

# Deep Learning with Low Precision by Half-wave Gaussian Quantization

Zhaowei Cai UC San Diego

zwcai@ucsd.edu

Xiaodong He Microsoft Research Redmond

xiaohe@microsoft.com

Jian Sun Megvii Inc.

sunjian@megvii.com

Nuno Vasconcelos UC San Diego

nuno@ucsd.edu

#### **Abstract**

The problem of quantizing the activations of a deep neural network is considered. An examination of the popular binary quantization approach shows that this consists of approximating a classical non-linearity, the hyperbolic tangent, by two functions: a piecewise constant sign function, which is used in feedforward network computations, and a piecewise linear hard tanh function, used in the backpropagation step during network learning. The problem of approximating the widely used ReLU non-linearity is then considered. An half-wave Gaussian quantizer (HWGQ) is proposed for forward approximation and shown to have efficient implementation, by exploiting the statistics of of network activations and batch normalization operations. To overcome the problem of gradient mismatch, due to the use of different forward and backward approximations, several piece-wise backward approximators are then investigated. The implementation of the resulting quantized network, denoted as HWGQ-Net, is shown to achieve much closer performance to full precision networks, such as AlexNet, ResNet, GoogLeNet and VGG-Net, than previously available low-precision networks, with 1-bit binary weights and 2-bit quantized activations.

# 1. Introduction

Deep neural networks have achieved state-of-the-art performance on computer vision problems, such as classification [21, 33, 34, 11, 12], detection [7, 31, 1], etc. However, their complexity is an impediment to widespread deployment in many applications of real world interest, where either memory or computational resource is limited. This is due to two main issues: large model sizes (50MB for GoogLeNet [34], 200M for ResNet-101 [12], 250MB for AlexNet [21], or 500M for VGG-Net [33]) and large computational cost, typically requiring GPU-based implementations. This generated interest in compressed models with smaller memory footprints and computation.

Several works have addressed the reduction of model size, through the use of quantization [3, 26, 24], low-rank

matrix factorization [18, 6], pruning [10, 9], architecture design [25, 16], etc. Recently, it has been shown that weight compression by quantization can achieve very large memory savings, reducing each weight to as little as 1 bit, at a marginal cost in classification accuracy [3, 26]. It is, however, less effective along the computational dimension, because the core network operation, implemented by each of its units, is the dot-product between a weight and an activation vector. Complementing binary or quantized weights with quantized activations would enable the replacement of expensive dot-products by logical and bit-counting operations. Substantial speed ups should thus be possible if, in addition to weights, the inputs of each unit were binarized or quantized to low-bit.

However, the quantization of activations is more difficult than that of weights. For example, [4, 30] have shown that, while it is possible to binarize weights with a marginal cost in model accuracy, additional quantization of activations incurs nontrivial losses for large-scale classification, such as on ImageNet [32]. The difficulty is that binarization or quantization of activations requires their processing with non-differentiable operators that create problems for the backpropagation algorithm. This iterates between a feedforward step that computes network outputs and a backpropagation step that computes the gradients required for learning. The difficulty is that binarization or quantization operators have step-wise responses that produce very weak gradient signals during backpropagation, compromising learning efficiency. So far, the problem has been addressed by using continuous approximations of the operator used in the feedforward step to implement the backpropagation step. This, however, creates a mismatch between the model that implements the forward computations and the derivatives used to learn it, leading to a sub-optimal model.

In this work, we view the quantization operator, used in the feedforward step, and the continuous approximation, used in the backpropagation step, as two functions that approximate the activation function of each network unit. We refer to these as the *forward* and *backward* approximation of the activation function. We start by considering the binary  $\pm 1$  quantizer, used in [4, 30], for which

these two functions can be seen as a discrete and a continuous approximation of a non-linear activation function, the hyperbolic tangent, frequently used in classical neural networks. This activation is, however, not commonly used in recent deep learning literature, where the ReLU nonlinearity [28, 36, 11] has achieved much greater preponderance. This is exactly because it produces much stronger gradient magnitudes. While the hyperbolic tangent or sigmoid nonlinearities are squashing non-linearities and mostly flat, the ReLU is an half-wave rectifier, of linear response to positive inputs. Hence, while the derivatives of the hyperbolic tangent are close to zero almost everywhere, the ReLU has unit derivative along the entire positive range of the axis.

