ABSTRACT

We formulate the preparation of a neural network for pruning and few-bit quantization as a variational inference problem. We introduce a quantizing prior that leads to a multi-modal, sparse posterior distribution over weights and further derive a differentiable KL approximation for this prior. After training with Variational Network Quantization (VNQ), weights can be replaced by deterministic quantization values with small to negligible loss of task accuracy (including pruning by setting weights to 0). Our method does not require fine-tuning after quantization. We show results for ternary quantization on LeNet-5 (MNIST) and DenseNet-121 (CIFAR-10).

1 INTRODUCTION

Parameters of a trained network commonly exhibit high degrees of redundancy (Denil et al., 2013) which implies an over-parametrization of the network. Network compression methods implicitly or explicitly aim at the systematic reduction of redundancy in neural network models while at the same time retaining a high level of task accuracy. Besides architectural approaches, such as SqueezeNet (Iandola et al., 2016) or MobileNets (Howard et al., 2017), many compression methods either perform some form of pruning or quantization. Pruning is the removal of irrelevant units (weights, neurons or convolutional filters) (LeCun et al. 1990). Relevance of weights is often determined by the absolute value (“magnitude based pruning” (Han et al. 2016, 2017, Guo et al. 2016), but more sophisticated methods have been known for decades, e.g. based on second-order (Optimal Brain Damage (LeCun et al. 1990) and Optimal Brain Surgeon Hassibi & Stork (1993)) or ARD (automatic relevance determination, a Bayesian framework for determining the relevance of weights, (MacKay 1995, Neal 1995, Karaletsos & Rätsch 2015)). Quantization is the reduction of the bit-precision of weights, activations or even gradients, which is particularly desirable from a hardware perspective (Sze et al. 2017). Methods range from fixed bit-width computation (e.g. 12-bit fixed point) to aggressive quantization such as binarization of weights and activations (Courbariaux et al. 2016, Rastegari et al. 2016, Zhou et al. 2016, Hubara et al. 2016). Few-bit quantization (2 to 6 bits) is often performed by k-means clustering of trained weights with subsequent fine-tuning of the cluster centers (Han et al. 2016). Pruning and quantization have been shown to work well in conjunction (Han et al. 2016). In so-called “ternary” networks, weights are either negative, zero or positive which also allows for simultaneous pruning and few-bit quantization (Li et al. 2016, Zhu et al. 2016).

Our work is closely related to some recent Bayesian methods for network compression (Ullrich et al. 2017, Molchanov et al. 2017, Louizos et al. 2017, Neklyudov et al. 2017) that learn a posterior distribution over network weights under a sparsity-inducing prior. The posterior distribution over network parameters allows identifying redundancies through three means: (1) weights with an expected value very close to zero and (2) weights with a large variance can be pruned as they do not contribute much to the overall computation. (3) the posterior variance over non-pruned parameters can be used to determine the required bit-precision (quantization noise can be made as large as implied by the posterior uncertainty). Additionally, variational Bayesian inference is known to automatically reduce parameter redundancy by penalizing overly complex models.

In this paper we present Variational Network Quantization, a Bayesian network compression method for simultaneous pruning and few-bit quantization of weights. We extend previous work introducing a multi-modal quantizing prior that penalizes weights of low variance unless they lie close to one of the target values for quantization. As a result, weights are either drawn to one of the quantization target values or they are assigned large variance values—see Fig. 1. After training, our method yields a
Bayesian Neural Network with a multi-modal posterior over weights (typically with one mode fixed at 0), which is the basis for subsequent pruning and quantization. However, posterior uncertainties can also be interesting for network introspection and analysis, as well as for obtaining uncertainty estimates over network predictions [Gal & Ghahramani (2015); Gal (2016); Depeweg et al. (2016, 2017)]. After pruning and hard quantization, and without the need for additional fine-tuning, our method yields a deterministic feed-forward neural network with heavily quantized weights. Our method is applicable to pre-trained networks but can also be used for training from scratch. Target values for quantization can either be manually fixed or they can be learned during training via hierarchical Bayesian inference. We demonstrate our results for the case of ternary quantization on LeNet-5 (MNIST) and DenseNet-121 (CIFAR-10).

![Figure 1: Distribution of weights (means $\theta$ and log-variance $\log \sigma^2$) before and after VNQ training of LeNet-5 on MNIST (validation accuracy before: 99.2% vs. after 195 epochs: 99.31%). Top row: scatter plot of weights (blue dots) per layer. Means were initialized from pre-trained deterministic network, variances with $\log \sigma^2 = -8$. Bottom row: corresponding density. Red shaded areas show the funnel-shaped “basins of attraction” induced by the quantizing prior. Target values for ternary quantization (the codebook) have also been learned. After training, weights with small (absolute) expected value or large variance (inside the area marked by the dotted line, corresponding to $\log \alpha_{ij} \leq \log T_n = 2$) are pruned and remaining weights are quantized without loss in accuracy.](image-url)

2 Preliminaries

2.1 Why Bayes for Compression?

Bayesian inference can be well motivated from an information-theoretic treatment of (lossy) compression [Cover & Thomas (2006); Tishby et al. (2000); Genewein et al. (2015)]. In particular, Bayesian inference automatically penalizes overly complex parametric models, an effect known as “Bayesian Occams Razor” in Bayesian model selection [MacKay (2003); Genewein & Braun (2014)]. The same effect leads to automatic regularization in variational Bayesian inference over model parameters (Grünewald 2007; Graves 2011) (see Molchanov et al. (2017), where the authors show that Sparse Variational Dropout (Sparse VD) successfully prevents a network from fitting an unstructured data, that is a random labeling). This is particularly interesting since regularization is the basis for compression and is thought to be key for generalization [MacKay (2003); Grünewald 2007]. The automatic regularization effect is based on maximizing model evidence, where model parameters are marginalized. A very complex model might have a parameter setting that achieves extremely good likelihood given the data, however, since the model evidence is based on the average or marginal likelihood, overly complex models are penalized for having many parameter settings with poor likelihood. The argument that Bayesian methods search for optimal model structure can also be made from an information-theoretic point-of-view by investigating the equivalence of variational inference and the Minimum description length (MDL) principle [Rissanen (1978); Grünewald (2007); Graves (2011); Louizos et al. (2017)]. The evidence lower bound (ELBO, see Eq. (1)), which is maximized

\[ \log p(x) \leq \log p(x|\theta) + \mathcal{KL}(q(\theta) \| p(\theta|x)) \]

