Group Whitening: Balancing Learning Efficiency and Representational Capacity

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Abstract

Batch normalization (BN) is an important technique commonly incorporated into deep learning models to perform standardization within mini-batches. The merits of BN in improving model’s learning efficiency can be further amplified by applying whitening, while its drawbacks in estimating population statistics for inference can be avoided through group normalization (GN). This paper proposes group whitening (GW), which elaborately exploits the advantages of the whitening operation and avoids the disadvantages of normalization within mini-batches. Specifically, GW divides the neurons of a sample into groups for standardization, like GN, and then further decorrelates the groups. In addition, we quantitatively analyze the constraint imposed by normalization, and show how the batch size (group number) affects the performance of batch (group) normalized networks, from the perspective of model’s representational capacity. This analysis provides theoretical guidance for applying GW in practice. Finally, we apply the proposed GW to ResNet and ResNeXt architectures and conduct experiments on the ImageNet and COCO benchmarks. Results show that GW consistently improves the performance of different architectures, with absolute gains of 1.02% ~ 1.49% in top-1 accuracy on ImageNet and 1.82% ~ 3.21% in bounding box AP on COCO.

1 Introduction

Batch normalization (BN) [20] represents a milestone technique in deep learning [13, 45, 51], and has been extensively used in various network architectures [13, 45, 56, 44, 15]. BN standardizes the activations within a mini-batch of data, which improves the conditioning of optimization and accelerates training [20, 3, 40]. The stochasticity of normalization introduced along the batch dimension is believed to benefit generalization [51, 41, 18]. However, this stochasticity also results in differences between the training distribution (using mini-batch statistics) and the test distribution (using estimated population statistics) [19], which is believed to be the main cause of BN’s small-batch-size problem — BN’s error increases rapidly as the batch size becomes smaller [51]. To address this issue, a number of approaches have been proposed [51, 37, 32, 19, 48, 43]. One representative method is group normalization (GN), which divides the neurons into groups and then performs the standardization operation over the neurons of each group, for each sample, independently. GN provides a flexible solution to avoid normalization along the batch dimension, and benefits on visual tasks limited to small-batch-size training [51].

As a widely used operation in data pre-processing, whitening not only standardizes but also decorrelates the data [26], which further improves the conditioning of optimization problem [26, 50, 16]. A whitened input has also been shown to make the gradient descent updates similar to the Newton updates for linear models [26, 50, 16]. Motivated by this, Huang et al. [16] proposed batch whitening (BW) for deep models, which performs whitening on the activations of each layer within a mini-batch. BW has been shown to achieve better optimization efficiency and generalization than BN [16, 18, 55]. However, BW further amplifies the disadvantage of BN in estimating the population statistics, where
the number of parameters to be estimated with BW is quadratic to the number of neurons/channels. Thus, BW requires a sufficiently large batch-size to work well.

To exploit whitening’s advantage in optimization, while avoiding its disadvantage in normalization along the batch dimension, this paper proposes group whitening (GW). For each sample, GW divides the neurons into groups for standardization over the neurons of each group (like GN), and then further decorrelates the groups. Unlike BW, GW has stable performance for a wide range of batch sizes, like GN, and thus can be applied to a variety of tasks. GW further improves the conditioning of optimization of GN with its whitening operation.

One important hyperparameter of GW is the group number. We observe that GW/GN has a significantly degenerated training performance when the group number is large, which is similar to the small-batch-size problem of BW/BN. We attribute this to the constraints on the output imposed by the normalization operation, which affect the model’s representational capacity. As such, this paper defines the constraint number of normalization (as will be discussed in Section C) to quantitatively measure the magnitude of the constraints provided by normalization methods. With the support of the constraint number, we analyze how the batch size (group number) affects the model’s representational capacity for batch (group) normalized networks. Our analysis presents a new viewpoint for understanding the small-batch-size problem of BN.

We apply the proposed GW to two representative deep network architectures (ResNet [13] and ResNeXt [52]) for ImageNet classification [39] and COCO object detection and instance segmentation [51]. GW consistently improves the performance for both architectures, with absolute gains of 1.02% ~1.49% in top-1 accuracy for ImageNet and 1.82% ~3.21% in bounding box AP for COCO.

2 Preliminaries

To simplify the discussion, we first consider the d-dimensional input vector x, which will be generalized to a convolutional input in the subsequent section. Let $X \in \mathbb{R}^{d \times m}$ be a data matrix denoting the mini-batch input of size $m$ in a given layer.

**Standardization.** During training, batch normalization (BN) [20] standardizes the layer input within a mini-batch, for each neuron, as\(^1\)

$$\hat{X} = \phi_{BN}(X) = \Lambda_d^{-\frac{1}{2}} (X - \mu_d 1^T). \quad (1)$$

Here, $\mu_d = \frac{1}{m} X 1$ and $\Lambda_d = \text{diag}(\sigma_1^2, \ldots, \sigma_d^2) + \epsilon I$, where $\sigma_i^2$ is the variance over mini-batches for the $i$-th neuron, $1$ is a column vector of all ones, and $\epsilon > 0$ is a small number to prevent numerical instability. During inference, the population statistics $\{\hat{\Lambda}_d^{-\frac{1}{2}}, \hat{\mu}_d\}$ are required for deterministic inference, and they are usually calculated by running average over the training iterations, as follows:

$$\begin{aligned}
\hat{\mu}_d &= (1 - \lambda) \mu_d + \lambda \mu_{td}, \\
\hat{\Lambda}_d^{-\frac{1}{2}} &= (1 - \lambda) \hat{\Lambda}_d^{-\frac{1}{2}} + \lambda \Lambda_d^{-\frac{1}{2}}.
\end{aligned} \quad (2)$$

Such an estimation process can limit BN’s usage in recurrent neural networks [25] [9], or harm the performance for small-batch-size training [19] [51].

