REAP: A Method for Pruning Convolutional Neural Networks with Performance Preservation*

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SUMMARY This paper presents a pruning method, Reconstruction Error Aware Pruning (REAP), to reduce the redundancy of convolutional neural network models for accelerating their inference. In REAP, we have the following steps: 1) Prune the channels whose outputs are redundant and can be reconstructed from the outputs of other channels in each convolutional layer; 2) Update the weights of the remaining channels by least squares method so as to compensate the error caused by pruning. This is how we compress and accelerate the models that are initially large and slow with little degradation. The ability of REAP to maintain the model performances saves us lots of time and labors for retraining the pruned models. The challenge of REAP is the computational cost for selecting the channels to be pruned. For selecting the channels, we need to solve a huge number of least squares problems. We have developed an efficient algorithm based on biorthogonal system to obtain the solutions of those least squares problems. In the experiments, we show that REAP can conduct pruning with smaller sacrifice of the model performances than several existing methods including the previously state-of-the-art one.

key words: pruning, REAP, biorthogonal system

1. Introduction

In various Computer Vision tasks, Convolutional Neural Networks (CNNs) have been showing the record-breaking performances. For example, some recent CNN models conduct object detection with high accuracy, and are expected to be used in assorted applications. Although, can we use those models as they are in the embedded systems, such as an object detection and classification system for in-vehicle camera? At this point, we need strong computational resources to use the large CNN models. For the future applications, it is important to produce fast CNN models without compromising their performances.

A major approach for producing a fast CNN model is to conduct pruning on a pretrained large model. Some early works aim for holistic network optimization based on the unified criteria across the layers, e.g. magnitude based criteria [2], loss function based one [3], and so on. However, most of these methods need a lot of mathematical approximations and/or use the heuristic criteria which do not guarantee the optimality of the pruning results. On the other hand, some recent works propose the pruning methods that focus on layer-wise optimization based on theoretically sound criteria [4]–[7] and those methods have turned out to be more effective than the earlier holistic methods. The weakness of the layer-wise methods is that the proper pruning ratio (the ratio of the number of the channels to be pruned) for each layer is not trivial and has to be determined manually. Although, in [8], reinforcement learning is employed to determine the pruning ratio for each layer, which enables us to achieve holistic network optimization using the layer-wise methods. Therefore, the layer-wise methods are becoming more important recently.

In this paper, we present a novel layer-wise pruning method, Reconstruction Error Aware Pruning (REAP), by extending Neuro-Unification (NU) [9] that we proposed in our previous work. The idea of NU is to unify a pair of neurons having highly correlated outputs so that the remaining one can reconstruct the outputs of the pruned one. We extend this idea from one-to-one unification to one-to-many. In REAP, we prune a neuron and reconstruct its outputs from all the remaining neuron’s outputs by solving the least squares problem.

However, our new approach requires tremendous computational cost. In REAP, we need to select the neuron to be pruned so that the output error is minimized. For this purpose, we have to repeat the following steps for each neuron: 1) prune a neuron, 2) solve the least squares problem to compute the error after reconstruction. This is computationally expensive when we have lots of neurons. Therefore, we also propose an efficient algorithm for this problem.

We conduct the experiments with several CNN models and datasets. The results show that our method can conduct pruning with smaller sacrifice of the model performances than the existing methods.

2. Related Works

The existing approaches on accelerating the CNN models can be categorized into 5 groups: 1) Factorization, 2) Sparsification, 3) Quantization, 4) Weight pruning, and 5) Channel pruning.

Factorization. The most fundamental method in this group is presented in [10]. They apply SVD to large weight matrix, and approximate it by the product of small matrices by discarding the components with small singular values. This results in reducing the parameters with small sacrifice of accuracy. For example, assume that a $m \times n$ matrix is approximated by the product of a $m \times o$ matrix and a $o \times n$ matrix. If $o \ll m, n$, the number of the parameters reduces...
from \(mn\) to \((m+n)\). Some other methods\cite{11,12} also belong here.

**Sparsification.** These methods make the weight tensors sparse by fine-tuning the models with L1 regularization\cite{13,14,15}. The theoretical weak point of sparsification is that L1 regularization shifts the global minimum of the cost function and sacrifices the model performances. Besides, in order to take the advantage of the sparsified models, the special hardwares and libraries are required for executing the computations on only non-zero weights.

**Quantization.** The methods in this group reduce the redundancy of each bit-wise operation\cite{16,17,18}, e.g., changing the floating point precision from 32-bit to 8-bit. The quantized models require the special hardwares and libraries when the pruned models are deployed.