1Kernel density estimate, with radial basis function kernels with a bandwidth of 0.05
in variational inference, is the sum of two terms: one, the average message length required to transmit outputs (labels) to a receiver that knows the inputs and the posterior over model parameters and two, the average message length to transmit the posterior parameters to a receiver that knows the prior over parameters:

\[ \mathcal{L} = \text{neg. reconstr. error} - \mathcal{L}^E + \text{neg. KL divergence} - \mathcal{L}^C = \text{entropy} - \text{cross entropy}, \]

compare Eq. (1). Maximizing the ELBO minimizes the total message length: \( \max \mathcal{L} = \min \mathcal{L}^E + \mathcal{L}^C, \) leading to an optimal trade-off between short description length of the data and the model (thus minimizing the sum of error cost \( \mathcal{L}^E \) and model complexity cost \( \mathcal{L}^C \)). Interestingly, MDL dictates the use of probabilistic models since they are in general “more compressible” compared to deterministic models: high uncertainty over parameters is rewarded by the entropy term in \( \mathcal{L}^C \)—higher uncertainty allows the quantization noise to be higher, thus requiring lower bit-precision for a parameter (the bits back argument \( \text{[Hinton & Van Camp][1993][Louizos et al.[1] 2017]} \)).

2.2 Variational Bayes and Reparameterization

Let \( D \) be a dataset of \( N \) pairs \((x_n, y_n)_{n=1}^{N}\) and \( p(y|x, w) \) be a parameterized model that predicts outputs \( y \) given inputs \( x \) and parameters \( w \). A Bayesian neural network models a (posterior) distribution over parameters \( w \) instead of just a point-estimate. The posterior is given by Bayes’ rule:

\[ p(w|D) = \frac{p(D|w)p(w)}{p(D)}, \]

where \( p(w) \) is the prior over parameters. Computation of the true posterior is in general intractable. Common approaches to approximate inference in neural networks are for instance: MCMC methods pioneered in \( \text{[Neal][1995]} \) and later refined e.g. via stochastic gradient Langevin dynamics \( \text{[Welling & Teh][2011]} \), or variational approximations to the true posterior \( \text{[Graves][2011]} \), Bayes by Backprop \( \text{[Blundell et al.][2015]} \), Expectation Backpropagation \( \text{[Soudry et al][2014]} \), Probabilistic Backprop \( \text{[Hernández-Lobato & Adams][2015]} \). In the latter methods the true posterior is approximated by a parameterized distribution \( q_\phi(w) \). Variational parameters \( \phi \) are optimized by minimizing the Kullback-Leibler divergence between the true and the approximate posterior \( D_{KL}(q_\phi(w)||p(w|D)) \). Since computation of the true posterior is intractable, minimizing this KL divergence is approximately performed by maximizing the so-called “evidence lower bound” (ELBO) or “negative variational free energy” \( \text{[Kingma & Welling][2014]} \):

\[
\mathcal{L}_{\text{SGVB}}(\phi) = \frac{N}{M} \sum_{m=1}^{M} \log p(y_m | \tilde{x}_m, f(\phi, \epsilon_m)) - D_{KL}(q_\phi(w)||p(w))
\]

where we have used the \text{Reparameterization Trick}[Kingma & Welling, 2014] in Eq. (2) to get an unbiased, differentiable, minibatch-based Monte Carlo estimator of the expected log likelihood \( L_{\mathcal{D}}(\phi) \). A mini-batch of data is denoted by \((\tilde{x}_m, y_m)_{m=1}^{M}\). Additionally, and in line with similar work \( \text{[Molchanov et al.][2017][Louizos et al.][2017][Neklyudov et al.][2017]} \), we use the \text{Local Reparameterization Trick}[Kingma et al.[2015] to further reduce variance of the stochastic ELBO gradient estimator, which locally marginalizes weights at each layer and instead samples directly from the distribution over pre-activations (which can be computed analytically). See Appendix A.2 for more details on the Local Reparameterization. Commonly, the prior \( p(w) \) and the parametric form of the posterior \( q_\phi(w) \) are chosen such that the KL divergence term can be computed analytically (e.g. a fully factorized Gaussian prior and posterior, known as the mean-field approximation). Due to the particular choice of prior in our work, a closed-form expression for the KL divergence cannot be obtained but instead we use a differentiable approximation (see Sec. 3.3).

2.3 Variational Dropout

Dropout \( \text{[Srivastava et al.][2014]} \) is a method originally introduced for regularization of neural networks, where activations are stochastically dropped (i.e. set to zero) with a certain probability \( p \). The trick is to use a deterministic differentiable function \( w = f(\phi, \epsilon) \) with \( \epsilon \sim \mathcal{N}(0, 1) \), instead of directly using \( q_\phi(w) \).
during training. It was shown that dropout, i.e. multiplicative noise on inputs, is equivalent to having noisy weights and vice versa (Wang & Manning, 2013; Kingma et al., 2015). Multiplicative Gaussian noise \( \xi_{ij} \sim \mathcal{N}(1, \alpha = \frac{1}{S_{ij}^2}) \) on a weight \( w_{ij} \) induces a Gaussian distribution