To improve the learning efficiency of quantized networks, we consider the design of forward and backward approximation functions for the ReLU. To discretize its linear component, we propose to use an optimal quantizer. By exploiting the statistics of network activations and batch normalization operations that are commonly used in the literature, we show that this can be done with an half-wave Gaussian quantizer (HWGQ) that requires no learning and is very efficient to compute. While some recent works have attempted similar ideas [4, 30], their design of a quantizer is not sufficient to guarantee good deep learning performance. We address this problem by complementing this design with a study of suitable backward approximation functions that account for the mismatch between the forward model and the back propagated derivatives. This study suggests operations such as linearization, gradient clipping or gradient suppression for the implementation of the backward approximation. We show that a combination of the forward HWGQ with these backward operations produces very efficient low-precision networks, denoted as HWGQ-Net, with much closer performance to continuous models, such as AlexNet [21], ResNet [12], GoogLeNet [34] and VGG-Net [33], than other available low-precision networks in the literature. To the best of our knowledge, this is the first time that a single low-precision algorithm could achieve success for so many popular networks. Using the arguments of [30], the HWGQ-Net (1-bit weights and 2-bit activations) could theoretically achieve  $\sim 32 \times$  memory and  $\sim 32 \times$  convolutional computation savings. This suggests that the HWGQ-Net can be very useful for the deployment of state-of-the-art neural networks in real world applications.

### 2. Related Work

The reduction of model size is a popular goal in the deep learning literature. One strategy is to exploit the widely known redundancy of neural network weights [5]. For example, [6, 18] proposed low-rank matrix factorization as a way to decompose a large weight matrix into several separable small matrices. An alternative procedure, known as connection pruning [10, 9], consists of removing unimportant

connections of a pre-trained model and retraining, showing considerable model reduction without noticable loss in accuracy. Another model compression strategy is to constrain the model architecture itself, e.g. by removing fully connected layers, using convolutional filters of small size, etc. Many state-of-the-art deep networks, such as NIN [25], GoogLeNet [34] and ResNet [12], rely on such design choices. For example, SqueezeNet [16] has been shown to achieve a parameter reduction of ~50 times, for accuracy comparable to that of AlexNet. Moreover, hash functions have also been used to compress model size [2].

Another branch of approaches for model compression is weight binarization [3, 30, 4] or quantization [26, 24, 8]. [35] used a fixed-point representation to quantize weights of pre-trained neural networks. [8] showed that vector quantization enables  $4\sim8$  times compression with minimal accuracy loss. [24] proposed a method for fixed-point quantization based on the optimal bit-width allocations across network layers. [22, 26] have shown that ternary weight quantization into levels  $\{-1,0,1\}$  can achieve  $16\times$  or  $32\times$  model compression with slight accuracy loss, even on large-scale classification tasks. Finally, [3] has shown that filter weights can be quantized to  $\pm1$  without noticeable loss of classification accuracy on CIFAR-10 [20].

Since quantization of activations enables further speedups and reduces training memory requirements, it has attracted some attention [35, 24, 4, 30, 37, 23, 26]. [35, 24] performed the quantization after network training, avoiding the issues of nondifferentiable optimization. Recently, [4, 30, 37] tried to tackle the nondifferentiable optimization issue by using a continuous approximation to the quantizer function in the backpropagation step. [23] proposed several potential solutions to the problem of gradient mismatch and [26, 37] showed that gradients can be quantized with a small number of bits during the backpropagation step. While some of these methods have produced good results on CIFAR-10, none has produced low precision networks competitive with full-precision models on large-scale classification tasks, such as ImageNet [32].

#### 3. Binary Networks

We start with a brief review of the issues involved in the binarization of a deep network.

#### **3.1. Goals**

Deep neural networks consist of layers of units that roughly model the computations of neurons in the mammalian brain. Each unit computes an activation function

$$z = g(\mathbf{w}^T \mathbf{x}),\tag{1}$$

where  $\mathbf{w} \in \mathbb{R}^{c \cdot w \cdot h}$  is a weight vector,  $\mathbf{x} \in \mathbb{R}^{c \cdot w \cdot h}$  an input vector, and  $g(\cdot)$  a non-linear function. A convolutional

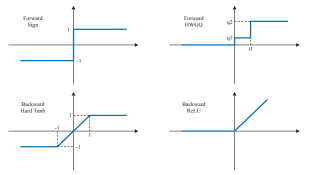


Figure 1. Forward and backward functions for binary sign (left) and half-wave Gaussian quantization (right) activations.

network implements layers of these units, where weights are usually represented as a tensor  $\mathbf{W} \in \mathbb{R}^{c \times w \times h}$ . The dimensions c, w and h are defined by the number of filter channels, width and height, respectively. Since modern networks have very large numbers of these units, the structure of (1) is the main factor in the complexity of the overall model. This complexity can be a problem for applications along two dimensions. The first is the large memory footprint required to store weights  $\mathbf{w}$ . The second is the computational complexity required to compute large numbers of dot-products  $\mathbf{w}^T\mathbf{x}$ . Both difficulties are compounded by the requirement of floating point storage of weights and floating point arithmetic to compute dot-products, which are not practical for many applications. This has motivated interest in low-precision networks [4, 30, 37].

