Graph Reparameterizations for Enabling 1000+ Monte Carlo Iterations in Bayesian Deep Neural Networks

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Abstract

Uncertainty estimation in deep models is essential in many real-world applications and has benefited from developments over the last several years. Recent evidence [Farquhar et al., 2020] suggests that existing solutions dependent on simple Gaussian formulations may not be sufficient. However, moving to other distributions necessitates Monte Carlo (MC) sampling to estimate quantities such as the KL divergence: it could be expensive and scales poorly as the dimensions of both the input data and the model grow. This is directly related to the structure of the computation graph, which can grow linearly as a function of the number of MC samples needed. Here, we construct a framework to describe these computation graphs, and identify probability families where the graph size can be **independent** or only weakly dependent on the number of MC samples. These families correspond directly to large classes of distributions. Empirically, we can run a much larger number of iterations for MC approximations for larger architectures used in computer vision with gains in performance measured in confident accuracy, stability of training, memory and training time.

1 INTRODUCTION

Motivated by the need to provide measures of uncertainty in the deployment of deep neural networks in mission critical and medical applications, there has been a strong recent interest in deep Bayesian learning. While deep Bayesian learning provides many methods to estimate posterior distributions, Variational Inference (VI) is a convenient choice for many problem settings [Blundell et al., 2015]. Many libraries such as Tensorflow Probability [Dillon et al., 2017] are also now available that offer a rich set of features.

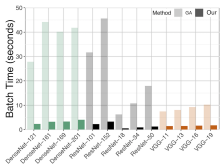
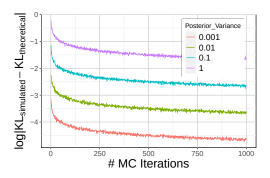


Figure 1: MC sampling is significantly slower in existing neural network libraries incorporating Gradient Accumulation (GA). In contrast, our proposed MC reparameterization reduces the compute time up to $14\times$ for some networks.

Denote the observed data as (x,y), where x is an input to the network, and y is a corresponding response (in autoencoder settings we may have y=x). When using VI in Bayesian Neural Networks (BNNs), one considers all weights $W=(W^1,\ldots,W^D)$ as a random vector and approximates the true unknown posterior distribution P(W|y,x) with an approximate posterior distribution Q_{θ} of our choice, which depends on learned parameters θ . Let $W_{\theta}=(W^1_{\theta},\ldots,W^D_{\theta})$ denote a random vector with a distribution Q_{θ} and pdf q_{θ} . VI seeks to find θ such that Q_{θ} is as close as possible to the real (unknown) posterior P(W|y,x), accomplished by minimizing the KL divergence between Q_{θ} and P(W|y,x). Given a prior pdf of weights p, along with a likelihood term p(y|W,x), and a common mean field assumption of independence for W^d and W^d_{θ} , for $d \in 1, \ldots, D$, i.e. $p(W) = \prod_{d=1}^D p^d(W^d)$ and $q_{\theta}(W_{\theta}) = \prod_{d=1}^D q^d_{\theta}(W^d_{\theta})$,

$$\boldsymbol{\theta}^* = \arg\min_{\boldsymbol{\theta}} KL(q_{\boldsymbol{\theta}}||p) - \mathbb{E}_{q_{\boldsymbol{\theta}}} \left[\ln p(y|W,x) \right]$$
 (1)

$$KL\left(q_{\theta}||p\right) = \sum_{d=1}^{D} \mathbb{E}_{q_{\theta}^{d}}\left[\ln q_{\theta}^{d}(w)\right] - \mathbb{E}_{q_{\theta}^{d}}\left[\ln p^{d}(w)\right] \quad (2)$$



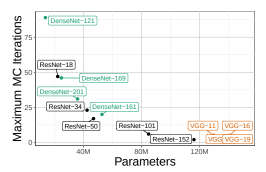


Figure 2: (left) Approximation error (\log) of simulated KL for the single-parameter Bayesian Neural Network at different variance values of approximate posterior distribution, (**right**) Maximum number of feasible MC iterations required for training Bayesian versions of different neural networks on a single GPU.

A key consideration in VI is the choice of prior p and the approximate posterior q_{θ} . This choice does not drastically change the computation of the likelihood term p(y|W,x)which is influenced more by the problem and the complexity of the network instead of W (e.g., it is Gaussian for regression problems). But it strongly impacts the computation of KL term. For example, a common choice for p, and q_{θ} is Gaussian, which allows calculating (2) in a closed form. However, there is emerging evidence [Farguhar et al., 2020, Fortuin et al., 2020] that the Gaussian assumption may not work well on medium/large scale Bayesian NNs. [Farquhar et al., 2020] attributes this to the probability mass in high-dimensional Gaussian distributions concentrating in a narrow "soap-bubble" far from the mean. Choosing a correct distribution is an open problem [Ghosh and Doshi-Velez, 2017, Farquhar et al., 2020, McGregor et al., 2019, Krishnan et al., 2019], and unfortunately, more complex distributions frequently lack closed form solutions for (2).