To avoid the estimation of population statistics shown in Eqn. 2, Ba et al. proposed layer normalization (LN) [3] to standardize the layer input within the neurons for each training sample, as:

$$\hat{X} = \phi_{LN}(X) = (X - 1 \mu_m^T) \Lambda_m^{-\frac{1}{2}}. \quad (3)$$

Here, $\mu_m = \frac{1}{m} X^T 1$ and $\Lambda_m = \text{diag}(\sigma_1^2, \ldots, \sigma_m^2) + \epsilon I$, where $\sigma_i^2$ is the variance over the neurons for the $i$-th sample. LN has the same formulation during training and inference, and is extensively used in natural language processing tasks [47] [55] [53], since it does not normalize within a mini-batch.

Group normalization (GN) [51] further generalizes LN, dividing the neurons into groups and performing the standardization within the neurons of each group independently, for each sample. Specifically, defining the group division operation as $\Pi : \mathbb{R}^{d \times m} \mapsto \mathbb{R}^{c \times gm}$, where $g$ is the group number and $d = gc$, GN can be represented as follows:

$$\hat{X} = \phi_{GN}(X; g) = \Pi^{-1}(\phi_{LN} (\Pi(X))). \quad (4)$$

\(^1\)BN and other normalization methods discussed in this paper all use extra learnable dimension-wise scale and shift parameters [20].
We propose group whitening (GW). Given a sample \( \mathbf{x} \in \mathbb{R}^d \), GW performs normalization as:

**Group division:**

\[
\mathbf{X}_G = \Pi(\mathbf{x}; g) \in \mathbb{R}^{g \times c},
\]

**Whitening:**

\[
\hat{\mathbf{x}}_G = \phi^W(\mathbf{X}_G) = \Sigma_d^{-\frac{1}{2}}(\mathbf{X}_G - \mu_d 1^T),
\]

**Inverse group division:**

\[
\hat{\mathbf{x}} = \Pi^{-1}(\hat{\mathbf{x}}_G) \in \mathbb{R}^d.
\]
Here, we define the constraint number $\Upsilon$ as an example, we have $9$ (Eqn. 6), GW costs $2$. The normalization operation can also be regarded as a way to find a solution $\hat{X}$, which should improve the conditioning, like BW, and benefit training. GW avoids normalization along the batch dimension, and it works stably across a wide range of batch sizes (Figure 1).

**Convolutional layer.** For the convolutional input $X \in \mathbb{R}^{d \times m \times H \times W}$, where $H$ and $W$ are the height and width of the feature maps, BN and BW consider each spatial position in a feature map as a sample and normalize over the unrolled input $X \in \mathbb{R}^{d \times m \times HW}$. In contrast, LN and GN view each spatial position in a feature map as a neuron and normalize over the unrolled input $X \in \mathbb{R}^{dHW \times m}$. Following GN, GW also views each spatial position as a neuron, i.e., GW operations (Eqs. 23, 24 and 25) are performed for each sample with unrolled input $x \in \mathbb{R}^{dHW}$.

**Computational complexity.** For a convolutional mini-batch input $X \in \mathbb{R}^{d \times m \times H \times W}$, GW using ZCA whitening (Eqn. 5) costs $2mHWd^g + mO(g^3)$. Using the more efficient ‘ItN’ operation (Eqn. 6), GW costs $2mHWd^g + mTg^3$, where $T$ is the iteration number, while the $3 \times 3$ convolution with the same input and output feature maps costs $9mHWd^2$. The relative cost of GW for a $3 \times 3$ convolution is $2g/(9d) + Tg^3/(9HWd^2)$.

**Difference from group-based BW.** Our method is significantly different from the group-based BW, in which the whitening operation is also performed within mini-batch data. Specifically, group-based BW has difficulty in estimating the population statistics, as discussed in Section 2. Note that group-based BW is reduced to BN if the channel number in each group $c = 1$, while GW is reduced to GN if the group number $g = 1$.

## 4 Constraint Analysis for Normalization

The normalization operation ensures that the normalized output $\hat{X} = \phi(X) \in \mathbb{R}^{d \times m}$ has a stable distribution. This stable distribution can be implicitly viewed as the constraints imposed on $\hat{X}$, which can be represented as a system of equations $T_\phi(\hat{X})$. For example, BN provides the constraints $T_{\phi,BN}(\hat{X})$ as:

$$
\sum_{j=1}^{m} \hat{X}_{ij} = 0 \text{ and } \sum_{j=1}^{m} \hat{X}_{ij}^2 = m, \text{ for each neuron } i = 1, \ldots, d. (11)
$$

Here, we define the constraint number of normalization to quantitatively measure the magnitude of the constraints provided by the normalization method.

**Definition 1** Given the input data $X \in \mathbb{R}^{d \times m}$, the constraint number of a normalization operation $\phi(\cdot)$, referred to as $\zeta(\phi; X)$, is the number of independent equations in $T_\phi(\hat{X})$.

As an example, we have $\zeta(\phi_{BN}; X) = 2d$ from Eqn. 11. Table 1 summarizes the constraint numbers of the main normalization methods discussed in this paper (please refer to the Appendix C for derivation details). We can see that the whitening operation provides significantly stronger constraints than the standardization operation.

The normalization operation can also be regarded as a way to find a solution $\hat{X}$ satisfying the constraints $T_\phi(\hat{X})$. To ensure the solution is feasible, it must satisfy the following condition:

$$
\zeta(\phi; X) \leq \chi(\hat{X}), (12)
$$

where $\chi(\hat{X}) = md$ is the number of variables in $\hat{X}$. Based on Eqn. 12, we have $m \geq 2$ for BN to ensure a feasible solution. We also provide the ranges of batch size/group number for other normalization methods in Table 1. Note that the batch size $m$ should be larger than/equal to $d$ to achieve a numerically stable solution for BW when using ZCA whitening in practice [16]. This also applies to GW, where $g$ should be less than/equal to $\sqrt{d}$.