# 3.2. Weight Binarization

An effective strategy to binarize the weights  $\mathbf{W}$  of convolutional filters, which we adopt in this work, has been proposed by [30]. This consists of approximating the full precision weight matrix  $\mathbf{W}$ , used to compute the activations of (1) for all the units, by the product of a binary matrix  $\mathbf{B} \in \{+1, -1\}^{c \times w \times h}$  and a scaling factor  $\alpha \in \mathbb{R}^+$ , such that  $\mathbf{W} \approx \alpha \mathbf{B}$ . A convolutional operation on input  $\mathbf{I}$  can then be approximated by

$$\mathbf{I} * \mathbf{W} \approx \alpha(\mathbf{I} \oplus \mathbf{B}),$$
 (2)

where  $\oplus$  denotes a multiplication-free convolution. [30] has shown that an optimal approximation can be achieved with  $\mathbf{B}^* = sign(\mathbf{W})$  and  $\alpha^* = \frac{1}{cwh} \|\mathbf{W}\|_1$ . While binary weights tremendously reduce the memory footprint of the model, they do not fully solve the problem of computational complexity. Substantial further reductions of complexity can be obtained by the binarization of  $\mathbf{I}$ , which enables the implementation of dot products in (2) with logical and bit-counting operations [4, 30].

### 3.3. Binary Activation Quantization

The use of binary activations has been suggested in [30, 4, 37]. It is usually implemented by replacing g(x) in (1) with the sign non-linearity

$$z = sign(x) = \begin{cases} +1, & \text{if } x \ge 0, \\ -1, & \text{otherwise} \end{cases}$$
 (3)

shown in Figure 1. This creates difficulties to the backpropagation algorithm used to learn the neural network, by minimizing a cost C with respect to the weights  $\mathbf{w}$ . Consider the unit of (1). The derivative of C with respect to  $\mathbf{w}$  is

$$\frac{\partial C}{\partial \mathbf{w}} = \frac{\partial C}{\partial z} g'(\mathbf{w}^T \mathbf{x}) \mathbf{x}.$$
 (4)

When g(x) is replaced by (3), the derivative  $g'(\mathbf{w}^T\mathbf{x})$  is zero almost everywhere and the gradient magnitudes tend to be very small. In result, the gradient descent algorithm does not converge to minima of the cost. To overcome this problem, [4] proposed to use an alternative function, *hard tanh*, which we denote by  $\widehat{sign}$ , in the backpropagation step. This function is shown in Figure 1, and has derivative

$$\widetilde{sign}'(x) = \begin{cases} 1, & \text{if } |x| \le 1\\ 0, & \text{otherwise.} \end{cases}$$
 (5)

In this work, we denote (3) as the forward and (5) as the backward approximations of the activation non-linearity g(x) of (1). These approximations have two main problems. The first is that they approximate the hyperbolic tangent (tanh), which is a squashing non-linearity. The saturating behavior of squashing non-linearities (such as the tanh or the sigmoid) emphasizes the problem of vanishing derivatives, compromising the effectiveness of backpropagation. The second is that the discrepancy between the approximation of g(x) by the forward sign and by the backward sign creates a mismatch between the feedforward model and the derivatives used to learn it. In result, backpropagation can be highly suboptimal. This is called the "gradient mismatch" problem [23].

# 4. Half-wave Gaussian Quantization

In this section, we propose an alternative quantization strategy, the approximation of the ReLU non-linearity.

#### **4.1. ReLU**

The ReLU is the half-wave rectifier defined by [28],

$$q(x) = \max(0, x). \tag{6}$$

It is now well known that, when compared to squashing non-linearities, its use in (1) significantly improves the efficiency of the backpropagation algorithm. It thus seems

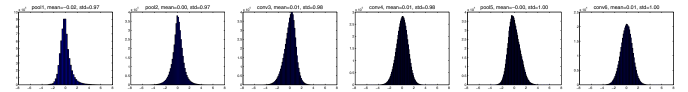


Figure 2. Dot-product distributions on different layers of AlexNet with binary weights and quantized activations (100 random images).

more sensible to rely on ReLU approximations for network quantization than those of the previous section. We propose a quantizer Q(x) to approximate (6) in the feedforward step and a suitable piecewise linear approximation  $\widetilde{Q}(x)$  for the backpropagation step.