**Weight pruning.** The idea of weight pruning is to remove the weights based on their saliencies\cite{3,7,14,17}. Similarly with sparsification methods, these methods aim to reduce the number of non-zero weights and do not aim to reduce the sizes of the weight tensors, therefore, the special hardwares and libraries are required when the pruned models are deployed.

**Neuron/Channel pruning.** The methods in this group prune the whole weights that belong to a certain neuron/channel\cite{4,5,6,19,20}. Therefore, the size of the weight tensor gets smaller, which results in saving the computational complexity for inference without any special hardwares and libraries.

Our method belongs to the channel pruning group. Nonetheless, it is possible to combine our method and the ones from other groups (e.g. quantization, factorization) to achieve further acceleration.

3. From NU to REAP

We explain NU, and show how this method can be improved by our new strategy. Then, we show the algorithms to accelerate the computation of REAP. We also explain how to apply REAP to convolutional layers.

3.1 Neuro-Unification (NU)

Let \(C\) and \(N\) denote the numbers of neurons in a layer and the next layer, and \(N\) denote the number of the input images. The forward propagation is formulated as

\[
Y = \sum_{n \in Z} x_n w_i^T
\]

where \(x_n \in \mathbb{R}^N\) denotes the outputs of the \(i\)-th neuron corresponding to \(N\) input images, \(w_i \in \mathbb{R}^C\) denotes the weights going from the \(i\)-th neuron to the ones in the next layer, \(Y \in \mathbb{R}^{N \times C}\) denotes the inner activation levels in the next layer, and \(Z = \{1, \cdots , c\}\) is the set of neuron indices. The goal is to reduce the number of neurons to the desired number while keeping \(Y\) as unchanged as possible.

**How to unify a given pair of neurons.** When we have a pair of neurons having linearly dependent outputs, we can prune one and merge it to the other one without error. For instance, if \(x_j = \alpha x_i\) holds, we prune the \(j\)-th neuron and update the \(i\)-th neuron’s weights as

\[
w_i^j \leftarrow w_i + \alpha w_j.
\]

Then, the error of \(Y\) is given by

\[
\Delta Y = x_i w_i^T - x_i w_i^T - x_j w_j^T = 0.
\]

This shows that \(Y\) is preserved because the \(i\)-th neuron compensates the error caused by pruning the \(j\)-th neuron.

In the case of unifying the neurons having linearly independent outputs, we first approximate \(x_j\) by a vector which is linearly dependent with \(x_i\):

\[
x_j \approx \alpha x_i.
\]

Then, we can prune the \(j\)-th neuron and update \(w_j\) by using Eq. (2). The question is how to determine \(\alpha\). The error of \(Y\) is given by

\[
\Delta Y = x_i w_i^T - x_i w_i^T - x_j w_j^T = - (x_j - \alpha x_i) w_i^T.
\]

Since we want to minimize the output error, we have

\[
\alpha^* = \arg \min_{\alpha} \left\| x_j - \alpha x_i \right\|_F^2
\]

\[
= \arg \min_{\alpha} \left\| x_j - \alpha x_i \right\|^2
\]

Note that we omitted \(w_i\) in Eq. (6) as it contains only constants. Thus, we can determine \(\alpha\) by computing orthogonal projection of \(x_j\) onto \(x_i\).

**How to select the pair of neurons to be unified.** We have to select the neurons to be unified when we have a lot of possible neuron pairs. As we want to minimize the error of \(Y\), we simply select the neuron pair that minimizes the output error:

\[
(i^*, j^*) = \arg \min_{(i,j)} \left\| x_j - \alpha x_i \right\|_F^2 \quad \text{s.t. } i \neq j.
\]

Note that we cannot omit \(w_j\) in Eq. (7) as it contains only constants.

3.2 Reconstruction Error Aware Pruning (REAP)

We extend the idea of NU from one-to-one unification to one-to-many.