\[
w_{ij} = \theta_{ij} \xi_{ij} = \theta_{ij}(1 + \sqrt{\alpha} \epsilon_{ij}) \sim \mathcal{N}(\theta_{ij}, \alpha \theta_{ij}^2)
\]

with \( \epsilon_{ij} \sim \mathcal{N}(0, 1) \). In standard (Gaussian) dropout training, the dropout rates \( \alpha \) (or \( p \) to be precise) are fixed and the expected log likelihood \( \mathcal{L}(\phi) \) (first term in Eq. (1)) is maximized with respect to the means \( \theta \). Kingma et al. (2015) show that Gaussian dropout training is mathematically equivalent to maximizing the ELBO (both terms in Eq. (1)), under a prior \( p(w) \) and fixed \( \alpha \) where the KL term does not depend on \( \theta \):

\[
\mathbb{E}_{q_\phi}[\mathcal{L}(\phi, \theta)] = \mathcal{L}(\alpha, \theta) = D_{KL}(q_\phi(w)||p(w)),
\]

where the dependencies on \( \alpha \) and \( \theta \) of the terms in Eq. (1) have been made explicit. The only prior that meets this requirement is the scale invariant log-uniform prior:

\[
p(\log |w_{ij}|) = \text{const.} \iff p(|w_{ij}|) \propto \frac{1}{|w_{ij}|}.
\]

Using this result, it is straightforward to learn individual dropout-rates \( \alpha_{ij} \) per weight, by including \( \alpha \) into the set of variational parameters \( \phi = (\theta, \alpha) \). This procedure was introduced in (Kingma et al., 2015) under the name “Variational Dropout”. With the choice of a log-uniform prior (Eq. (3)) and a factorized Gaussian approximate posterior \( q_\phi(w_{ij}) = \mathcal{N}(\theta_{ij}, \alpha \theta_{ij}^2) \) (Eq. (3)) the KL term in Eq. (1) is not analytically tractable, but the authors of (Kingma et al., 2015) present an approximation

\[
-D_{KL}(q_\phi(w_{ij})||p(w_{ij})) \approx \text{const.} + 0.5 \log \alpha + c_1 \alpha + c_2 \alpha^2 + c_3 \alpha^3,
\]

see original publication for numerical values of \( c_1, c_2, c_3 \). Note that due to the mean-field approximation, where the posterior over all weights factorizes into a product over individual weights \( q_\phi(w) = \prod q_\phi(w_{ij}) \), the KL divergence factorizes into a sum of individual KL divergences \( D_{KL}(q_\phi(w_{ij})||p(w_{ij})) = \sum D_{KL}(q_\phi(w_{ij})||p(w_{ij})) \).

### 2.4 Pruning Units with Large Dropout Rates

Learning dropout rates is interesting for network compression since neurons or weights with very high dropout rates \( p \to 1 \) can very likely be pruned without loss in accuracy. However, as the authors of Sparse Variational Dropout (sparse VD) (Molchanov et al., 2017) report, the approximation in Eq. (6) is only accurate for \( \alpha < 1 \) (corresponding to \( p \leq 0.5 \)). For this reason, the original variational dropout paper restricted \( \alpha \) to values smaller or equal to 1, which are unsuitable for pruning. Molchanov et al. (2017) propose an improved approximation, which is very accurate on the full range of \( \log \alpha \):

\[
-D_{KL}(q_\phi(w_{ij})||p(w_{ij})) \approx \text{const.} + k_1 \log(1 + \alpha) - 0.5 \log(1 + \alpha^{-1}) = F_{KL,LU}(\theta_{ij}, \sigma_{ij}),
\]

with \( k_1 = 0.63576 \), \( k_2 = 1.87320 \) and \( k_3 = 1.48695 \) and \( S \) denoting the sigmoid function. Additionally, the authors propose to use an additive, instead of a multiplicative noise reparameterization, which significantly reduces variance in the gradient \( \frac{\partial \mathcal{L}_{SGSVB}}{\partial \sigma_{ij}} \) for large \( \alpha_{ij} \). To achieve this, the multiplicative noise term is replaced with an exactly equivalent additive noise term \( \sigma_{ij} \epsilon_{ij} \) with \( \sigma_{ij}^2 = \alpha_{ij} \theta_{ij}^2 \) and the set of variational parameters becomes \( \phi = (\theta, \sigma) \):

\[
w_{ij} = \theta_{ij} \left( 1 + \sqrt{\alpha} \epsilon_{ij} \right) = \theta_{ij} + \sigma_{ij} \epsilon_{ij} \sim \mathcal{N}(\theta_{ij}, \sigma_{ij}^2), \quad \epsilon \sim \mathcal{N}(0, 1).
\]

After Sparse VD training, pruning is performed by thresholding \( \alpha_{ij} = \frac{\sigma_{ij}^2}{\theta_{ij}^2} \), which translates into a threshold for the variance-to-mean ratio (also known as the index of dispersion, a limit-case of the Fano factor). In Molchanov et al. (2017) a threshold of \( \log \alpha \approx 3 \) is used, which roughly corresponds to \( p > 0.95 \). Pruning weights that lie above a threshold of \( T_\alpha \) leads to

\[
\frac{\sigma_{ij}^2}{\theta_{ij}^2} \geq T_\alpha \iff \sigma_{ij}^2 \geq T_\alpha \theta_{ij}^2.
\]
which means effectively that weights with large variance but also weights of lower variance and a mean $\theta_{ij}$ close to zero are pruned. A visualization of the pruning threshold can also be seen in Fig 1 (the “central funnel”, i.e. the area marked by the red dotted lines for a threshold for $T_a = 2$). Sparse VD training can be performed from random initialization or with pre-trained networks by initializing the means $\theta_{ij}$ accordingly. In Bayesian Compression (Louizos et al., 2017) and Structured Bayesian Pruning (Neklyudov et al., 2017), Sparse VD has been extended to include group-sparsity constraints, which allows for pruning of whole neurons or convolutional filters (via learning their corresponding dropout rates).