### 4.2. Forward Approximation

A quantizer is a piecewise constant function

$$Q(x) = q_i, \quad if \quad x \in (t_i, t_{i+1}],$$
 (7)

that maps all values of x within quantization interval  $(t_i,t_{i+1}]$  into a quantization level  $q_i\in\mathbb{R}$ , for  $i=1,\cdots,m$ . In general,  $t_1=-\infty$  and  $t_{m+1}=\infty$ . This generalizes the sign function, which can be seen as a 1-bit quantizer. A quantizer is denoted uniform if

$$q_{i+1} - q_i = \Delta, \quad \forall i, \tag{8}$$

where  $\Delta$  is a constant quantization step. The quantization levels  $q_i$  act as reconstruction values for x, under the constraint of reduced precision. Since, for any x, it suffices to store the quantization index i of (7) to recover the quantization level  $q_i$ , non-uniform quantization requires  $\log_2 m$  bits of storage per activation x. However, more than  $\log_2 m$  bits are needed to represent x in arithmetic operations, since these use  $q_i$ , not the index i. For a uniform quantizer, where  $\Delta$  is a universal scaling factor that can be placed in evidence,  $\log_2 m$  bits are enough for both storage and arithmetic computation.

Optimal quantizers are usually defined in the meansquared error sense, i.e.

$$Q^*(x) = \arg\min_{Q} E_x[(Q(x) - x)^2]$$

$$= \arg\min_{Q} \int p(x)(Q(x) - x)^2 dx$$
(9)

where p(x) is the probability density function of x. Hence, the optimal quantizer of the dot-products of (1) depends on their statistics. While the optimal solution  $Q^*(x)$  of (9) is usually non-uniform, a uniform solution  $Q^*(x)$  is available by adding the uniform constraint of (8) to (9). Given dot product samples, the optimal solution of (9) can be obtained by Lloyd's algorithm [27]. This, however, is an iterative algorithm. Since a different quantizer must be designed per

network unit, and this quantizer changes with the backpropagation iteration, the straightforward application of this procedure is computationally intractable.

This difficulty can be avoided by exploiting the statistical structure of the activations of deep networks. For example, [15, 17] have noted that the dot-products of (1) tend to have a symmetric, non-sparse distribution, that is close to Gaussian. Taking into account the fact that the ReLU is a half-wave rectifier, this suggests the use of the half-wave Gaussian quantizer (HWGQ),

$$Q(x) = \begin{cases} q_i, & \text{if } x \in (t_i, t_{i+1}], \\ 0, & x \le 0, \end{cases}$$
 (10)

where  $q_i \in \mathbb{R}^+$  for  $i=1,\cdots,m$  and  $t_i \in \mathbb{R}^+$  for  $i=1,\cdots,m+1$  ( $t_1=0$  and  $t_{m+1}=\infty$ ) are the optimal quantization parameters for the Gaussian distribution. The adoption of the HWGQ guarantees that these parameters only depend on the mean and variance of the dot-product distribution. However, because these can vary across units, it does not eliminate the need for the repeated application of Lloyd's algorithm across the network.

This problem can be alleviated by resorting to batch normalization [17]. This is a widely used normalization technique, which forces the responses of each network layer to have zero mean and unit variance. We apply this normalization to the dot-products, with the result illustrated in Figure 2, for a number of AlexNet units of different layers. Although the distributions are not perfectly Gaussian and there are minor differences between them, they are all close to Gaussian with zero mean and unit variance. It follows that the optimal quantization parameters  $q_i^*$  and  $t_i^*$  are approximately identical across units, layers and backpropagation iterations. Hence, Lloyd's algorithm can be applied once, with data from the entire network. In fact, because all distributions are approximately Gaussian of zero mean and unit variance, the quantizer can even be designed from samples of this distribution. In our implementation, we drew  $10^6$ samples from a standard Gaussian distribution of zero mean and unit variance, and obtained the optimal quantization parameters by Lloyd's algorithm. The resulting parameters  $t_i^*$ and  $q_i^*$  were used to parametrize a single HWGQ that was used in all layers, after batch normalization of dot-products.

### 4.3. Backward Approximation

Since the HWGQ is a step-wise constant function, it has zero derivative almost everywhere. Hence, the approximation of g(x) by Q(x) in (4) leads to the problem of vanishing derivatives. As in Section 3, a piecewise linear function  $\widetilde{Q}(x)$  can be used during the backpropagation step to avoid weak convergence. In summary, we seek a piece-wise function that provides a good approximation to the ReLU and to the HWGQ. We next consider three possibilities.