## 4.1 Analysis on Representational Capacity

It is believed that the constraints introduced by normalization affect the representational capacity of neural networks [20], while the batch size of the optimization algorithm significantly affects the performance of batch normalized networks [51, 18]. Here, we provide a unified analysis based on the
constraint number of normalization, and show how the batch size $m$ of the optimization algorithm affects the representational capacity of the model using BN/BW, but not GN/GW. Our analysis is based on the following assumption.

**Assumption 1** The constraint number of the normalization and the representational capacity of the normalized model have a negative correlation.

### Batch normalized networks.
Given training data $\mathcal{D}$ of size $N$, we consider the optimization algorithm with batch size $m$ (we assume $N$ is divisible by $m$). We calculate the constraint number of normalization over the entire training data $\zeta(\phi; \mathcal{D})$. The results for different normalization methods are shown in Table 1. We find that $\zeta(\phi_{BN}; \mathcal{D})$ or $\zeta(\phi_{BW}; \mathcal{D})$ is inversely proportional to $m$, which suggests that the representational capacity of batch normalized networks decreases with decreasing batch size, based on Assumption 1. For example, the normalized outputs of BN are constrained to be $(1, -1)$ or $(-1, 1)$ when $m = 2$, which heavily reduces the representational capacity of the model and results in significantly degenerated training performance, as shown in Figure 1. To the best of our knowledge, our analysis is the first to show how the batch size of an optimization algorithm affects the model’s representational capacity for batch normalized networks.

### Group normalized networks.
We also observe that $\zeta(\phi_{GN}; \mathcal{D})$ or $\zeta(\phi_{GW}; \mathcal{D})$ is not related to $m$, which demonstrates that the batch size of an optimization algorithm does not affect the model’s representational capacity for group normalized networks. Our analysis provides a new understanding of why GN is not sensitive to batch size [31], from the perspective of a model’s representational capacity. Although unrelated to $m$, $\zeta(\phi_{GN}; \mathcal{D})$ and $\zeta(\phi_{GW}; \mathcal{D})$ are positively proportional to $g$, which suggests that the representational capacity of group normalized networks will decrease with increasing group number, according to Assumption 1. Note that a large group number contributes to the distribution stability of the normalized output, which may benefit training. Therefore, there exists a trade-off between the reduced model representational capacity and the increased learning efficiency, when increasing the group number.

We conduct experiments on MNIST with random labels [58]. The results are shown in Figure 2. We observe that the model with GN/GW has significantly degenerated training accuracy when $g$ is too large, which means that a large group number heavily limits the model’s representational capacity. We note that GW is more sensitive to the group number than GN. The main reason is that $\zeta(\phi_{GW}; \mathcal{D})$ is quadratic to $g$, while $\zeta(\phi_{GN}; \mathcal{D})$ is linear to it, from Table 1. We also observe that the best training accuracy of GW is higher than that of GN (85.80% vs. 81.40%). We attribute this to the fact that the whitening operation is better for improving the conditioning of optimization, compared to standardization.

### 4.2 Discussion of Previous Work

Previous analyses on BN are mainly derived from the perspective of optimization [40, 29, 24, 6]. One argument is that BN can improve the conditioning of the optimization problem [40, 6, 11, 22, 10], either by avoiding the rank collapse of pre-activation matrices [10] or alleviating the pathological sharpness of the landscape [40, 22]. This argument has been further investigated by computing the spectrum of the Hessian for a large-scale dataset [11]. The improved conditioning enables large learning rates, thus improving the generalization [10, 33]. Another argument is that BN is scale invariant [20, 3], enabling it to adaptively adjust the learning rate [1, 6, 59, 28], which stabilizes and further accelerates training [20, 3]. Other analyses focus on investigating the signal and gradient propagation, either by exploiting mean-field theory [54, 49], or a neural tangent kernel (NTK) [21].

Different from these works, we are the first to investigate how BN/GN affects a model’s representational capacity, which opens new doors in analyzing and understanding normalization methods. We

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2This assumption is intuitively reasonable, since the constraint number indicates the constraints (represented by equations) imposed on the normalized output in neural networks. The representational capacity of the model should be reduced, when provided with more constraints.
Figure 2: Effects of group number for group normalized networks. We train a four-layer MLP with 1280 neurons in each layer for MNIST, with random labels [58]. We train the model for 100 epochs with a batch size of 16. We vary the group number of GN/GW and evaluate the training accuracy.

Figure 3: Effects of group number of GW/GN on ResNet-50 for ImageNet classification. We evaluate the top-1 training and validation accuracies.

Further investigate how batch size affects the training performance of batch normalized networks (Figure 1), from the perspective of a model’s representational capacity. Several works [41, 18, 17] have shown that batch size is related to the magnitude of stochasticity [2, 46] introduced by BN, which also affects the model’s training performance. However, the stochasticity analysis [18] is specific to normalization along the batch dimension, and cannot explain why GN with a large group number has significantly worse performance (Figure 2), while our work provides a unified analysis for batch and group normalized networks.

5 Experiments on Large-scale Visual Recognition Tasks

We investigate the effectiveness of our proposed GW on large-scale ImageNet classification [39], as well as COCO object detection and segmentation [31]. We use the more efficient and numerically stable ‘ItN’ (with $T = 5$) [18] to calculate the whitening matrix for both GW and BW, in all experiments. Our implementation is based on PyTorch [36].