2.5 Sparsity Inducing Priors

The prior $p(w)$ can be used to induce sparsity into the posterior by having high density at zero and heavy tails. There is a well known family of such distributions: scale-mixtures of normals (Andrews & Mallows, 1974; Louizos et al., 2017; Ingraham & Marks, 2017):

$$w \sim N(0, z^2); \quad z \sim p(z),$$

where the scales of $w$ are random variables. A well-known example is the spike-and-slab prior (Mitchell & Beauchamp, 1988), which has a delta-spike at zero and a slab over the real line. Gal & Ghahramani (2015); Kingma et al. (2015) show how Dropout (Srivastava et al., 2014) implies a spike-and-slab distribution over weights. The log uniform prior used in Sparse VD (Eq. (5)) can also be derived as a marginalized scale-mixture of normals

$$p(w_{ij}) \propto \int \frac{1}{|z|} N(w_{ij}|0, z^2) dz = \frac{1}{|w_{ij}|}; \quad p(z) \propto \frac{1}{|z|}, \quad (10)$$

also known as the normal-Jeffreys prior (Figueiredo, 2002). Louizos et al. (2017) discuss how the log-uniform prior can be seen as a continuous relaxation of the spike-and-slab prior and how the alternative formulation through the normal-Jeffreys distribution can be used to couple the scales of weights that belong together and thus learn dropout rates for whole neurons or convolutional filters, which is the basis for Bayesian Compression (Louizos et al., 2017) and Structured Bayesian Pruning (Neklyudov et al., 2017).

3 Variational Network Quantization

We formulate the preparation of a neural network for a post-training quantization step as a variational inference problem. To this end, we introduce a multi-modal, quantizing prior and train by maximizing the ELBO (Eq. (2)) under a mean-field approximation of the posterior (i.e. a fully factorized Gaussian). The goal of our algorithm is to achieve soft quantization, that is learning a posterior distribution such that the accuracy-loss introduced by post-training quantization is small. Our variational posterior approximation and training procedure is similar to Kingma et al. (2015) and Molchanov et al. (2017) with the crucial difference of using a quantizing prior that drives weights towards the target values for quantization.

3.1 A Quantizing Prior

The log uniform prior (Eq. (5)) can be viewed as a continuous relaxation of the spike-and-slab prior with a spike at location 0 (Louizos et al., 2017). We use this insight to formulate a quantizing prior, a continuous relaxation of a “multi-spike-and-slab” prior which has multiple spikes at locations $c_k, \ k \in \{1..K\}$. Each spike location corresponds to one target value for subsequent quantization. The quantizing prior allows weights of low variance only at the locations of the quantization target values $c_k$. The effect of using such a quantizing prior during Variational Network Quantization is shown in Fig. 1. After training, most weights of low variance are distributed very closely around the quantization target values $c_k$ and can thus be replaced by the corresponding value without significant loss in accuracy. Weights of large variance can be pruned. Additionally, we typically fix one of the spike locations to zero, e.g. $c_2 = 0$, which allows to prune weights with an $\alpha_{ij}$ threshold (see Eq. (9)) as in sparse Variational Dropout (Molchanov et al., 2017). Following the interpretation of the log uniform prior $p(w)$ as a marginal over the scale-hyperparameter $z$, we extend Eq. (10) with a hyper-prior over locations

$$p(w_{ij}) = \int N(w_{ij}|m, z)p_z(z)p_m(m) dzm = \sum_k p_k \delta(m - c_k), \quad (11)$$
with \( p(z) \propto |z|^{-1} \). The location prior \( p_m(m) \) is a mixture of weighted delta distributions located at the quantization values \( c_k \). Marginalizing over \( m \) yields the quantizing prior

\[
p(w_{ij}) \propto \sum_k p_k \int \frac{1}{|z|} \mathcal{N}(w_{ij}|c_k, z) \, dz = \sum_k p_k \frac{1}{|w_{ij} - c_k|}.
\]

In our experiments we use \( K = 3, p_k = 1/K \ \forall k \) and \( c_2 = 0 \) unless indicated otherwise.

### 3.2 Post-Training Quantization

Equation (9) implies that using a threshold on \( \alpha_{ij} \) as a pruning criterion is equivalent to pruning weights whose value does not differ significantly from zero:

\[
\theta_{ij}^2 \leq \frac{\sigma_{ij}^2}{T_\alpha} \Leftrightarrow \theta_{ij} \in (-\frac{\sigma_{ij}}{\sqrt{T_\alpha}}, \frac{\sigma_{ij}}{\sqrt{T_\alpha}}).
\]

To be precise, \( T_\alpha \) specifies the width of a scaled standard-deviation band \( \pm \sigma_{ij}/\sqrt{T_\alpha} \) around the mean \( \theta_{ij} \). If the value zero lies within this band, the weight is assigned the value 0. For instance, a pruning threshold which implies \( p \geq 0.95 \) corresponds to a variance band of approximately \( \sigma_{ij}/4 \).

An equivalent interpretation is that a weight is pruned if the likelihood for the value 0 under the weight posterior exceeds the threshold given by the standard-deviation band (Eq. (13)):

\[
\mathcal{N}(0|\theta_{ij}, \sigma_{ij}^2) \geq \mathcal{N}(\theta_{ij} \pm \frac{\sigma_{ij}}{\sqrt{T_\alpha}}|\theta_{ij}, \sigma_{ij}^2) = \frac{1}{\sqrt{2\pi\sigma_{ij}}} e^{-\frac{1}{2}\left(\frac{\theta_{ij} - \theta_{ij}}{\sigma_{ij}}\right)^2}.
\]

Following this interpretation we can design a maximum a-posteriori (MAP) quantization scheme: to each weight we assign the quantized values \( c_k \) with the highest likelihood under the posterior. Since weight posteriors are Gaussian, this translates into minimizing the squared distance between the mean \( \theta_{ij} \) and the quantized values \( c_k \):

\[
\arg \max_k \mathcal{N}(c_k|\theta_{ij}, \sigma_{ij}^2) = \arg \max_k \exp \left( -\frac{(c_k - \theta_{ij})^2}{2\sigma_{ij}^2} \right) = \arg \min_k (c_k - \theta_{ij})^2
\]

Additionally, the pruning rate can be increased by assigning a hard 0 to all weights that exceed the pruning threshold \( T_\alpha \) (see Eq. (9)) before performing the MAP assignment to quantize the non-pruned weights described above.