#### 4.3.1 Vanilla ReLU

Since the ReLU of (6) is already a piece-wise linear function, it seems sensible to use the ReLU itself, denoted the *vanilla ReLU*, as the backward approximation function. This corresponds to using the derivative

$$\widetilde{Q}'(x) = \begin{cases} 1, & \text{if } x > 0, \\ 0, & \text{otherwise} \end{cases}$$
 (11)

in (4). The forward and backward approximations Q(x) and  $\widetilde{Q}(x)$  of the ReLU are shown in Figure 1. Note that, while the backward approximation is exact, it is not equal to the forward approximation. Hence, there is a gradient mismatch. For x>0, the approximation of Q(x) by the ReLU has error |Q(x)-x|. This is upper bounded by  $(t_{i+1}-q_i)$  for  $x\in (t_i,t_{i+1}]$  but unbounded when  $x\in (t_m,\infty)$ . Hence, the mismatch is particularly large for large values of x. Since these are the values on the tail of the distribution of x, the ReLU is said to have a large mismatch with Q(x) "on the tail." When the ReLU is used to approximate g'(x) in (4), it can originate very inaccurate gradients for large dot-products. In our experience, this can make the learning algorithm unstable.

This is a classical problem in the robust estimation literature, where outliers can unduly influence the performance of a learning algorithm [14]. For quantization, where Q(x) assumes that values of x beyond  $q_m$  have very low probability, large dot-products are effectively outliers. The classical strategy for outlier mitigation is to limit the growth rate of the error function, in this case |Q(x)-x|. Hence, the problem is the monotonicity of the ReLU beyond  $x=q_m$ . To address it, we investigate alternative backwards approximation functions of slower growth rate.

## 4.3.2 Clipped ReLU

The first approximation, denoted the *clipped ReLU*, is identical to the vanilla ReLU in  $(-\infty, q_m]$  but constant beyond  $x=q_m$ ,

$$\widetilde{Q}_c(x) = \begin{cases} q_m, & x > q_m, \\ x, & x \in (0, q_m], \\ 0, & \text{otherwise.} \end{cases}$$
 (12)

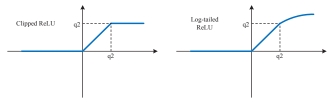


Figure 3. Backward piece-wise activation functions of clipped ReLU and log-tailed ReLU.

Its use to approximate  $g'(\mathbf{w}^T\mathbf{x})$  in (4) guarantees that there is no mismatch on the tail. Gradients are non-zero only for dot-products in the interval  $(0, q_m]$ . As illustrated in Figure 3, the clipped ReLU is a better match for the HWGQ than the vanilla ReLU. In our experiments, ReLU clipping proved very useful to guarantee a stable optimization. This is similar to previous observations that gradient clipping robustifies the learning of very deep networks [29].

#### 4.3.3 Log-tailed ReLU

Ideally, a network with quantized activations should approach the performance of a full-precision network as the number of quantization levels m increases. The sensitivity of the vanilla ReLU approximation to outliers limits the performance of low precision networks. While the clipped ReLU alleviates this problem, it can impair network performance due to the loss of information in the clipped interval  $(q_m, \infty)$ . An intermediate solution is to use, in this interval, a function whose growth rate is in between that of the clipped ReLU (zero derivative) and the ReLU (unit derivative). One possibility is to enforce logarithmic growth on the tail, according to

$$\widetilde{Q}_{l}(x) = \begin{cases}
q_{m} + \log(x - \tau), & x > q_{m}, \\
x, & x \in (0, q_{m}], \\
0, & x \leq 0,
\end{cases} (13)$$

where  $\tau = q_m - 1$ . This is denoted the *log-tailed ReLU* and is shown in Figure 3. It has derivative

$$\widetilde{Q}'_{l}(x) = \begin{cases}
1/(x-\tau), & x > q_{m}, \\
1, & x \in (0, q_{m}], \\
0, & x \le 0.
\end{cases}$$
(14)

When used to approximate g'(x) in (4), the log-tailed ReLU is identical to the vanilla ReLU for amplitudes smaller than  $q_m$ , but gives decreasing weight to amplitudes larger than this. It behaves like the vanilla ReLU (unit derivative) for  $0 < x \le q_m$  but converges to the clipped ReLU (zero derivative) as x grows to infinity.

### 5. Experimental Results

The proposed HWGQ-Net was evaluated on ImageNet (ILSVRC2012) [32], which has ~1.2M training images

Table 1. Full-precision Activation Comparison for AlexNet.

	Full	$FW+\widetilde{sign}$	$FW+\widetilde{Q}$	$BW+\widetilde{sign}$	$BW+\widetilde{Q}$		
Top-1	55.7	46.7	55.7	43.9	53.9		
Top-5	79.3	71.0	79.3	68.3	77.3		

Table 2. Low-bit Activation Comparison.