5.1 ImageNet Classification

We experiment on the ImageNet dataset with 1000 classes [39]. We use the official 1.28M training images as a training set, and evaluate the top-1 accuracy on a single-crop of 224x224 pixels in the validation set with 50k images. We investigate the ResNet [13] and ResNeXt [52] models.

5.1.1 Ablation Study on ResNet-50

We follow the same experimental setup as described in [13], except that we use two GPUs and train over 100 epochs. We apply SGD with a mini-batch size of 256, momentum of 0.9 and weight decay of 0.0001. The initial learning rate is set to 0.1 and divided by 10 at 30, 60 and 90 epochs. Our baseline is the 50-layer ResNet (ResNet-50) trained with BN [20].

Effects of group number. We investigate the effects of group number for GW/GN, which we use to replace the BN of ResNet-50. We vary the group number $g$ ranging in $\{8, 16, 32, 64, 128\}$ (we use the channel number if it is less than the group number in a given layer), and report the training and validation accuracies in Figure 5. We can see GW has consistent improvement over GN in training accuracy, across all values of $g$, which indicates the advantage of the whitening operation.
We also perform experiments using BW [18] and GW after the last average pooling (before the full connection layer). We also investigate the effect of inserting a GW/BW layer after the last average pooling (before the full connection layer). We believe there is a trade-off between GW and BN, in terms of computational cost vs. performance compared to BW [17]). Our baselines are the original networks trained with BN, and we also provide the results trained with GN.

### Table 2: Effects of position when applying GW on ResNet-50 for ImageNet classification.

| Method | S1-B1 | S1-B2 | S1-B3 | S1-B12 |
|--------|-------|-------|-------|--------|
| Baseline (BN) | 76.23 | 76.23 | 76.23 | 76.23 |
| GW | 76.76 | 77.62 | 77.72 | 77.45 |

Note that our implementations are based on the APIs provided by PyTorch and are not finely optimized. For more discussion on time costs, please refer to the Appendix D.

### Table 3: Comparison of validation accuracy on ResNets [13] and ResNeXts [52] for ImageNet. Note that we use an additional layer for BWΣ to learn the decorrelated feature, as recommended in [17].

| Method | ResNet-50 | ResNet-101 | ResNetXt-50 | ResNetXt-101 |
|--------|-----------|------------|-------------|-------------|
| Baseline (BN) | 76.23 | 77.69 | 77.01 | 79.29 |
| GN | 75.71 (10.52) | 77.20 (10.49) | 75.69 (11.32) | 78.00 (11.29) |
| BWΣ | 77.21 (10.98) | 78.27 (10.58) | 77.29 (10.28) | 79.43 (10.14) |
| GW | 77.72 (11.49) | 78.71 (11.02) | 78.43 (11.42) | 80.43 (11.14) |

Here, we investigate the position at which to apply GW (g=64) on ResNet-50. ResNet and ResNeXt are both mainly composed of a stem layer and multiple bottleneck blocks [13]. We consider: 1) replacing the BN in the stem layer with GW (referred to as ‘S1’); 2) replacing the 1st, 2nd, 1st & 2nd, and 3rd BNs in all the bottleneck blocks, which are referred to as ‘B1’, ‘B2’, ‘B12’ and ‘B3’, respectively. We investigate five architectures, S1, S1-B1, S1-B2, S1-B3 and S1-B12, which have 1, 17, 17, 17 and 33 GW modules, respectively. We also perform experiments using BW [18] and BWΣ (employing a covariance matrix to estimate the population statistics of BW) for contrast.

We report the results in Table 2. GW/BWΣ improve the BN counterpart on all architectures by a clear margin, which demonstrates the advantage of the whitening operation over standardization [18]. GW provides significant improvements over BW/BWΣ on S1-B1, S1-B2, S1-B3 and S1-B12 (an absolute improvement of 0.9% on average). We attribute this to the advantage of GW in avoiding the estimation of population statistics. We also observe that GW has a slightly worse performance on S1-B12 than on S1-B1/S1-B2. We believe there is a trade-off between GW and BN, in terms of affecting the model’s representational capacity, optimization efficiency and generalization.

We also investigate the effect of inserting a GW/BW layer after the last average pooling (before the last linear layer) to learn the decorrelated feature representations, as proposed in [18]. This can slightly improve the performance (0.10% on average) when using GW, though the net gain is smaller than using BW (0.22%) or BWΣ (0.43%). Please refer to the Appendix D for details.

### 5.1.2 Validation on Larger Models

In this section, we further validate the effectiveness of GW on ResNet-101 [13], ResNeXt-50 and ResNeXt-101 [52]. We apply GW (g=64) in these models following the S1-B2 architecture, which achieves the best performance (Table 2) without significantly increasing the computational cost (it is only increased by roughly 23%). For comparison, we also apply BWΣ following the ‘S1-B2’ architecture, combining the learning of decorrelated features [17] (BWΣ has a slightly improved performance compared to BW [17]). Our baselines are the original networks trained with BN, and we also provide the results trained with GN.

The results are shown in Table 3. We can see that 1) our method improves the baseline (BN) by a significant margin (between 1.02% and 1.49%); 2) BWΣ has consistently better performance than BN.

