student network is generated and is trained from scratch using attention transfer with the teacher. Our BlockSwap students are WRN-40-2 nets where each of its 18 blocks are determined using our method with the blocks outlined in Section 4. We compare against the following students:

1. Reduced width/depth versions of the teacher: WRN-16-1, WRN-16-2, WRN-40-1
2. Single block-type teacher reductions: every block in the teacher is swapped out for (i) the MBConv6 block from MobileNet v2 (Sandler et al., 2018) and (ii) the block discovered in DARTS (Liu et al., 2019a).
3. Pruned teacher reductions: we compare against (i) the magnitude-based pruning methodology from Han et al. (2016) and (ii) SNIP (Lee et al., 2019) versions of the teacher.

We additionally compare against CondenseNet-86 (Huang et al., 2018), distilled with the Born-Again strategy described by Furlanello et al. (2018).

First, we train three teacher networks independently. These are used to train all of our students; each student network is trained three times, once with each of these teachers. Figure 3 shows the mean test errors of BlockSwap students at various parameter counts, compared to the alternatives listed above. Full results with standard deviations are listed in Table 2, along with the number of Multiply-Accumulate (MAC) operations each network uses.

Our results show that block cheapening is more effective than simple downscaling schemes (reducing width/depth); not only are the reduced models (WRN-40-1, WRN-16-2, WRN-16-1) in exible in parameter range, they perform significantly worse than our models at the parameter budgets they can map to. We also show that BlockSwap finds more accurate networks than the other top-down approaches (l-pruning and SNIP) across the full parameter spectrum. Note that we are being

Table 2: CIFAR-10 top-1 test error for student nets, with parameter count (in thousands, as P.(K)) and total MAC operations (in millions, as Ops(M)). D-W specifies the number of layers and the width multiplier of the student. We compare BlockSwap to reductions that rely on reducing depth (D) and width (W), single-blocktype networks (MBConv6, DARTS, CondenseNet), and pruning via SNIP (Lee et al., 2019). Further comparisons are given in Appendix C. We do not report Ops for SNIP since this is dependent on choice of sparse representation format. BlockSwap is able to choose the networks with the lowest mean error for all parameter budgets considered.

D-W Method	P. (K)	Ops (M)	Err.	D-W Method	P. (K)	Ops (M)	Err.
40-2 Teacher	2243.5	328.3	3.96	40-2 SNIP	404.2	-	4.67 0.11
16-2 D-Scaled	691.7	101.4	4.78	40-2 SNIP	289.2	-	5.04 0.06
40-1 W-Scaled	563.9	83.6	4.57	40-2 SNIP	217.0	-	5.53 0.08
16-1 D-W-Scaled	175.1	26.8	7.32	40-2 SNIP	162.2	-	6.00 0.16
40-2 MBConv6	1500.9	231.7	5.32	40-2 BlockSwap	811.4	132.5	3.79 0.01
40-2 DARTS	321.5	52.8	4.49	40-2 BlockSwap	556.0	89.5	4.17 0.22
86- CondenseNet	520.2	65.2	4.95	40-2 BlockSwap	404.2	92.8	4.21 0.13
40-2 SNIP	811.4	-	4.13	40-2 BlockSwap	289.2	65.9	4.45 0.18
40-2 SNIP	556.0	-	4.32	40-2 BlockSwap	217.0	38.8	4.68 0.37
				40-2 BlockSwap	162.2	33.9	5.17 0.00

Figure 3: CIFAR-10 top-1 test error of students versus parameters. BlockSwap (the blue line) models give lower error for each parameter budget when compared to depth/width reduced or pruned models. They also outperform single blocktype networks (MBConv, DARTS, CondenseNet-86).

particularly generous to the sparse methods by counting only non-zero parameters (the weight tensors remain the same size as the original teacher).

The mixed block architectures that BlockSwap generates are more accurate at these reduced budgets than all of the single blocktype alternatives we considered. While the performance of DARTS is very similar to BlockSwap, it is worth noting that the DARTS architecture search required 24 GPU hours, while BlockSwap needs only 8 minutes on a single CPU.

Given the strength of random baselines in architecture search settings (Liu et al., 2019a; Sciuto et al., 2019; Li & Talwalkar, 2019), we also compare BlockSwap against randomly generated block configurations in Appendix C. BlockSwap consistently outperforms these, proving that our aggregated Fisher potential metric is an effective means for choosing optimal block structures.