Model	Full	BW	FW+Q	BW+sign	BW+Q	
AlexNet	Top-1	55.7	52.4	49.5	39.5	46.8
Alexinet	Top-5	79.3	75.9	73.7	63.6	71.0
ResNet-18	Top-1	66.3	61.3	37.5	42.1	33.0
KCSINCI-10	Top-5	87.5	83.6	61.9	67.1	56.9
VGG-Variant	Top-1	68.6	65.5	48.3	50.1	44.1
v GG- variant	Top-5	88.9	86.5	72.3	74.3	68.7

from 1K categories and 50K validation images. The evaluation metrics were top-1 and top-5 classification accuracy. Several popular networks were tested: AlexNet [21], ResNet [12], a variant of VGG-Net [33, 11], and GoogLeNet [34]. Our implementation uses Caffe [19], see source code at https://github.com/zhaoweicai/hwgq.

### 5.1. Implementation Details

In all experiments, training images were resized to  $256 \times 256$ , and a  $224 \times 224$  ( $227 \times 227$  for AlexNet) crop was randomly sampled from an image or its horizontal flip. Batch normalization [17] was applied before each quantization layer, as discussed in Section 4.2. The ratio of dropout [13] was set as 0.1 for networks with binary weights and full activations, but no dropout was used for networks with quantized activations. All networks were learned from scratch with SGD. No data augmentation was used other than standard random image flipping and cropping. No bias term was used for binarized weights. As in [30], networks with quantized activations used max-pooling before batch normalization. This is denoted "layer re-ordering". As in [30, 37], the first and last network layers had full precision. Evaluation was based solely on central  $224 \times 224$  crops.

On AlexNet [21] experiments, the mini-batch size was 256, weight decay 0.0005, and learning rate started at 0.01. For ResNet, we used the parameters of [12]. For the variant of VGG-Net, denoted VGG-Variant, a smaller version of model-A in [11], only 3 convolutional layers were used for input size of 56, 28 and 14, and the "spp" layer was removed. The mini-batch size was 128, and learning rate started at 0.01. For GoogLeNet [34], the branches for side losses were removed, in the inception layers, max-pooling was removed and the channel number of the "reduce"  $1\times1$ convolutional layers was increased to that of their following  $3\times3$  and  $5\times5$  convolutional layers. Weight decay was 0.0002 and the learning strategy was similar to ResNet [12]. For all networks tested, momentum was 0.9, and when mini-batch size was 256 (128), the learning rate was divided by 10 after every 50K (100K) iterations, 160K (320K) in total. Only AlexNet, ResNet-18 and VGG-Variant were explored in the following ablation studies. In all tables and fig-

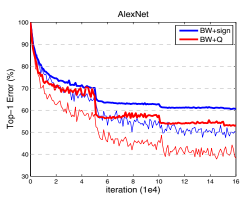


Figure 4. The error curves of training (thin) and test (thick) for sign(x) and Q(x) (HWGQ) activation functions.

ures, "FW" indicates full-precision weights, "BW" binary weights, and "Full" full-precision weights and activations.

### 5.2. Full-precision Activation Comparison

Before considering the performance of the forward quantized activation functions sign(x) and Q(x), we compared the performance of the continuous  $\widetilde{sign}(x)$  (hard tanh) and  $\widetilde{Q}(x)$  (ReLU) as activation function. In this case, there is no activation quantization nor forward/backward gradient mismatch. AlexNet results are presented in Table 1, using identical settings for  $\widetilde{sign}(x)$  and  $\widetilde{Q}(x)$ , for fair comparison. As expected from the discussion of Sections 3 and 4,  $\widetilde{Q}(x)$  achieved substantially better performance than  $\widetilde{sign}(x)$ , for both FW and BW networks. The fact that these results upper bound the performance achievable when quantization is included suggests that sign(x) is not a good choice for quantization function. Q(x), on the other hand, has a fairly reasonable upper bound.

#### 5.3. Low-bit Activation Quantization Results

We next compared the performance achieved by adding the sign and HWGQ Q(x) (backward vanilla ReLU) quantizers to the set-up of the previous section. The results of AlexNet, ResNet-18 and VGG-Variant are summarized in Table 2. Notice, first, that BW has weaker performance than BW+Q of AlexNet in Table 1, due to the impact of the layer re-ordering [30] introduced in Section 5.1. Next, comparing BW to FW+Q, where the former binarizes weights only and the latter quantizes activations only, it can be seen that weight binarization causes a minor degradation of accuracy. This is consistent with the findings of [30, 4]. On the other hand, activation quantization leads to a nontrivial loss. This confirms that the latter is a more difficult problem.