### 3.3 KL Divergence Approximation

Under the quantizing prior (Eq. (12)) the KL divergence between the mean-field posterior and prior \( D_{KL}(q_\theta(w)||p(w)) \) is analytically intractable. Similar to Kingma et al. (2015); Molchanov et al. (2017) we use a differentiable approximation \( F_{KL}((\theta, \sigma, c)) \) composed of a small number of differentiable functions to keep the computational effort low during training. We now present the approximation for a reference codebook \( c = [-r, 0, r], r = 0.2 \), however later we show how the approximation can be used for arbitrary ternary, symmetric codebooks as well. The basis of our approximation is the approximation \( F_{KL,LU} \) introduced by Molchanov et al. (2017) for the KL divergence between a log uniform prior and a Gaussian posterior (see Eq. (7)) which is centered around zero. We observe that a weighted mixture of shifted versions of \( F_{KL,LU} \) can be used to approximate the KL divergence for our multi-modal quantizing prior (Eq. (12)) (which is composed of shifted versions of the log uniform prior). In a nutshell, we shift one version of \( F_{KL} \) to each codebook entry \( c_k \) and then use \( \theta \)-dependent Gaussian windowing functions \( \Omega(\theta) \) to mix the shifted approximations (see more details in the Appendix A.3). The approximation for the KL divergence between a Gaussian posterior and our multi-modal quantizing prior is given as

\[
F_{KL}(\theta, \sigma, c) = \sum_{k: c_k \neq 0} \Omega(\theta - c_k)F_{KL,LU}(\theta - c_k, \sigma) + \Omega(\theta)F_{KL,LU}(\theta, \sigma)
\]
with
\[
\Omega(\theta) = \exp\left(-\frac{1}{2} \frac{\theta^2}{\tau^2}\right) \quad \Omega_0(\theta) = 1 - \sum_{k: c_k \neq 0} \Omega(\theta - c_k)
\]  
\tag{17}
\]

We use \(\tau = 0.075\) in our experiments. Illustrations of the approximation, including a comparison against the ground-truth computed via Monte Carlo sampling are shown in Fig. [2]. Over the range of \(\theta\) - and \(\sigma\)-values relevant to our method, the maximum absolute deviation from the ground-truth is 1.07. See Fig. [4] in the Appendix for a more detailed quantitative evaluation of our approximation.

![Figure 2](image)

**Figure 2**: Approximation to the analytically intractable KL divergence \(D_{KL}(q_\phi || p)\), constructed by shifting and mixing known approximations to the KL divergence between the posterior a log uniform prior. Top row: Shifted versions of the known approximation (Eq. (7)) in color and the ground truth KL approximation (computed via Monte Carlo sampling) \(D_{KL}^{MC}(q_\phi || p)\) in black. Middle row: weighting functions \(\Omega(\theta)\) that mix the shifted known approximation to form the final approximation \(F_{KL}\), shown in the bottom row (gold), compared against the ground-truth (MC sampled). Each column corresponds to a different value of \(\sigma\). A comparison between ground-truth and our approximation over a large range of \(\sigma\) and \(\theta\) values is shown in the Appendix in Fig. [4]. Note that since the priors are improper, KL approximation and ground-truth can only be compared up to an additive constant \(C\) - the constant is irrelevant for network training but has been chosen in the plot such that ground-truth and approximation align for large values of \(\theta\).

This KL approximation in Eq. (16), developed for the reference codebook \(c_r = [-r, 0, r]\), can be reused for any symmetric ternary codebook \(c_a = [-a, 0, a]\), \(a \in \mathbb{R}^+\), since \(c_a\) can be represented with the reference codebook and a positive scaling factor \(s\), \(c_a = sc_r\), \(s = a/r\). As derived in the Appendix (A.4), this re-scaling translates into a multiplicative re-scaling of the variational parameters \(\theta\) and \(\sigma\). The KL divergence between the posterior \(q_\phi(w)\) and a prior based on the codebook \(c_a\) is thus given by \(D_{KL}(q_\phi(w)||p_{c_a}(w)) \approx F_{KL}(\theta/s, \sigma/s, c_r)\). This result allows learning the quantization level \(a\) during training as well.

4 **Experiments**

In our experiments we train with VNQ and then first prune via thresholding \(\log \alpha_{ij} \geq \log T_a = 3\). Remaining weights are then quantized by minimizing the squared distance to the quantization values \(c_k\) (corresponding to a MAP quantization, see Sec. 3.2). We use warm-up \(\theta\) boosting \citep{Sønderby2016}, that is we multiply the KL divergence term (Eq. (4)) with a factor \(\beta\), where \(\beta = 0\) during the first few epochs and then linearly ramps up to \(\beta = 1\). To improve stability of VNQ training we ensure through clipping that \(\log \sigma_{ij}^2 \in (-10, 1)\) and \(\theta_{ij} \in (0.223 \sigma \pm a)\) (which corresponds to a shifted \(\log \alpha\) threshold of 3, that is we clip \(\theta_{ij}\) if it lies left of the \(-a\) funnel or right of the \(+a\) funnel, compare Fig. [1]). When learning codebook values \(a\) during training, we use a lower learning rate for
adjusting the codebook, otherwise we observe a tendency for codebook values to collapse in early stages of training (a similar observation was made by Ullrich et al. (2017)). Additionally, we ensure $a \leq 0.05$ by clipping.