**How to prune a neuron and reconstruct its outputs.** In NU, the output of the pruned neuron is reconstructed from another neuron. In REAP, we use all the remaining neurons for reconstruction, as shown in Fig. 1. This can be formulated by

\[
\{v_{ij} \mid i \in Z \setminus \{j\}\} = \arg \min_{v_{ij}} \left\| x_j - \sum_{i \in Z \setminus \{j\}} v_{ij} x_i \right\|_2^2.
\]
where the $v$-s denote the coefficients for reconstructing $x_j$ from the other $x$-s. Similarly with Eq. (2), the weights of the remaining neurons are updated as follows for each $i \in Z \setminus \{j\}$.

\[ w'_i = w_i + v_{ji}^* w_j. \]

(9)

**How to select the neuron to be pruned.** We should select the neuron to be pruned so as to minimize the reconstruction error of $Y$:

\[
\begin{align*}
    j^* &= \arg\min_j \left\| Y - \sum_{i \in Z \setminus \{j\}} x_i w_i'^\top \right\|_F^2 \\
    &= \arg\min_j \left\| Y - \sum_{i \in Z \setminus \{j\}} x_i (w_i + v_{ji}^* w_j)'^\top \right\|_F^2.
\end{align*}
\]

(10)

We prune the neurons one by one in a greedy fashion so that the error of $Y$ is as small as possible.

### 3.3 Algorithm for Efficient Computation

The problem of REAP is its computational cost. In order to solve Eq. (10), we need to solve Eq. (8) for each $j$, which is computationally expensive. Therefore, we have developed some efficient algorithms.

#### 3.3.1 How to Solve Eq. (8) for Each $j$ in One Shot.

We solve Eq. (8) for each $j$ in one shot by using biorthogonal system.

Let $r_j$ represent the residual of $x_j$ reconstructed from the other $x$-s:

\[ r_j = x_j - \sum_{i \in Z \setminus \{j\}} v_{ji}^* x_i. \]

(11)

As $r_j$ is the residual of $x_j$, $r_j$ is orthogonal to all other $x$-s. In other words, $r_j$ is orthogonal to the subspace spanned by $\{x_i|i \in Z \setminus \{j\}\}$, as shown in Fig. 2.

We compute $r_j$ by using biorthogonal system. Let $\{\bar{x}_i|i \in Z\}$ denote the dual bases of $\{x_i|i \in Z\}$. The biorthogonal system is defined by

\[
\langle x_j, \bar{x}_i \rangle = \begin{cases} 1 & (i = j) \\ 0 & (\text{otherwise}) \end{cases}
\]

(12)

\[
\bar{x}_j = \sum_{i \in Z \setminus \{j\}} v_{ji}^* x_i
\]

\[
U_{\{Z \setminus \{j\}\}} \sum_{i \in Z \setminus \{j\}} v_{ji}^* x_i
\]

Thus, we can solve Eq. (15) for $r_j$ by computing the orthogonal projection of $x_j$ onto $\bar{x}_j$, as shown in Fig. 2. Thus, the following must hold:

\[
\begin{align*}
    r_j &= \frac{\langle x_j, \bar{x}_j \rangle}{\|\bar{x}_j\|^2} \bar{x}_j = \bar{x}_j / \|\bar{x}_j\|^2. \\
    \bar{x}_j &= (X^g)^\top,
\end{align*}
\]

(15)

(16)

By using Eq. (15), we can compute $r_j$ for each $j \in Z$ in one-shot. Let $X = [x_1 \cdots x_c]$ and $\bar{X} = [\bar{x}_1 \cdots \bar{x}_c]$. By definition of dual bases, $\bar{X}$ can be computed as

\[
\bar{X} = (X^g)^\top
\]

(17)

where $X^g$ denotes the generalized inverse of $X$. Let $D$ denote the diagonal matrix whose $(i,i)$ entry is given by $1/\|\bar{x}_i\|^2$, and $R = [r_1 \cdots r_c]$. Since we have Eq. (15), we also have

\[
R = \bar{X} D.
\]

Let $V^*$ denote a matrix, and its $(j,i)$ entry is $v_{ji}^*$. Because we

![Fig. 1](image1.png) **Fig. 1** Concepts of NU and REAP. Left: original network. Center: network pruned with NU. Only one neuron is used for reconstruction. Right: network pruned with REAP. All other neurons are used for reconstructing the outputs of the pruned one.