Note that our implementations are based on the APIs provided by PyTorch and are not finely optimized. For more discussion on time costs, please refer to the Appendix D.
Table 4: Detection results (%) on COCO using the Faster R-CNN framework implemented in [34]. We use ResNet-50 as the backbone, combined with FPN. All models are trained by 1x lr scheduling (90k iterations), with a batch size of 16 on eight GPUs.

| Method | APbbox 50 | APbbox 75 | APbbox 75 train | APbbox 50 | APbbox 75 | APbbox 75 train |
|--------|-----------|-----------|-----------------|-----------|-----------|-----------------|
| BN†    | 42.24     | 63.00     | 46.19           | 37.53     | 59.82     | 39.96           |
| GN     | 42.18(0.06) | 63.22(0.22) | 46.00(0.19)     | 37.54(0.01) | 60.18(0.36) | 39.99(0.03)    |
| GW     | 44.41(1.17) | 65.36(2.96) | 48.67(2.48)     | 39.17(1.64) | 62.13(3.21) | 41.95(1.99)    |

Table 5: Detection and segmentation results (%) on COCO using the Mask R-CNN framework implemented in [34]. We use ResNeXt-101 as the backbone, combined with FPN. All models are trained by 2x lr scheduling (180k iterations), with a batch size of 8 on eight GPUs.

| Method | APbbox 50 | APbbox 75 | APbbox 75 train | APbbox 50 | APbbox 75 | APbbox 75 train |
|--------|-----------|-----------|-----------------|-----------|-----------|-----------------|
| BN†    | 36.31     | 58.39     | 38.83           | 36.39     | 57.22     | 39.56           |
| GN     | 36.62(0.31) | 58.91(0.52) | 39.32(0.49)     | 37.86(1.47) | 58.96(1.74) | 40.76(1.20)    |
| GW     | 38.13(1.82) | 60.63(2.24) | 41.08(2.25)     | 39.60(1.23) | 61.12(3.90) | 43.25(3.69)    |

but the net gain is reduced on wider networks (ResNeXt-50 and ResNeXt-101), which is probably caused by the difficulty in estimating the population statistics.

5.2 Object Detection and Segmentation on COCO

We fine-tune the models trained on ImageNet for object detection and segmentation on the COCO benchmark [31]. We experiment on the Faster R-CNN [38] and Mask R-CNN [12] frameworks using the publicly available codebase ‘maskrcnn-benchmark’ [34]. We train the models on the COCO train2017 set and evaluate on the COCO val2017 set. We report the standard COCO metrics of average precision (AP), AP50, and AP75, for bounding box detection (APbbox) and instance segmentation (APmask) [31]. For BN, we use its frozen version (indicated by $BN^\dagger$) when fine-tuning for object detection [51].

Results on Faster R-CNN. For the Faster R-CNN framework, we use the ResNet-50 models pre-trained on ImageNet (Table 4) as the backbones, combined with the Feature Pyramid Network (FPN) [40]. We consider two setups: 1) we use the box head consisting of two fully-connected layers (‘2fc’) without a normalization layer, as proposed in [30]; 2) following [51], we replace the ‘2fc’ box head with ‘4conv1fc’, which can better leverage GN, and apply GN/GW to the FPN and box head. We use the default hyperparameter configurations from the training scripts provided by the codebase [34] for Faster R-CNN. The results are reported in Table 4. The GW pre-trained model improves $BN^\dagger$ and GN by 1.82% and 1.51% AP, respectively. By adding GW/GN to the FPN and ‘4conv1fc’ head box, GW improves $BN^\dagger$ and GN by 3.21% and 1.74% AP, respectively.

Results on Mask R-CNN. For the Mask R-CNN framework, we use the ResNeXt-101 [52] models pre-trained on ImageNets (Table 3) as the backbones combined with FPN. We use the ‘4conv1fc’ box head, and apply GN/GW to the FPN, box head and mask head. We again use the default hyperparameter configurations from the training scripts provided by the codebase for Mask R-CNN [34]. The results are shown in Table 5. GW achieves 44.41% in box AP and 39.17% in mask AP, an improvement over $BN^\dagger$ of 2.17% and 1.64%, respectively.

6 Conclusion and Further Work

In this paper, we proposed group whitening (GW), which combines the advantages of normalization within a group of channels and the whitening operation. The effectiveness of GW was validated on large-scale visual recognition tasks. Furthermore, we also provided a constraint analysis for normalization methods, enabling further understanding on how the batch size (group number) affects the performance of batch (group) normalized networks from the perspective of representational capacity. This constraint analysis can provide theoretical guidance for applying GW and other normalization methods in practice.

In the future, it would be interesting to investigate normalization along other dimensions (e.g., positional normalization [27] and divisive normalization [37]) or other normalization operations (e.g., scaling only [7,57]), using our constraint analysis. We hope our analysis will provide a new means of understanding the behaviors of normalization methods.
**Broader Impact**

The proposed group whitening method could be applied to better train deep neural networks (DNNs), since it achieves high performance on visual recognition tasks, as validated by our experiments. Further, the proposed constraint analysis provides a new tool for analyzing and understanding normalization methods, which may contribute to the design of new DNN architectures and better understanding of DNN behaviors. As commonly acknowledged, DNNs are essential tools in a wide range of applications, e.g., computer vision tasks and natural language process tasks. We thus believe the work in this paper will have a broad impact for researchers and engineers in these fields in particular. Finally, while our work may bring many important benefits, we also recognize that it could be leveraged by someone to carry out the research/projects that violate ethical standards, e.g., generating biased or offensive texts and images using deep generative models, and detecting and tracking people without permission using deep discriminative models.

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Algorithm 1 The forward pass of group whitening.

1: **Input:** a input sample \( \mathbf{x} \in \mathbb{R}^d \).
2: **Hyperparameters:** \( \epsilon \), group number \( g \).
3: **Output:** \( \hat{\mathbf{x}} \in \mathbb{R}^d \).
4: Group division: \( \mathbf{X}_G = \Pi(\mathbf{x}; g) \in \mathbb{R}^{g \times \epsilon} \).
5: \( \mu = \frac{1}{G} \mathbf{X}_G 1 \).
6: \( \mathbf{X}_C = \mathbf{X}_G - \mu 1^T \).
7: \( \Sigma = \frac{1}{2} \mathbf{X}_C \mathbf{X}_C^T + \epsilon I \).
8: Calculate whitening matrix: \( \Sigma^{-\frac{1}{2}} = \psi^f(\Sigma) \).
9: \( \hat{\mathbf{X}}_G = \Sigma^{-\frac{1}{2}} \mathbf{X}_C \).
10: Inverse group division: \( \hat{\mathbf{x}} = \Pi^{-1}(\hat{\mathbf{X}}_G) \in \mathbb{R}^d \).