Implementation Details: Networks are trained for 200 epochs using SGD with momentum 0.9. The initial learning rate of 0.1 is reduced by a factor of 5 every 60 epochs. Minibatches of size 128 are used with standard crop + flip data augmentation and Cutout (DeVries & Taylor, 2017). The weight decay factor is set to 0.0005. For attention transfers set to 1000.

## 6 IMAGENET CLASSIFICATION

Here, we demonstrate that students chosen by BlockSwap succeed on the more challenging ImageNet dataset (Russakovsky et al., 2015). We use a pretrained ResNet-34 (16 blocks) as a teacher, and compare students at two parameter budgets (3M and 8M). We train a Blockswap student at each of these budgets and compare their validation errors to those of a reduced depth/width student (ResNet18 and ResNet-18-0.5—a ResNet-18 where the channel width in the last 3 sections has been halved) and a single-blocktype student (ResNet-34 with G(4) and G(N) blocks). The student networks found by BlockSwap for these two budgets are illustrated in Appendix D. Top-1 and top-5 validation errors are presented in Table 3. At both budgets, BlockSwap chooses networks that outperform its comparators. At 8M parameters it even surpasses the teacher by quite a margin. Specifically, it beats the teacher by 0.49% in top-1 error and 0.82% in top-5 error despite using almost fewer parameters.

Implementation Details: Networks are trained with a cross-entropy loss for 100 epochs using SGD with momentum 0.9. The initial learning rate of 0.1 is reduced by every 30 epochs. Minibatches of size 256 are used with standard crop + ip augmentation. The weight decay factor is set to 0.0001. For attention transfer is set to 750 using the output of each of the four sections of network.

Table 3: Top-1 and Top-5 classification errors (%) on the validation set of ImageNet for students trained with attention transfer from a ResNet-34. We can see that for a similar number of parameters, the student found from BlockSwap outperforms its counterparts, and in one instance, the teacher.

Model	Params	MACs	Top-1 err	Top-5 err
ResNet-34Teacher	21.8M	3.669G	26.73	8.57
ResNet-18	11.7M	1.818G	29.18	10.05
ResNet-34-G(4)	8.1M	1.395G	26.58	8.43
BlockSwap	8.1M	1.242G	26.24	7.75
ResNet-18-0.5	3.2M	909M	37.20	15.02
ResNet-34-G(N)	3.1M	559M	30.16	10.66
BlockSwap	3.1M	812M	29.57	10.20

## 7 COCO DETECTION

Thus far, we have used BlockSwap for image classification problems. Here we observe whether it extends to object detection on the COCO dataset (Lin et al., 2014)—specifically, training on 2017 train, and evaluating on 2017 val. We consider a Mask R-CNN (He et al., 2017) with a ResNet-34 backbone, and apply BlockSwap using COCO images to obtain a mixed-blocktype backbone with 3M parameters. We compare this to a single-blocktype ResNet-34-G(N) backbone which uses the same number of parameters. To avoid contamination with ImageNet, we train everything from scratch. The results can be found in Table 4. We can see that the BlockSwap backbone again outperforms its single-blocktype counterpart.

Implementation Details: Networks are trained using the default Mask R-CNN settings in Torchvision. We use a batch-size of 16 split across 8 GPUs. All models are trained from scratch, and we forgo distillation due to memory constraints.

Table 4: Average Precisions (%) for COCO-2017 val detection for Mask-RCNNs using a ResNet-34-G(N) and BlockSwap backbone (each using 3M parameters).

Backbone	AP	AP <sub>50</sub>	AP <sub>75</sub>	AP <sub>S</sub>	AP <sub>M</sub>	AP <sub>L</sub>
ResNet34-G(N)	22.3	38.9	23.0	12.9	22.9	30.7
BlockSwap	23.4	40.0	24.7	13.6	24.2	31.1

## 8 CONCLUSION

We have developed BlockSwap: a fast, simple method for reducing large neural networks to exible parameter targets based on block substitution. We have verified that these reduced networks make for excellent students, and have performed a comprehensive ablation study. Future work could use BlockSwap to choose networks based on inference time, or energy cost instead of parameter count.



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## A BLOCK REPLACEMENTS

Table 5: Block substitutions used in this paper: Conv refers to a convolution. GConv is a grouped  $k \times k$  convolution and Conv1x1 is a pointwise convolution. We assume that the input to each block has  $N$  channels and that channel size doesn't change unless written explicitly as  $(y)$ . Where applicable  $g$  is the number of groups and  $b$  is the bottleneck contraction. The convolutional and batch-norm (BN) costs are given, although the latter is significantly smaller. BN+ReLU precedes each convolution for WideResNets and follows each convolution for standard ResNets.