![Fig. 2](image2.png) **Fig. 2** Illustration of the projection of $x_j$ onto a subspace $U_{\{Z \setminus \{j\}\}}$ spanned by $\{x_i|i \in Z \setminus \{j\}\}$. Computing the orthogonal projection of $x_j$ onto $U_{\{Z \setminus \{j\}\}}$ is equivalent to solving the problem of reconstructing $x_j$ from $\{x_i|i \in Z \setminus \{j\}\}$ by the least squares method. The residual $r_j$ is linearly dependent on $\bar{x}_j$, the dual basis for $x_j$. Where $\langle \cdot, \cdot \rangle$ denotes inner product. The biorthogonal expansion for $r_j$ is given by

\[
r_j = \sum_{i \in Z} \langle r_j, x_i \rangle \bar{x}_i.
\]

(13)

Obviously, $\langle r_j, x_i \rangle = 0$ holds for each $i \in Z \setminus \{j\}$, because $r_j$ is orthogonal to the subspace spanned by $\{x_i|i \in Z \setminus \{j\}\}$. Thus, Eq. (13) can be rewritten as

\[
r_j = \langle r_j, x_j \rangle \bar{x}_j + \sum_{i \in Z \setminus \{j\}} \langle r_j, x_i \rangle \bar{x}_i = \langle r_j, x_j \rangle \bar{x}_j.
\]

(14)

Equation (14) means that $r_j$ is linearly dependent on $\bar{x}_j$. Therefore, we can obtain $r_j$ by computing the orthogonal projection of $x_j$ onto $\bar{x}_j$, as shown in Fig. 2. Thus, the following must hold:

\[
r_j = \frac{\langle x_j, \bar{x}_j \rangle}{\|\bar{x}_j\|^2} \bar{x}_j = \frac{\bar{x}_j}{\|\bar{x}_j\|^2}.
\]

(15)
have Eq. (11), the following must hold:
\[ X = R + XV^*\top. \] (18)
Then, we have
\[ V^*\top = I - XV^*\top R \] (19)
This is equivalent to the solution of Eq. (8) for each \( j \).

Substitute approach. We have developed another algorithm for computing the reconstruction error of each neuron by using Gram-Schmidt process. The detail of this substitute approach is provided in Appendix A.

3.3.2 How to Solve Eq. (8) when We Prune Another Neuron

Assume that we have pruned the \( j \)-th neuron. When we prune another neuron, we just need to repeat the same procedures mentioned in Sect. 3.3 with the remaining neurons. We have to solve the following problem for each \( k \).

\[ \{u^*_k|i \in Z\{j,k\}\} = \arg\min_{u_k} \left\| x_k - \sum_{i \in Z\{j,k\}} u_k x_i \right\|^2, \] (20)
Then, we solve the following problem for selecting the next neuron to be pruned:
\[ k^* = \arg\min_{k} \left\| Y - \sum_{i \in Z\{j,k\}} \left( w^*_{ij} x_i + u^*_k w^*_{kj}\right) \right\|^2_{F}. \] (21)

As we already have the \( w^*\)-s, we only need to compute the \( u^*\)-s for solving Eq. (21). We may use biorthogonal system based algorithm again for solving Eq. (20). Although, by using the solution of Eq. (8), Eq. (20) can be solved even faster. We already have

\[ x_j = r_j + u^*_k x_k + \sum_{i \in \{j,k\}} v^*_ji x_i, \] (22)
\[ x_k = r_k + u^*_k x_j + \sum_{i \in \{j,k\}} v^*_ki x_i. \] (23)

After pruning both the \( j \)-th and the \( k \)-th neurons, we cannot use \( x_j \) for reconstructing \( x_k \). Thus, we substitute Eq. (22) to Eq. (23) and get
\[ x_k = r_k + u^*_k x_j + \sum_{i \in \{j,k\}} v^*_ki x_i. \] (24)

Obviously, we have \( \langle x_i, r_j \rangle = 0 \) and \( \langle x_i, r_k \rangle = 0 \) for each \( i \in Z\{j,k\} \). Therefore, \( \langle x_i, r_k + u^*_k x_i \rangle = 0 \) holds for each \( i \in \{j,k\} \), which means the first term of the RHS of Eq. (24) denotes the residual of \( x_k \) reconstructed from \( \{x_i|i \in Z\{j,k\}\} \) and the second term denotes the projection of \( x_k \) onto the subspace spanned by \( \{x_i|i \in Z\{j,k\}\} \). Thus, the coefficients of the \( x \)-s in the second term is equivalent to the solution of Eq. (20):

\[ u^*_k = \frac{v^*_ki + v^*_k j v^*_ji}{1 - v^*_k v^*_kj}. \] (25)

As above, we can compute \( u^\top\)-s without directly solving least squares problems. After computing \( u^\top\)-s, we can solve Eq. (21) easily.