Algorithm 2 The corresponding backward pass of Algorithm 1.

1: **Input:** gradient of a sample: \( \frac{\partial \mathcal{L}}{\partial \mathbf{x}} \in \mathbb{R}^d \), and auxiliary data from respective forward pass: (1) \( \mathbf{X}_G \); (2) \( \Sigma^{-\frac{1}{2}} \).
2: **Output:** gradient with respect to the input: \( \frac{\partial \mathcal{L}}{\partial \mathbf{x}} \in \mathbb{R}^d \).
3: Group division: \( \frac{\partial \mathcal{L}}{\partial \mathbf{X}_G} = \Pi(\frac{\partial \mathcal{L}}{\partial \mathbf{x}}; g) \in \mathbb{R}^{g \times \epsilon} \).
4: \( \frac{\partial \mathcal{L}}{\partial \Sigma^{-\frac{1}{2}}} = \frac{\partial \mathcal{L}}{\partial \mathbf{X}_G} \mathbf{X}_C^T \).
5: Calculate gradient with respect to the covariance matrix: \( \frac{\partial \mathcal{L}}{\partial \Sigma} = \psi^b(\frac{\partial \mathcal{L}}{\partial \Sigma^{-\frac{1}{2}}} \) as follows:
6: \( f = \frac{1}{\epsilon} \frac{\partial \mathcal{L}}{\partial \mathbf{X}_G} 1 \).
7: \( \frac{\partial \mathcal{L}}{\partial \mathbf{X}_G} \Sigma^{-\frac{1}{2}} (\frac{\partial \mathcal{L}}{\partial \mathbf{X}_G} - f 1^T) + \frac{1}{2}(\frac{\partial \mathcal{L}}{\partial \Sigma} + \frac{\partial \mathcal{L}}{\partial \Sigma}^T) \mathbf{X}_C \).
8: Inverse group division: \( \frac{\partial \mathcal{L}}{\partial \mathbf{x}} = \Pi^{-1}(\frac{\partial \mathcal{L}}{\partial \mathbf{X}_G}) \in \mathbb{R}^d \).

A Algorithms

The forward pass of the proposed group whitening (GW) method is shown in Algorithm 1 and its corresponding backward pass is shown in Algorithm 2. Note that we need to specify the method to calculate the whitening matrix \( \Sigma^{-\frac{1}{2}} \psi^f(\Sigma) \) in Line 8 of Algorithm 1 as well as its backward operation \( \frac{\partial \mathcal{L}}{\partial \Sigma} = \psi^b(\frac{\partial \mathcal{L}}{\partial \Sigma^{-\frac{1}{2}}} \) shown in Line 5 of Algorithm 2. As stated in the submitted paper, we use zero-phase component analysis (ZCA) whitening and its efficient approximation by Newton’s iteration (‘ItN’) [18]. Here, we provide the details.

**ZCA whitening.** ZCA whitening [16] calculates the whitening matrix by eigen decomposition as:
\[ \Sigma^{-\frac{1}{2}} = \psi^{ZCA}(\Sigma) = \Lambda^{-\frac{1}{2}} \mathbf{D}^T, \]
where \( \Lambda = \text{diag}(\sigma_1, \ldots, \sigma_d) \) and \( \mathbf{D} = [d_1, \ldots, d_d] \) are the eigenvalues and associated eigenvectors of \( \Sigma \), i.e. \( \Sigma = \mathbf{D} \Lambda \mathbf{D}^T \).

The corresponding backward operation \( \frac{\partial \mathcal{L}}{\partial \Sigma} = \psi^{ZCA}_b(\frac{\partial \mathcal{L}}{\partial \Sigma^{-\frac{1}{2}}} \) is as follows:
\[
\begin{align*}
\frac{\partial \mathcal{L}}{\partial \Lambda} &= \mathbf{D}^T \left( \frac{\partial \mathcal{L}}{\partial \Sigma^{-\frac{1}{2}}} \right) \mathbf{D} \left( -\frac{1}{2} \Lambda^{-3/2} \right) \\
\frac{\partial \mathcal{L}}{\partial \mathbf{D}} &= \left( \frac{\partial \mathcal{L}}{\partial \Sigma^{-\frac{1}{2}}} \right) \mathbf{D} \Lambda^{-1/2} \\
\frac{\partial \mathcal{L}}{\partial \Sigma} &= \mathbf{D} \left( \mathbf{K}^T \odot \left( \mathbf{D}^T \frac{\partial \mathcal{L}}{\partial \mathbf{D}} \mathbf{K} \right) + \left( \frac{\partial \mathcal{L}}{\partial \Lambda} \right) \right) \mathbf{D}^T,
\end{align*}
\]
where \( (\frac{\partial \mathcal{L}}{\partial \Lambda})_{\text{diag}} \) sets the off-diagonal elements of \( \frac{\partial \mathcal{L}}{\partial \Lambda} \) as zero.