### 4.1 LeNet-5 on MNIST

We demonstrate our method with LeNet-5 \citep{LeCun1998} on the MNIST handwritten digits dataset. Images are pre-processed by subtracting the mean and dividing by the standard-deviation over the training set. For the pre-trained network we run 5 epochs on a randomly initialized network (Glorot initialization, Adam optimizer), which leads to a validation accuracy of 99.2\%. We initialize means $\theta$ with the pre-trained weights and variances with $\log \sigma^2 = -8$. The warm-up factor $\beta$ is linearly increased from 0 to 1 during the first 15 epochs. VNQ training runs for a total of 195 epochs with a batch-size of 128, the learning rate is linearly decreased from 0.001 to 0 and the learning rate for adjusting the codebook parameter $a$ uses a learning rate that is 100 times lower. Results are shown in Table 1, a visualization of the distribution over weights after VNQ training is shown in Fig. 1.

| Method                  | val. error | $|w|/|\theta| \%- | bits |
|-------------------------|------------|-----------------|------|
| Original                | 0.8        | 100             | 32   |
| VNQ (no P\&Q)           | 0.69       | 100             | 32   |
| VNQ + P\&Q              | 0.83       | 24.5            | 2    |
| Deep Compression (P\&Q) | 0.74       | 8               | 10-13|
| Soft weight-sharing (P\&Q) | 0.97     | 3               | 3    |
| Sparse VD (P)           | 0.75       | 0.7             | -    |
| Bayesian Comp. (P)      | 1.0        | 0.6             | 7-18 |
| Structured BP (P)       | 0.86       | -               | -    |

We find that VNQ training sufficiently prepares a network for pruning and quantization with negligible loss in accuracy and without requiring subsequent fine-tuning. Compared to pruning methods that do not consider few-bit quantization in their objective, we achieve significantly lower pruning rates. This is an interesting observation since our method is based on a similar objective (e.g. compared to Sparse VD) but with the addition of forcing non-pruned weights to tightly cluster around the quantization levels. Few-bit quantization severely limits network capacity. Perhaps this capacity limitation must be countered by pruning fewer weights. Our pruning rates are roughly in line with other papers on ternary quantization, e.g. \cite{Zhu2016}, who report sparsity levels between 30\% and 50\% with their ternary quantization method. Note that a direct comparison between pruning, quantizing and ternarizing methods is difficult and depends on many factors such that a fair computation of the compression rate that does not implicitly favor certain methods is hardly possible within the scope of this paper. For instance, compression rates for pruning methods are typically reported under the assumption of a CSC storage format which would not fully account for the compression potential of a sparse ternary matrix. We thus chose not to report any measures for compression rates, however for the methods listed in Table 1 they can easily be found in the literature.

### 4.2 DenseNet-121 on CIFAR-10

Our second experiment uses a modern DenseNet-121 \citep{Huang2017} (k=12, with bottleneck) on CIFAR-10 \citep{Krizhevsky2009}. The training procedure is identical to the procedure on MNIST with the following exceptions: we use a batch-size of 64 epochs, warmup is linearly ramped

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4 the Caffe version, see \url{https://github.com/BVLC/caffe/blob/master/examples/mnist/lenet_train_test.prototxt}
up from 0 to 1 over the first 20 epochs, the learning rate of 0.005 is kept constant for the first 50 epochs and then linearly decreased until training stops at epoch 300. Results are shown from epoch 150. We pre-train a deterministic DenseNet (reaching validation accuracy of 93.19\%) to initialize VNQ training. Results are shown in Table 2. A visualization of the distribution over weights after VNQ training is shown in the Appendix Fig. 3.

Table 2: Results on DenseNet-121 (CIFAR-10), showing the error on the validation set, the percentage of non-pruned weights and the bit-precision per parameter. Original denotes the pre-trained network. We show results after VNQ training without pruning and quantization (weights were deterministically replaced by the (full-precision) means $\theta$), and VNQ with subsequent pruning and quantization (in the condition (w/o 1) we use full-precision means for the weights in the first layer and do not prune and quantize this layer).

| Method                   | val error [%] | $|w|\neq0$ [%] | $|w|$ bits |
|--------------------------|---------------|---------------|-----------|
| Original                 | 6.81          | 100           | 32        |
| VNQ (no P&Q)             | 8.45          | 100           | 32        |
| VNQ + P&Q (w/o 1)        | 8.52          | 55            | 2 (32)    |
| VNQ + P&Q                | 10.92         | 55            | 2         |

We generally observe lower levels of sparsity for DenseNet, compared to LeNet. This might be due to the fact that DenseNet already has an optimized architecture which removed a lot of redundant parameters from the start. In line with previous publications we find that the first and last layer of the network are most sensitive to pruning and quantization. However, in contrast to many other methods that do not quantize these layers (e.g. Zhu et al. (2016)), we find that after sufficient training, the final layer can be pruned and quantized without loss in accuracy and the first layer can also be pruned and quantized with a small loss in accuracy (see Table 2).