3.3.3 Algorithm

To sum up, our algorithm can be described as below.
1. Initialize \( Z^* \leftarrow Z \), compute \( v^*_i \) and for each \( (i,j) \).
2. Solve Eq. (10) to select \( j \) and update
   \[ Z^* \leftarrow Z^* \setminus \{j\}, \]
   \[ w^*_{ij} \leftarrow w^*_{ij} + v^*_i j v^*_ji \] for each \( i \in Z^* \),
   \[ v^*_ki \leftarrow (v^*_ki + v^*_k j v^*_ji)/(1 - v^*_k v^*_kj) \] for each \( i, k \in Z^*, i \neq k. \)
3. If \( |Z^*| > c^* \) (\( c^* \) is the desired number of the neurons), go to step 2.

3.4 Applying REAP to Convolutional Layers

REAP can be applied to the convolutional layers with a minor modification. By expanding the feature maps and the kernels in the convolutional layer into matrices, we can deal with the convolutional layers in the same manner with the fully connected layers.

Same with “im2col” method implemented in cuDNN [21], we can describe the sliding window operations in the convolutional layers by the sum of the matrix multiplications. Let \( a \) and \( A \) denote the numbers of the input channels and the output channels, \( h \) denotes the width and the height of the feature maps, \( s \) denote the width and the height of the weight tensor. The sliding window operations with a \( R^{N \times A \times s \times h} \) tensor, which denotes the feature maps corresponding to the \( N \) input images, and a \( A \times a \times s \times s \) tensor, which denotes the weights, can be alternatively written as

\[ Y = \sum_{i \in B} \Phi_i \Psi_i^\top = \sum_{i \in B, m \in \mathbb{Z}} \Phi_{im} \Psi_{im}^\top, \] (26)

where \( B = \{1, \ldots, a\} \) denotes the set of the input channel indices, \( \Phi_i \in \mathbb{R}^{Nh \times s} \) denotes the \( i \)-th channel of the reshaped input feature maps, \( \Psi_i \in \mathbb{R}^{As \times s} \) denotes the reshaped weight tensor, \( T = \{1, \ldots, s^2\} \) denotes the set of the column indices of \( \Phi \)s and \( \Psi \)s, \( \Phi_{im} \) and \( \psi_{im} \) denote the \( m \)-th columns of \( \Phi_i \) and \( \psi_i \).

We can regard that this layer is equivalent to the fully connected layer where \( a s^2 \) neurons exists, the behaviors of those neurons are given by the \( \phi \)-s, and the outgoing weights are given by the \( \psi \)-s. Pruning the \( i \)-th channel in the convolutional layer is equivalent to pruning \( s^2 \) corresponding neurons in this converted form.

3.5 Relation to CP

CP [4] is also a layer-wise pruning method that conducts reconstruction with least squares method. Although, its strategy for neuron selection is different from ours. They select
the neurons to be pruned by solving the following Lasso Regression problem:

$$\beta^* = \arg\min_{\beta} \left\| Y - \sum_{i=1}^{d} \beta_i x_i w_i^\top \right\|_F^2 + \lambda \| \beta \|_1$$

subject to $\| \beta \|_0 \leq c'$, where $c'$ denotes the desired number of neurons and $\beta = (\beta_1, \ldots, \beta_e)^\top$ denotes a vector used for neuron selection. If $\beta_j = 0$, the $j$-th neuron can be pruned. Then, the reconstruction is done in the same way with ours.

The weak point of CP is that it selects the neurons to be pruned based on the error before reconstruction, which does not guarantee the minimal error after reconstruction. On the other hand, REAP selects the neurons to be pruned based on the reconstruction error. Because of this difference, REAP performs better than CP, as we will show in experiments.

4. Experiments

We conducted the experiments with several CNN models and datasets (VGG16 [22], ResNet-56 [24] on cifar-10 [25], and DenseNet-121 [26] on Stanford Dogs [27]) to verify REAP.

Even though REAP is an extended method of NU, we mainly compare REAP with CP, because REAP is the most similar with CP in theory, and CP is one of the state-of-the-art methods recently.

4.1 VGG16 on ImageNet

We conducted experiments with VGG16 on ImageNet. We pruned the convolutional layers until the theoretical speed-up ratio (the ratio of the floating point multiplications before and after the pruning) became $\times 2$ and $\times 5$. For pruning, we used 5000 images randomly selected from ImageNet training dataset. The images were resized so that the shorter side became 256, then $224 \times 224$ random crop was applied to the training images and $224 \times 224$ center crop was applied to the test images. The pruned models were fine-tuned for 10 epochs with the learning rate $10^{-5}$. The momentum was set to 0.9, the minibatch size was set to 128, and the dropout rate in the fully connected layers was set to 0.5. The rest of the setups, including the pruning ratio for each layer, were set to the same values with [4].