When weight binarization and activation quantization were combined, recognition performance dropped even further. For AlexNet, the drop was much more drastic for BW+sign (backward hard tanh) than for BW+Q (back-

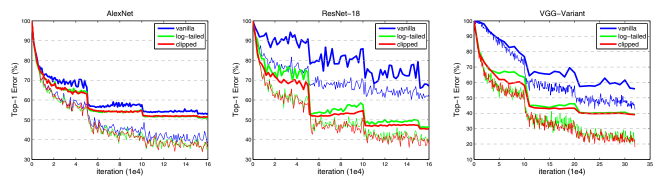


Figure 5. The error curves of training (thin) and test (thick) for alternative backward approximations.

Table 3. Backward Approximations Comparison.

		1.1			1		
Model	BW	no-opt	vanilla	clipped	log-tailed		
AlexNet	Top-1	52.4	30.0	46.8	48.6	49.0	
Alexinet	Top-5	75.9	53.6	71.0	72.8	73.1	
ResNet-18	Top-1	61.3	34.2	33.0	54.5	53.5	
Kesnet-10	Top-5	83.6	59.6	56.9	78.5	77.7	
VGG-Variant	Top-1	65.5	42.8	44.1	60.9	60.6	
voo-variant	Top-5	86.5	68.3	68.7	83.2	82.9	

ward vanilla ReLU). These results support the hypotheses of Section 3 and 4, as well as the findings of Table 1. The training errors of BW+sign and BW+Q for AlexNet are shown in Figure 4. Note the much lower training error of Q(x), suggesting that it enables a much better approximation of the full precision activations than sign(x). Nevertheless, the gradient mismatch due to the use of Q(x)as forward and the vanilla ReLU as backward approximators made the optimization somewhat instable. For example, the error curve of BW+Q is bumpy during training. This problem becomes more serious for deeper networks. In fact, for the ResNet-18 and VGG-Variant, BW+Q performed worse than BW+sign. This can be explained by the fact that the sign has a smaller gradient mismatch problem than the vanilla ReLU. Substantial improvements are possible by correcting the mismatch between the forward quantizer Q(x) and its backward approximator.

#### 5.4. Backward Approximations Comparison

We next considered the impact of the backward approximators of Section 4.3. Table 3 shows the performance under the different approximations. In all cases, weights were binarized and the HWGQ was used as forward approximator (quantizer). "no-opt" refers to the quantization of activations of pre-trained BW networks. This requires no non-differentiable approximation, but fails to account for the quantization error. We attempted to minimize the impact of cumulative errors across the network by recomputing the means and variances of all batch normalization layers. Even after this, "no-opt" had significantly lower accuracy than the full-precision activation networks.

Substantial gains were obtained by training the activa-

Table 4. Bit-width Comparison of Activation Quantization.

quantization type			non-uniform				uniform		none
# levels			2	3	7	15	3*	7*	BW
AlovN	AlexNet	Top-1	48.6	50.6	52.4	52.6	50.5	51.9	52.4
Alexiv		Top-5	72.8	74.3	75.8	76.2	74.6	75.7	75.9
PacNat	ResNet-18	Top-1	54.5	57.6	60.3	60.8	56.1	59.6	61.3
Kesivet		Top-5	78.5	81.0	82.8	83.4	79.7	82.4	83.6

tion quantized networks from scratch. Although the vanilla ReLU had reasonable performance as backwards approximator for AlexNet, much better results were achieved with the clipped ReLU of (12) and the log-tailed ReLU of (13). Figure 5 shows that the larger gradient mismatch of the vanilla ReLU created instabilities in the optimization, for all networks. However, these instabilities were more serious for the deeper networks, such as ResNet-18 and VGG-Variant. This explains the sharper drop in performance of the vanilla ReLU for these networks, in Table 3. Note, in Figure 5, that the clipped ReLU and the log-tailed ReLU enabled more stable learning and reached a much better optimum for all networks. Among them, the log-tailed ReLU performed slightly better than the clipped ReLU on AlexNet, but slightly worse on ResNet-18 and VGG-Variant. To be consistent, "clipped ReLU" was used in the following sections.

## 5.5. Bit-width Impact

The next set of experiments studied the bit-width impact of the activation quantization. In all cases, weights were binarized. Table 4 summarizes the performance of AlexNet and ResNet-18 as a function of the number of quantization levels. While the former improved with the latter, there was a saturation effect. The default HWGQ configuration, also used in all previous experiments, consisted of two non-uniform positive quantization levels plus a "0". This is denoted as "2" in the table. For AlexNet, this very low-bit quantization sufficed to achieve recognition rates close to those of the full-precision activations. For this network, quantization with seven non-uniform levels was sufficient to reproduce the performance of full-precision activations. For ResNet-18, however, there was a more noticeable gap be-

Table 5. HWGQ implementation of various popular networks.