5 RELATED WORK

Our method is an extension of Sparse VD (Molchanov et al., 2017), originally used for network pruning. In contrast we use a quantizing prior, leading multi-modal posterior suitable for few-bit quantization and pruning. Bayesian Compression and Structured Bayesian Pruning (Louizos et al., 2017; Neklyudov et al., 2017) extend Sparse VD to prune whole neurons or filters via group-sparsity constraints. Additionally, in (Louizos et al., 2017) the required bit-precision per layer is determined via posterior uncertainty. In contrast to our method, Bayesian Compression does not explicitly enforce clustering of weights during training and thus requires bit-widths in the range between 5 and 18 bits. Extending our method to include group-constraints for pruning is an interesting direction for future work. Another Bayesian method for simultaneous network quantization and pruning is soft weight-sharing (SWS) (Ullrich et al., 2017), which uses a Gaussian mixture model prior (and a KL term without trainable parameters such that the KL term reduces to the prior entropy). SWS acts like a probabilistic version of k-means clustering with the advantage of automatic collapse of unnecessary mixture components. Similar to learning the codebooks in our method, soft weight-sharing learns the prior from the data, a technique known as empirical Bayes (see also ARD (Karaletsos & Rätsch, 2015)). We cannot directly compare against soft weight-sharing since the authors do not report results on ternary networks. (Gal et al., 2017) learn dropout rates by using a continuous relaxation of dropout’s discrete masks (via the concrete distribution). The authors learn layer-wise dropout rates, which does not allow for dropout-rate-based pruning. We have experimented with using the concrete distribution for learning codebooks for quantization with promising early results but we have generally observed lower pruning rates or lower accuracy compared to VNQ. A non-probabilistic state-of-the-art method for network ternarization is Trained Ternary Quantization (Zhu et al., 2016) which uses full-precision shadow weights during training, but quantized forward passes. Additionally it learns a (non-symmetric) scaling per layer for the non-zero quantization values, similar to our learned quantization level $a$. While the method achieves impressive accuracy, the sparsity and thus pruning rates are rather low (between 50\% and 70\% sparsity) and the first and last layer need to be kept with full precision.
6 DISCUSSION

A potential shortcoming of our method is the KL divergence approximation (Sec. 3.3). While the approximation is reasonably good on the relevant range of $\theta$- and $\sigma$-values, there is still room for improvement which could have the benefit that weights are drawn even more tightly onto the quantization levels, resulting in lower accuracy loss after quantization and pruning. Since the KL approximation only needs to be computed once and an arbitrary amount of ground-truth data can be produced, it should be possible to improve upon the approximation presented here, at least by some brute-force function approximation, e.g. a neural network, polynomial or kernel regression. The main difficulty is that the resulting approximation must be differentiable and must not introduce significant computational overhead since the approximation is evaluated once for each network parameter in each gradient step.

Compared to similar methods that only consider network pruning, our pruning rates are significantly lower. This does not seem to be a particular problem of our method since other papers on network ternarization report similar sparsity levels (Li et al. (2016) roughly achieve between 30% and 50% sparsity). The reason for this might be that heavily quantized networks have much lower capacity compared to full-precision networks. This limited capacity might require that the network compensates by effectively using more weights such that the pruning rates become significantly lower. Similar trends have also been observed with binary networks, where drops in accuracy could be prevented by increasing the number of neurons (with binary weights) per layer. Principled experiments to test the trade-off between low bit-precision and sparsity rates would be an interesting direction for future work. One starting point could be to test our method with more quantization levels (e.g. 5, 7 or 9) and investigate how this affects the pruning rate.

REFERENCES


Figure 3: Visualization of distribution over DenseNet-121 weights after training on CIFAR-10 with VNQ. Each panel shows one layer, starting in the top-left corner with the input and ending with the final layer in the bottom-right panel (going row-wise, that is first moving to the right as layers increase). The validation accuracy of the network shown is 91.2\%.
A.2 Local Reparameterization

We follow Sparse VD [Molchanov et al. (2017)] and use the Local Reparametrization trick [Kingma et al. (2015)] and Additive Noise Reparametrization to optimize the stochastic gradient variational lower bound $L_{SVGD}$ (Eq. (2)). We optimize posterior means and log-variances $(\theta, \log \sigma^2)$ and in some experiments additionally the codebook level $\alpha$. We apply Variational Network Quantization to fully connected and convolutional layers. Denoting inputs to a layer with $A^{M \times l}$ outputs of a layer with $B^{M \times O}$ and using local reparametrization we get:

$$b_{mj} \sim \mathcal{N}(\gamma_{mj}, \delta_{mj}); \; \gamma_{mj} = \sum_{i=1}^{l} a_{mi} \theta_{ij}, \; \delta_{mj} = \sum_{i=1}^{l} \sigma_{mi}^2 \sigma_{ij}^2$$

for a fully connected layer. Similarly activations for a convolutional layer are computed as follows

$$\text{vec}(b_{mk}) \sim \mathcal{N}(\gamma_{mk}, \delta_{mk}); \; \gamma_{mk} = \text{vec}(A_{m} * \theta_{k}), \; \delta_{mk} = \text{diag}(\text{vec}(A_{m}^2 * \sigma_{k}^2)),$$

where $(\cdot)^2$ denotes an element-wise operation, $\ast$ is the convolution operation and $\text{vec}(\cdot)$ denotes reshaping of a matrix/tensor into a vector.

A.3 KL Approximation for Quantizing Prior

Under the quantizing prior (Eq. (12)) the KL divergence between the mean-field posterior and prior $D_{KL}(q_{\theta}(w_{ij})||p(w_{ij}))$ is analytically intractable. [Molchanov et al. (2017)] presented an approximation for the KL divergence under a (zero-centered) log uniform prior (Eq. (5)). Since our quantizing prior is essentially a composition of shifted log uniform priors, we construct a composition of the approximation for the KL divergence under a (zero-centered) log uniform prior (Eq. (5)).

$$D_{KL}(q_{\theta}(w_{ij})||p(w_{ij})) = D_{KL}(q_{\theta}(\gamma_{mj}, \delta_{mj})||p(\gamma_{mj}, \delta_{mj}))$$

After the shift to the posterior parameter $\theta$

$$D_{KL}(q_{\theta}(\gamma_{mj}, \delta_{mj})||p(\gamma_{mj}, \delta_{mj})) = D_{KL}(q_{\theta}(\gamma_{mj} - \gamma_{mj}, \delta_{mj})||p(\gamma_{mj} - \gamma_{mj}, \delta_{mj})) + \tilde{C} \quad (18)$$