The results are shown in Table 1. REAP performs consistently better than the existing methods. In the accuracy after retraining, we marginally outperform the other methods at $\times 2$ speed-up. At $\times 5$ speed-up, the existing methods suffer even larger accuracy drop than we do.

An important observation is that we only suffer 9.4% accuracy drop at $\times 5$ speed-up before retraining. On the other hand, CP suffers 22.0% drop and ThiNet spoiled the model performance. This is because we use consistent strategy for channel selection and reconstruction to preserve the performances of the pruned models. As we show better performances before retraining, we can achieve higher accuracy after retraining as well. To put this observation differently, REAP enables us to achieve a certain accuracy with fewer epochs of retraining, which means that we can save time and labors for retraining by pruning with REAP.

It is also worth noting that the model pruned by REAP, at $\times 2$ speed-up, after retraining, is better than original VGG16. This is most likely because we removed the redundant weights, the remaining weights had smaller chance of being trapped in the local minima during the retraining.

(1) Close analysis on performance difference between CP and REAP.

As already mentioned, REAP is the most similar with CP in theory. The only difference of them is the channel selection criteria. As REAP selects the channels to be pruned in a greedy fashion, which probably raises the following questions:

1. Do the channels selected in REAP actually cause smaller reconstruction error than the channels selected by Lasso Regression in CP?
2. Although REAP takes an efficient algorithm for channel selection, does the computation finishes within reasonable time?

In order to answer these questions, we conducted additional experiments with VGG16 [22]. We pruned $conv1-1$, $conv2-1$, $conv3-1$, $conv4-1$’ with several pruning ratios, and observe the layer-wise reconstruction errors ($\|\Delta Y\|_F^2$) and measure the computational time spent on channel selection. For tuning the hyper-parameter in CP (the coefficient for Lasso regression), we use binary search algorithm as we found

| Speed-up ratio | Method   | Acc. before rt | Acc. after rt | Retrain epoch # |
|----------------|----------|----------------|---------------|-----------------|
|                | REAP     | -2.0%          | +0.2%         | 10              |
|                | NU [9]   | -5.0%          | -             | -               |
| ×2             | CP [4]   | -2.7%          | 0.0%          | 10              |
|                | ThiNet [5] | -65.0%        | -1.0%         | 10              |
|                | SPP [19] | -              | 0.0%          | -               |
| ×5             | REAP     | -9.4%          | -1.3%         | 10              |
|                | CP [4]   | -22.0%         | -1.7%         | 10              |
|                | ThiNet [5] | -88.8%        | -3.4%         | 10              |
|                | SPP [19] | -              | -2.0%         | -               |

*VGG16 has 4 blocks that have several convolutional layers, and $convX-Y$ represents the $Y$-th layer of the $X$-th block.
Fig. 3  Layer-wise analysis for VGG16 on ImageNet. The channels selected by REAP cause consistently smaller reconstruction error than the channels selected by CP.

Table 2  Time (sec.) spent for channel selection in Conv1-1, Conv2-1, Conv3-1 and Conv4-1 (channels# in the parentheses), at the pruning ratios of 0.25, 0.5, 0.75.

| Method | Conv1-1 (64) | Conv2-1 (128) | Conv3-1 (256) | Conv4-1 (512) |
|--------|--------------|---------------|---------------|---------------|
|        | 0.25 0.50 0.75 | 0.25 0.50 0.75 | 0.25 0.50 0.75 | 0.25 0.50 0.75 |
| REAP   | 21 22 24     | 35 44 53      | 116 168 229   | 777 1307 1872 |
| CP     | 90 84 83     | 117 130 120   | 209 226 220   | 385 434 418   |

Table 3  ResNet-56 on cifar-10. The changes of top-1 accuracy (baseline: 93.4%) are reported (The greater, the better.). *our implementation. **results taken from [4].

| Speed-up ratio | Method | Acc. before rt | Acc. after rt | Retrain epoch# |
|----------------|--------|----------------|---------------|----------------|
| ×2             | REAP   | -1.9%          | -0.5%         | 100            |
|                | NU[4]  | -13.3%         | -1.9%         | 10             |
|                | CP[4]  | -3.7% (*-2.0%) | -0.9% (*-1.0%)| 100            |
|                | *ThiNet[5] | -56.9%       | -1.9%         | 100            |
|                | DCP[6] | -              | -0.3%         | 400            |

out it was the fastest. All the methods are implemented with python3.6 and tested on Intel(R) Core(TM) i7-2600K CPU @ 3.40GHz.