Block	S	G(g)	B(b)	BG(b; g)
Structure	Conv Conv	GConv(g) Conv1x1 GConv(g) Conv1x1	Conv1x1( $\frac{N}{b}$ ) Conv Conv1x1( $\frac{N}{b}$ )	Conv1x1( $\frac{N}{b}$ ) GConv(g) Conv1x1( $\frac{N}{b}$ )
Conv Params	$2N^2k^2$	$2N^2(\frac{k^2}{g} + 1)$	$N^2(\frac{k^2}{b^2} + \frac{2}{b})$	$N^2(\frac{k^2}{gb^2} + \frac{2}{b})$
BN Params	$4N$	$8N$	$N(2 + \frac{4}{b})$	$N(2 + \frac{4}{b})$

## B UNDERSTANDING BLOCK PLACEMENT

To understand what makes for “good” block placement, we begin with the more tractable question: if we were only to cheapen one block, which one should we choose? In Figure 4 we show what happens when we substitute a single standard block (G) for each of the possible blocks for our WideResNet. We plot the final test error after training to convergence, as well as the total parameter count after the substitution (the higher the bar, the higher the error, therefore the more sensitive the block).

Notice that in Figure 4 (a), substituting the start and end of a stage appears to have a very detrimental effect on accuracy, whereas mid-stage blocks can be substituted without hurting performance. For example, block 12 has few parameters, as shown in Figure 4(b), but substituting it for a cheaper alternative results in a drastic drop in accuracy. This implies that it is safest to substitute blocks in the middle of a stage but that blocks at the start and end of each stage should have their representational capacity preserved as much as possible.

We used this insight to analyse the block configurations of “good” and “bad” amongst the random architectures at each parameter budget. With several blocktypes, however, we need a method to rank the capacity of blocktypes; we make the simplifying assumption that training the blocktype on its own is a good indicator of this (though this is not always true, since blocks give varying performance with different input sizes). An example of a good and bad network judged this way can be seen in Figure 5; while they do roughly follow our observations, there are clearly more complex interactions at play (perhaps unsurprising given the random nature of their construction). We found that overall, the models in our ablation set adhere to the pattern described above (for each stage: high capacity blocks at the start and end, increasing capacity throughout), with some variance.

Figure 4: Each bar shows a WideResNet-40-2 with one block substituted (G), where (a) shows final CIFAR-10 error and (b) shows resulting number of parameters, with blocks coloured by network stage. This shows us the sensitivity of each block; for example, block 12 has relatively few parameters, but substituting it for a cheaper block is very detrimental to final error.

Figure 5: An example of (a) a bad architecture and (b) a good architecture for a given parameter budget (200K), with blocks coloured by network stage. Capacity score is given by the accuracy that the selected blocktype achieves after being trained on its own. We can see that the good model (b) follows the pattern observed in Figure 4, while the bad model (a) does not.

### C COMPARING TO RANDOM CONFIGURATIONS

Table 6: Comparing BlockSwap to randomly configured networks and pruned versions of the original teacher on the CIFAR-10 dataset. Total MAC operations are not reported for pruning because calculating this is highly dependent on choice of sparse tensor representation format.

D-W	Method	Parameters (K)	MAC Ops (M)	Error ( )
40-2	BlockSwap	811.4	132.5	3.790.01
40-2	BlockSwap	556.0	89.5	4.170.22
40-2	BlockSwap	404.2	92.8	4.210.13
40-2	BlockSwap	289.2	65.9	4.450.18
40-2	BlockSwap	217.0	38.8	4.680.37
40-2	BlockSwap	162.2	33.9	5.170.00
40-2	Random	795.7	81.6	4.260.04
40-2	Random	551.1	76.7	4.540.18
40-2	Random	397.4	59.5	4.910.12
40-2	Random	285.1	56.7	4.800.12
40-2	Random	217.6	46.9	5.130.24
40-2	Random	168.3	33.6	5.750.09
40-2	\`1-pruning	894.7	-	4.12 0.03
40-2	\`1-pruning	760.5	-	4.32 0.06
40-2	\`1-pruning	671.1	-	4.51 0.13
40-2	\`1-pruning	536.8	-	4.98 0.21
40-2	\`1-pruning	447.4	-	5.60 0.23
40-2	\`1-pruning	313.2	-	7.70 0.18
40-2	\`1-pruning	223.7	-	11.93 0.77

## D BLOCKSWAP NETWORKS FOR IMAGENET

Figure 6: ResNet-34 block substitutions chosen by BlockSwap at 3M (top) and 8M (bottom) parameters on ImageNet.