**‘ItN’ whitening.** ‘ItN’ whitening [18] calculates the whitening matrix by Newton’s iteration as:
\[
\Sigma^{-\frac{1}{2}} = \psi^{\text{ItN}}(\Sigma) = \sqrt{\text{tr}(\Sigma)} \mathbf{P}_T,
\]
where \( \text{tr}(\Sigma) \) indicates the trace of \( \Sigma \) and \( \mathbf{P}_T \) is calculated iteratively as:
\[
\begin{cases}
\mathbf{P}_0 = \mathbf{I} \\
\mathbf{P}_k = \frac{1}{2} (3 \mathbf{P}_{k-1} - \mathbf{P}_{k-1}^3 \Sigma_N^k), & k = 1, 2, \ldots, T.
\end{cases}
\]

For GW, we also use the extra learnable dimension-wise scale and shift parameters, like BN [20]. We omit this in the algorithms for simplicity.
def GroupWhitening(X, gamma, beta, g, T=5, eps=1e-5):
    # X input feature with size [m, d] or [m, d, H, W]
    # gamma, beta: the learnable affine
    # g: the group number of group whitening
    # T: the iteration number of Newton's iteration
    size = X.size()
    X_G = X.view( size[0], g, -1)  # group division
    m, g, c = X_G.size()
    # centering
    mean = X_G.mean(-1, keepdim=True)
    X_G_mean = X_G - mean
    # approximate ZCA whitening by Newton's iteration
    P = [torch.Tensor([]) for _ in range(T+1)]
    sigma = X_G_mean.matmul(X_G_mean.transpose(1, 2)) / c
    P[0] = torch.eye(d).to(x).expand(sigma.shape)
    M_zero = sigma.clone().fill_(0)
    trace_inv = torch.addcmul(M_zero, sigma, P[0]).sum((1, 2), keepdim=True).reciprocal_()
    sigma_N = torch.addcmul(M_zero, sigma, trace_inv)
    for k in range(T):
        P[k+1] = torch.baddbmm(1.5, P[k], -0.5, torch.matrix_power(P[k], 3), sigma_N)
    wm = torch.addcmul(M_zero, P[T], trace_inv.sqrt())
    y = wm.matmul(X_G_mean)
    output = y.view_as(X)  # inverse group division
    return output * gamma + beta

Figure 4: Python code of GW using ItN whitening, based on PyTorch.

Here, $\Sigma_d^N = \Sigma_d / tr(\Sigma_d)$. The corresponding backward operation $\frac{\partial L}{\partial \Sigma} = \psi_{ItN}(\frac{\partial L}{\partial \Sigma^N} + \frac{1}{2})$ is as follows:

\[
\frac{\partial L}{\partial P_T} = \frac{1}{\sqrt{tr(\Sigma)}} \frac{\partial L}{\partial \Sigma^N} \frac{1}{2}
\]

\[
\frac{\partial L}{\partial \Sigma_N} = -\frac{1}{2} \sum_{k=1}^{T} (P_{k-1}^T)^T \frac{\partial L}{\partial P_k}
\]

\[
\frac{\partial L}{\partial \Sigma} = \frac{1}{tr(\Sigma)} \frac{\partial L}{\partial \Sigma_N} - \frac{1}{(tr(\Sigma))^2} tr((\frac{\partial L}{\partial \Sigma_N}^T \Sigma)I
\]

\[- \frac{1}{2(tr(\Sigma))^{3/2}} tr((\frac{\partial L}{\partial \Sigma_N}^T P_T)^T)I.
\]

(17)

Here, $\frac{\partial L}{\partial P_k}$ can be calculated by the following iterations:

\[
\frac{\partial L}{\partial P_{k-1}} = \frac{3}{2} \frac{\partial L}{\partial P_k} - \frac{1}{2} \frac{\partial L}{\partial P_k} (P_{k-1}^T \Sigma_N) - \frac{1}{2} (P_{k-1}^T)^T \frac{\partial L}{\partial P_k} \Sigma_N^T
\]

\[- \frac{1}{2} (P_{k-1})^T \frac{\partial L}{\partial P_k} (P_{k-1}^T \Sigma_N)^T, \ k = T, ..., 1.
\]

(18)

We also provide the python code of GW using ItN whitening, based on PyTorch [36], in Figure 4.

B More Results on Effects of Batch Size

In Figure 1 of the submitted paper, we show the effects of batch size for different normalization methods, where the results are obtained with a learning rate of 0.1. Here, we provide more results using different learning rates, shown in Figure 5. We obtain similar observations.
Figure 5: Effects of batch size for different normalization methods. We train a four-layer multilayer perceptron (MLP) with 256 neurons in each layer, for MNIST classification. We compare BN, BW, group-based BW with 16 neurons/channels in each group (‘BW-C16’), and GW (we use a group number of 16). We vary the batch size and evaluate the training (thick ‘plus’ with solid line) and validation (thin ‘plus’ with dashed line) accuracy at the end of 50 training epochs. These results are obtained using a learning rate of (a) 0.01 and (b) 0.5.

C Derivation of Constraint Number of Normalization Methods

In Section 4 of the submitted paper, we define the constraint number of a normalization operation, and summarize the constraint number of different normalization methods in Table 1 at the submitted paper. Here, we provide the details for deriving the constraint number of batch whitening (BW), group normalization (GN) and our proposed GW, for the mini-batch input $X \in \mathbb{R}^{d \times m}$.

**Constraint number of BW.** BW [16] ensures that the normalized output is centered and whitened, which has the constraints $\Upsilon_{\phi_{BW}}(\hat{X})$ as:

\[
\hat{X}1 = 0_d, \quad \text{and} \quad \hat{X}^T - mI = 0_{d \times d},
\]

where $0_d$ is a $d$-dimensional column vector of all zeros, and $0_{d \times d}$ is a $d \times d$ matrix of all zeros. Note that there are $d$ independent equations in the system of equations $\hat{X}1 = 0_d$. Let’s denote $M = \hat{X}^T - mI$. We have $M^T = M$, and thus $M$ is a symmetric matrix. Therefore, there are $d(d + 1)/2$ independent equations in the system of equations $\hat{X}^T - mI = 0_{d \times d}$. We thus have $d(d + 1)/2 + d$ independent equations in $\Upsilon_{\phi_{BW}}(\hat{X})$, and the constraint number of BW is $d(d + 3)/2$.