As shown in Fig. 3, REAP suffers smaller error than CP. Besides, the trend is that the higher the pruning ratios are, the larger the error gaps between REAP and CP are. Because REAP selects channels based on the reconstruction error, we suffer consistently smaller reconstruction error than CP, despite we use greedy algorithm for channel selection.

Table 2 shows the results of computational time measurements. In conv4-1 that has 512 channels, REAP consumes more computational time than CP. However, REAP requires 1872 seconds at the pruning ratio of 0.75, which we think is acceptable enough, considering that REAP saves us time for retraining the pruned model and that the training typically takes much more time (e.g. 1 epoch takes over 8 hours on NVIDIA GeForce GTX 1080 Ti). When the number of channels is no greater than 256 (in conv1-1, conv2-1, and conv3-1), REAP is faster than or as fast as CP.

4.2 ResNet-56 on Cifar-10

We pruned the pretrained model taken from [28]. For pruning, we used 6400 randomly sampled training images. ResNet-56 has 27 residual units that have 2 sequential convolutional layers and a shortcut path. The pruning ratios in the first 9 units, the second 9 ones and the rest were set to 3 : 2 : 1. The pruned models were retrained for 100 epochs, beginning with the learning rate $10^{-2}$ and dividing it by 10 every 25 epochs. The rest of the training setups were aligned with [24]. Since we wanted to compare REAP with CP in the same conditions, we tried to evaluate CP on our own and put the results reported in [4] just for reference (because the pretrained model used in [4] is not available, and the experimental setups are not mentioned in [4]).

The results are shown in Table 3. Before retraining, we could easily outperform the existing methods. We suffer only 1.9% accuracy drop without retraining, while CP suffers 3.7%. After retraining, we are slightly worse than,
although competitive with another state-of-the-art pruning method DCP [6]. However, while DCP needs 400 epochs of retraining to achieve this result, we only need 100 epochs to achieve the competitive result. In this way, the fact that we can save efforts on retraining is a strength of REAP.

4.3 DenseNet-121 on Stanford Dogs

Finally, we conducted the experiments on DenseNet-121 fine-tuned with StanfordDogs dataset. For transfer learning, we set the learning rate to $10^{-2}$ for the first 30 epochs and set it to $10^{-3}$ for 20 more epochs. For retraining after pruning, we set the learning rate as $10^{-3}$ and trained the models for 10 epochs, then repeated another 10 epochs with the learning rate $10^{-4}$. We set the pruning ratios in Block1, Block2, Block3 and Block4 to 5 : 5 : 4 : 3. The rest of the setups were aligned with Sect. 4.1.

The results are shown in Table 4. Similarly with other experiments, REAP could preserve the model accuracy better than the other methods did. It is also remarkable that the model pruned by REAP without retraining is as accurate as the model after retraining. REAP preserves the model performances so well that we sometimes do not even need to retrain the pruned models.

Table 4 DenseNet-121 on Stanford Dogs. The changes of top-1 accuracy (baseline: 84.6%) are reported (The greater, the better). *our implementation.

| Speed-up ratio | Method | Acc. before rt | Acc. after rt | Retrain epoch# |
|----------------|--------|----------------|---------------|---------------|
| ×2             | REAP   | -3.1%          | -3.3%         | 20            |
|                | CP[4]  | -4.7%          | -3.5%         | 20            |
|                | ThiNet[5] | -63.7%     | -4.9%         | 20            |

5. Conclusion

We have proposed REAP, a channel pruning method to accelerate the inference of CNNs. REAP is an extension of the currently state-of-the-art pruning method CP. REAP prunes the channels based on the reconstruction error of the outputs, then reconstructs the outputs by the least squares method. Thus, REAP can reduce the computational complexity of the CNN models while maintaining their performances, which not only makes it possible to produce a fast, compact, and accurate models but also saves time and labors required for retraining the pruned models. On the experiments, we could confirm these strengths of REAP.

6. Future Works

As the nature of a layer-wise pruning method, the performance of REAP depends on the pruning ratio that is manually determined for each layer. We plan to extend REAP to overcome this weak point. It should be possible to determine the proper pruning ratio for a layer based on the reconstruction error (e.g. imposing a threshold on the maximum reconstruction error in each layer). Then, we can conduct holistic optimization with REAP and achieve even better pruning performances.