For small posterior variances $\sigma_{ij}^2$ ($\sigma_{ij} \ll r)$ and means near the quantization levels (i.e., $|\theta_{ij} \approx r$), the KL divergence is dominated by the mixture prior component located at the respective quantization level $r$. For these values of $\theta$ and $\sigma$, the KL divergence can be approximated by shifting the approximation $F_{LU,KL}(\theta, \sigma)$ to the quantization level $r$, i.e. $F_{LU,KL}(\theta \pm r, \sigma)$. For small $\sigma$ and values of $\theta$ near zero or far away from any quantization level, as well as for large values of $\sigma$ and arbitrary $\theta$, the KL divergence can be approximated by the original non-shifted approximation $F_{LU,KL}(\theta, \sigma)$. Based on these observations we construct our KL approximation by properly mixing shifted versions of $F_{LU,KL}(\theta \pm r, \sigma)$. We use Gaussian window functions $\Omega(\theta \pm r)$ to perform this weighting (to ensure differentiability). The remaining $\theta$ domain is covered by an approximation located at zero and weighted such that this approximation is dominant near zero and far away from the quantization levels, which is achieved by introducing the constraint that all window functions sum up to one on the full $\theta$ domain. See Fig. 2 for a visual representation of shifted approximations and their respective window functions.

We evaluate the quality of our KL approximation (Eq. (16)) by comparing against a ground-truth Monte Carlo approximation on a dense grid over the full range of relevant $\theta$ and $\sigma$ values. Results of this comparison are shown in Fig. 4。

A.4 Reusing the KL Approximation for Arbitrary Codebooks

We show that the KL approximation (Eq. (16)), developed for a fixed reference codebook, can be reused for arbitrary codebooks as long as codebook learning is restricted to learning a multiplicative scaling factor. Without loss of generality we consider the case of ternary, symmetric codebooks:

$$c_r = [-r, 0, r]; \quad p_{c_r}(w) \propto \sum_{k=1}^{3} \frac{p_{c_r}(w - c_{r,k})}{|w - c_{r,k}|} \quad (19)$$

Note that indices $ij$ have been dropped for notational brevity from the whole section. However, throughout the section we refer to individual weights $w_{ij}$ and their variational parameters $\theta_{ij}$ and $\sigma_{ij}$.
where $r \in \mathbb{R}^+$ is the quantization level value and $p_{cr}$ denotes a sparsity-inducing, quantizing prior over weights (sparsity is induced because one of the codebook entries is fixed to 0). We denote $c_r$ as the reference codebook for which we design the KL approximation $D_{KL}(q_\phi(w)||p_{cr}) = F_{KL}(\theta, \sigma, c_r)$ (Eq. (16)). This approximation can be reused for any symmetric ternary codebook $c_a = [-a, 0, a]$ with quantization level $a \in \mathbb{R^+}$. The latter can be seen by representing $c_a$ with the reference codebook and a positive scaling factor $s > 0$ as $c_a = sc_r$, $s = a/r$. This re-scaling translates into a multiplicative re-scaling of the variational parameters $\theta$ and $\sigma$. To see this, consider the prior $p_{ca}$, based on codebook $c_a$:

$$
p_{ca}(w) = \frac{1}{Z} \sum_{k=1}^{3} \frac{p_k}{|w - c_{a,k}|} = \frac{1}{Z} \sum_{k=1}^{3} \frac{p_k}{|w - sc_{r,k}|}.
$$

(20)

The KL divergence between the posterior $q_\phi(w)$ and a prior based on the codebook $c_a$ is given by

$$
D_{KL}(q_\phi(w)||p_{ca}(w)) = \int q_\phi(w) \log \frac{q_\phi(w)}{\frac{1}{s} \sum_{k=1}^{3} \frac{p_k}{|w - sc_{r,k}|}} \ dw + C
$$

$$
= \int q_\phi(sz) \log \frac{q_\phi(sz)}{\frac{1}{s} \sum_{k=1}^{3} \frac{p_k}{|z - c_{r,k}|}} \ dz + C.
$$

(21)

Since $q_\phi(sz)$ is Gaussian, the scaling $s$ can be transferred into the variational parameters $\phi = (\theta, \sigma)$:

$$
q_\phi(sz) = \mathcal{N}(sz; \theta, \sigma^2) = \frac{1}{s} \mathcal{N}(z; \frac{\theta}{s}, \frac{\sigma^2}{s^2}) = \frac{1}{s} q_\phi(z).
$$
with \( \hat{\phi} = (\frac{\theta}{s}, \frac{\sigma}{s}) \). Inserting into Eq. \((21)\) yields:

\[
D_{KL}(q_{\hat{\phi}}(w)||p_{ca}(w)) = \int \frac{1}{s} q_{\hat{\phi}}(z) \log \frac{1}{s} \sum_{k=1}^{3} \frac{p_k}{|z-c_{r,k}|} \, sdz + C.
\]

\[
= \int q_{\hat{\phi}}(z) \log \sum_{k=1}^{3} \frac{p_k}{|z-c_{r,k}|} \, dz + C.
\]

\[
= D_{KL}(q_{\hat{\phi}}(w)||p_{ca}(w)). \tag{22}
\]

Thus, \( D_{KL}(q_{\hat{\phi}}(w)||p_{ca}(w)) = D_{KL}(q_{\hat{\phi}}(w)||p_{ca}(w)) \approx F_{KL}(\theta/s, \sigma/s, c_r) \), where \( F_{KL} \) is given by Eq. \((16)\). This means that the KL approximation can be used for arbitrary ternary, symmetric codebooks of the form \( c_a = [-a, 0, a] = sc_r \) because the scaling factor \( s \) translates into a re-scaling of the variational parameters \( \hat{\phi} = (\frac{\theta}{s}, \frac{\sigma}{s}) \).