**Constraint number of GN.** Given a sample $x \in \mathbb{R}^d$, GN divides the neurons into groups: $Z = \Pi(x) \in \mathbb{R}^{g \times c}$, where $g$ is the group number and $d = gc$. The standardization operation is then performed on $Z$ as:

\[
\hat{Z} = \Lambda_g^{-\frac{1}{2}}(Z - \mu_g 1^T),
\]

where, $\mu_g = \frac{1}{g}Z1$ and $\Lambda_g = \text{diag}(\sigma_1^2, \ldots, \sigma_g^2) + \epsilon I$. This ensures that the normalized output $\hat{Z}$ for each sample has the constraints:

\[
\sum_{j=1}^c \hat{Z}_{ij} = 0 \text{ and } \sum_{j=1}^c \hat{Z}_{ij}^2 = c, \text{ for each group } i = 1, \ldots, g.
\]

In the system of equations the number of independent equations is $2g$. Therefore, the constraint number of GN is $2dm$, when given $m$ samples.

**Constraint number of GW.** Given a sample $x \in \mathbb{R}^d$, GW performs normalization as:

\[
\text{Group division: } X_G = \Pi(x; g) \in \mathbb{R}^{g \times c},
\]

\[
\text{Whitening: } \hat{X}_G = \phi^W(X_G) = \Sigma_g^{-\frac{1}{2}}(X_G - \mu_g 1^T),
\]

\[
\text{Inverse group division: } \hat{x} = \Pi^{-1}(\hat{X}_G) \in \mathbb{R}^d.
\]
The normalization operation ensures that \( \hat{X}_G \in \mathbb{R}^{g \times c} \) has the following constraints:

\[
\hat{X}_G 1 = 0, \quad \text{and} \quad (26)
\]

\[
\hat{X}_G X_G^T = c1 = 0. \quad (27)
\]

Following the analysis for BW, the number of independent equations is \( g(g + 3)/2 \) from Eqns. (26) and (27). Therefore, the constraint number of GW is \( mg(g + 3)/2 \), when given \( m \) samples.

### D Learning Decorrelated Feature Representations

As described in Section 5.1.1 of the submitted paper, we investigate the effect of inserting a GW/BW layer after the last average pooling (before the last linear layer) to learn the decorrelated feature representations, as proposed in [18]. We provide the results in Table 6. This can slightly improve the performance (0.10% on average) when using GW (comparing Table 6 to Table 2 of the submitted paper). We note that BW benefits the most from this kind of architecture.

### E Running Time Comparison

In this section, we compare the wall-clock time of the models described in Section 5.1 of the submitted paper. We run the experiments on GPUs (NVIDIA Tesla V100). All implementations are based on the API provided by PyTorch, with CUDA (version number: 9.0). We use the same experimental setup as described in Section 5.1 of the submitted paper. We evaluate the training time for each architecture, averaged over 100 iterations. The ResNets-50 baseline (BN) costs 419 ms. Replacing the BNs of ResNet-50 with our GWs \((g=64)\) costs 796 ms, a 90% additional time cost on ResNet-50. This is one factor that drives us to investigate the position at which to apply GW.

Table 7 shows the time costs of five architectures, \( S1, S1-B1, S1-B2, S1-B3 \) and \( S1-B12 \), which have 1, 17, 17, 17 and 33 GW modules, respectively. Note that applying GW in the \( S1-B3 \) architecture results in a clearly increased computational cost, compared to \( S1-B1/S1-B2 \). This is because the channel number of the third normalization layer is \( 4 \times \) larger than that of the first/second normalization layer, in the bottleneck blocks [13].

Table 8 shows the time costs of ResNets [13] and ResNeXts [52] (the corresponding models in Table 3 of the submitted paper) for ImageNet classification.

### Table 6: Effects of inserting a GW/BW/BW\(_2\) layer after the last average pooling of ResNet-50 to learn decorrelated feature representations for ImageNet classification. We evaluate the top-1 validation accuracy on five architectures (\( S1, S1-B1, S1-B2, S1-B3 \) and \( S1-B12 \)), described in the submitted paper. Note that we also use an extra BN layer after the last average pooling for the Baseline (BN).

| Method | ResNet-50 | ResNet-101 | ResNeXt-50 | ResNeXt-101 |
|--------|-----------|------------|------------|-------------|
| Baseline (BN) [20] | 419 | 672 | 574 | 912 |
| BW [18] | 550 (\(\Delta 31.3\)% | 882 (\(\Delta 30.9\)% | 798 (\(\Delta 39.0\)% | 1587 (\(\Delta 74.0\)% |
| GW | 514 (\(\Delta 22.7\)% | 810 (\(\Delta 20.5\)% | 738 (\(\Delta 28.6\)% | 1180 (\(\Delta 29.3\)% |

### Table 7: Time costs (\(ms\)) of five architectures when applying GW on ResNet-50 (\( S1, S1-B1, S1-B2, S1-B3 \) and \( S1-B12 \)). Note that \( \Delta x\% \) indicates the additional time cost is \( x\% \), compared to the baseline.

| Method | ResNet-50 | ResNet-101 | ResNeXt-50 | ResNeXt-101 |
|--------|-----------|------------|------------|-------------|
| Baseline (BN) | 419 | 672 | 574 | 912 |
| BW [18] | 550 (\(\Delta 31.3\)% | 882 (\(\Delta 30.9\)% | 798 (\(\Delta 39.0\)% | 1587 (\(\Delta 74.0\)% |
| GW | 514 (\(\Delta 22.7\)% | 810 (\(\Delta 20.5\)% | 738 (\(\Delta 28.6\)% | 1180 (\(\Delta 29.3\)% |