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Appendix A: Substitute approach for computing re-
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Algorithm 1

| Input: | $x_i \in \{1, \cdots, c\}$ and iterations $d$ |
| Output: | $r_i \in Z$ and $v_{ji}, j \in Z$ |
| Initialize: | $r_i \leftarrow x_i$ and $v_{ji} \leftarrow 0$ for each $i, j \in Z$ |
| while the number of iterations $\leq d$ do |
| for $(j, i) \in Z \times Z$ do |
| if $i \neq j$ then |
| $v_{ji} \leftarrow v_{ji} + f(r_j, x_i)$ |
| $r_j \leftarrow r_j - f(r_j, x_i)x_i$ |
| end if |
| end for |
| end while |

\[
r_j = x_j - \sum_{i \in Z \backslash \{j\}} v_{ji}^* x_i |
\]
\[
= x_j - M_{Z(\backslash j)} M_{Z(\backslash j)}^T x_j,
\]

where $M_{Z(\backslash j)}$ represents the orthogonal basis of $U_{Z(\backslash j)}$. However, $M_{Z(\backslash j)}$ is typically a large matrix, and computing $M_{Z(\backslash j)}$ for every possible $j$ is not memory-efficient. We need some more tricks to accelerate the computation.

We obtain the approximate solution of the $r$-s and the $v$-s by applying Gram-Schmidt process iteratively. Let $\leftarrow$ denote assignment and $f$ denote the function to compute the orthogonal projection coefficient:

\[
f(r, x) = \frac{\langle r, x \rangle }{||x||^2}.
\]

Then, the idea of this algorithm is illustrated in Fig. A.1 and can be described as follows.

- We first initialize: $r_j \leftarrow x_j$, $v_{ji} \leftarrow 0$ for each $i \in Z \backslash \{j\}$.
- We apply Gram-Schmidt process to make $r_j$ orthogonal to $x_i$ for each $i \in Z \backslash \{j\}$ alternately, such that $r_j \leftarrow r_j - f(r_j, x_i)x_i$. At the same time, we also update the coefficient $v_{ji}$ such that $v_{ji} \leftarrow v_{ji} + f(r_j, x_i)$.
- After repeating this sequential orthogonalizations many times, $r_j$ becomes a vector which is approximately orthogonal to all the vectors in $\{x_i | i \in Z \backslash \{j\}\}$, in other words, $r_j^\star$ becomes approximately orthogonal to $U_{Z(\backslash j)}$. Then, $\{v_{ji} | i \in Z \backslash \{j\}\}$ is the approximate solution of Eq. (8).

The detailed procedure is summarized in Algorithm 1.

The theoretical computational order of this algorithm is $O(c^2)$. However, it substantially reduces to $O(c^2)$ by parallel computing. In this algorithm, we do not need any large tensors except for the $x$-s, the $r$-s, and the $v$-s. In CUDA, they can be stored in the shared memory and passed to each thread by reference. Therefore, each thread suffers little memory consumption, and $r_j$ and $v_{ji}$ for $i, j \in Z$ can be computed parallelly.

Appendix A: Substitute approach for computing re-
construction error of each neuron

We try to accelerate the computation for (8) by parallel com-
puting. However, it is not smart to implement (8) as is, be-
cause the naive solution of the least squares problem is not
memory-efficient, and the degree of parallelism would be
limited because of the heavy memory requirement. There-
fore, we have developed a memory efficient algorithm that
uses Gram-Schmidt process.

As already mentioned, solving (8) is equivalent to com-
puting the orthogonal projection of $x_j$ onto the subspace
$U_{Z(\backslash j)}$ spanned by $\{x_i | i \in Z \backslash \{j\}\}$, as shown in Fig. 2. Then, we have
Fig. A-1  An example of the Gram-Schmidt process based algorithm. We want to obtain $r_j$, the residual of the projection of $x_j$ onto the subspace spanned by $x_k$ and $x_l$. a) $r_j$ is initialized to $x_j$. b) $r_j$ is projected onto $x_l$ and is replaced by the residual. c) $r_j$ is projected onto $x_k$ and is replaced by the residual. Note that $r_j$ becomes NOT orthogonal to $x_l$ again, as $x_k$ and $x_l$ are not orthogonal. d) After repeating b) and c) lots of times, $r_j$ converges into the vector that is orthogonal to both of $x_k$ and $x_l$.

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