Model	Reference	Full	HWGQ	
AlexNet	Top-1	57.1	58.5	52.7
Alexinet	Top-5	80.2	81.5	76.3
ResNet-18	Top-1	69.6	67.3	59.6
ICSINCI-10	Top-5	89.2	87.9	82.2
ResNet-34	Top-1	73.3	69.4	64.3
ICSI (CI-54	Top-5	91.3	89.1	85.7
ResNet-50	Top-1	76.0	71.5	64.6
Resiret-50	Top-5	93.0	90.5	85.9
VGG-Variant	Top-1	-	69.8	64.1
voo-variant	Top-5	-	89.3	85.6
GoogLeNet	Top-1	68.7	71.4	63.0
Googleitei	Top-5	88.9	90.5	84.9

tween low-bit and full-precision activations. These results suggest that increasing the number of quantization levels is more beneficial for ResNet-18 than for AlexNet.

Table 4 also shows the results obtained with uniform quantization, with superscript "\*". Interestingly, for the same number of quantization levels, the performance of the uniform quantizer was only slightly worse than that of its non-uniform counterpart. This is, however, not a completely fair comparison since, as discussed in Section 4.2, non-uniform quantization requires more bits for arithmetic operations. For the same bit width, e.g. both "2" and "3\*" require a 2-bit representation for arithmetic computation, the uniform quantizer was noticeably superior to the non-uniform quantizer.

### 5.6. Comparison to the state-of-the-art

Table 5¹ presents a comparison between the full precision and the low-precision HWGQ-Net of many popular network architectures. In all cases, the HWGQ-Net used 1-bit binary weights, a 2-bit uniform HWGQ as forward approximator, and the clipped ReLU as backwards approximator. Comparing to the previous ablation experiments, the numbers of training iterations were doubled and polynomial learning rate annealing (power of 1) was used for HWGQ-Net. Table 5 shows that the HWGQ-Net approximates well all popular networks, independently of their complexity or depth. The top-1 accuracy drops from full- to low-precision are similar for all networks (5~9 points), suggesting that low-precision HWGQ-Net will achieve improved performance as better full-precision networks become available.

Training a network with binary weights and low-precision activations from scratch is a new and challenging problem, only addressed by a few previous works [4, 30, 37]. Table 6 compares the HWGQ-Net with the recent XNOR-Net [30] and DOREFA-Net [37], on the ImageNet classification task. The DOREFA-Net result is for a model of binary weights, 2-bit activation, full precision

Table 6. Comparison to state-of-the-art low-precision methods. Top-1 gap to the corresponding full-precision network is also reported.

Model		AlexNet	ResNet-18		
Model	XNOR	DOREFA	HWGQ	XNOR	HWGQ
Top-1	44.2	47.7	52.7	51.2	59.6
Top-5	69.2	-	76.3	73.2	82.2
Top-1 gap	-12.4	-8.2	-5.8	-18.1	-7.7

gradient and no pre-training. For AlexNet, the HWGQ-Net outperformed the XNOR-Net and the DOREFA-Net by a large margin. Similar improvements over the XNOR-Net were observed for the ResNet-18, where DOREFA-Net results are not available. It is worth noting that the gaps between the full-precision networks and the HWGQ-Net (-5.8 for AlexNet and -7.7 for ResNet-18) are much smaller than those of the XNOR-Net (-12.4 for AlexNet and -18.1 for ResNet-18) and the DOREFA-Net (-8.2 for AlexNet). This is strong evidence that the HWGQ is a better activation quantizer. Note that, in contrast to the experimentation with one or two networks by [4, 30, 37], the HWGQ-Net is shown to perform well for various network architectures. To the best of our knowledge, this is the first time that a single low-precision network is shown to successfully approximate many popular networks.

#### 6. Conclusion

In this work, we considered the problem of training high performance deep networks with low-precision. This was achieved by designing two approximators for the ReLU non-linearity: a half-wave Gaussian quantizer in the feedforward computations, and a piece-wise continuous function in the backpropagation step. This design overcomes the learning inefficiency of the popular binary sign quantization procedure. To minimize the problem of gradient mismatch, we have studied several backwards approximation functions, including clipped ReLU and log tailed ReLU approximators. The proposed network, denoted HWGQ-Net, was shown to significantly outperform previous efforts at deep learning with low precision, for various state-of-the-art networks. These promising results suggest that the HWGQ-Net can be very useful for the deployment of state-of-the-art neural networks in real world applications.

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<sup>&</sup>lt;sup>1</sup>The reference performance of AlexNet and GoogLeNet is at https://github.com/BVLC/caffe, and of ResNet is at https://github.com/facebook/fb.resnet.torch. Our worse ResNet implementations are probably due to fewer training iterations and no further data augmentation.

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