Numerical approximations. When the integrals for these expectations cannot be solved in closed form, an approximation is used [Ranganath et al., 2014, Paisley et al., 2012, Miller et al., 2017]. One strategy is Monte Carlo (MC) sampling, which gives an unbiased estimator with variance $O(\frac{1}{M})$ where M is number of samples. For a function $g(\cdot)$:

$$\mathbb{E}_{q_{\theta}}\left[g(w)\right] = \int g(w)q_{\theta}(w)dw \approx \frac{1}{M} \sum_{i=1}^{M} g(w_{i}),$$
where $w_{i} \sim Q_{\theta}$. (3)

Expected value terms in (2) can be estimated by applying the scheme in (3) and in fact, even if a closed form expression can be computed, with enough samples an MC approximation may perform similarly [Blundell et al., 2015]. Unfortunately, MC procedures are costly, and may need many samples (i.e., iterations) for a good estimation as the model size grows: [Miller et al., 2017] shows this relationship for small networks, and demonstrates that using fewer samples leads to large variances in the approximation. In general, for deep BNNs, computation of both KL and expectation of

log-likelihood requires numerical approximation with MC sampling, but for now, we will only focus on the KL term.

How does M affect the KL approximation necessary for large scale VI? Consider a standard Gaussian distribution for the approximate posterior q_{θ} and prior p for the weights of an arbitrary BNN, and also consider an MC approximation of the KL term in (2). In this case, we have a closed form solution for KL, which allows checking the approximation quality: the gap between the MC approximation \widehat{KL} and the closed form KL.

(a) Figure 2 (left) shows this gap for different variances of the approximate posterior for a BNN. While decreasing the variance of the posterior distribution indeed reduces the variance of an estimator, with such a small variance on weights, the model is essentially deterministic. Clearly by increasing M, we decrease the error. However, in current DNNs, increasing the number of MC iterations not only slows down computation, but severely limits GPU memory. **(b)** Figure 2 (right) presents the maximum number of iterations possible on a single GPU (Nvidia 2080 TI) with a direct implementation of MC approximation for Bayesian versions of popular DNN architectures: ResNet, DenseNet and VGG (more details in §3). Extrapolating Figure 2, we see clearly that Bayes versions of these networks will result in large variances. This raises the question: is there a way to increase the number of MC iterations for deep networks without sacrificing performance, memory, or time?

Contributions. This work makes two contributions. (a) We propose a new framework to construct an MC estimator for the KL term, which significantly decreases GPU memory needs and improves runtime. Memory savings allow us to run up to $1000 \times$ more MC iterations on a single GPU, resulting in smaller variances of the MC estimators, improving both training convergence and final accuracy, especially on subsets of data where the model is not confident. We show feasibility for popular architectures including ResNets [He et al., 2016], DenseNets [Huang et al., 2017], VGG [Simonyan and Zisserman, 2014] and U-Net [Ronneberger

Figure 3: Proposed MC reparameterization presented as an API. Only a minimal change in an existing programming interface is required to incorporate our method. See the supplement for details.

et al., 2015] – strategies for successfully training Bayesian versions of many of these (deep) networks remain limited [Dusenberry et al., 2020]. (b) From the user perspective, we provide a simple interface for implementing and estimating BNNs (Figure. 3). (c) On the technical side, we obtain a scheme under which we can determine whether our *reparameterization* can be applied. The result covers a broad class of distributions used in VI as an approximate posterior and prior. Inspired by the Pitman–Koopman–Darmois theorem [Koopman, 1936], we show that our method is effective when an exponential family is used as a prior on weights in deep BNNs estimated via VI, and the approximate posterior is modeled as location-scale or certain other distributions, expanding the range of distributions that can be used.

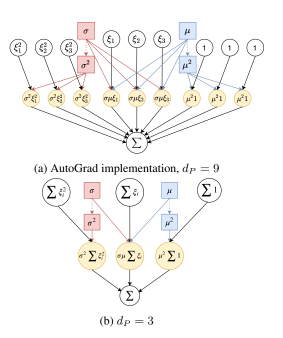


Figure 4: Two computation graphs of the same MC expression $\sum_{i=1}^3 (\mu + \sigma \xi_i)^2$, with different parameterizations. Filled squares represent elements of the vector function $n(\theta)$, clear circles represent functions of auxiliary variables $t(\xi)$, yellow circles represent the Hadamard product $n(\theta) \circ t(\xi)$. Clearly, parameterization affects the size of the graph, and there exists parameterization (b) where the size is independent of the number of MC iterations M. Note: We slightly modify the computational graph presentation for space and clarity. Actual computation graphs from PyTorch convey the same message.

2 RELATED WORK

In addition to VI, the literature provides a broad range of ways to estimate posterior predictive distributions. Ensemble methods [Lakshminarayanan et al., 2017, Pearce et al., 2018, Newton et al., 2018] can be applied to common networks with minimal modifications; however, they require many forward passes, often similar in terms of space/time to a standard gradient accumulation schemes (we provide a PyTorch code snippet in Figure 5). Figure 1 provides experimental results showing that gradient accumulation is much slower. Other methods like Deterministic Variational Inference [Wu et al., 2019] and Probabilistic Backpropagation [Hernández-Lobato and Adams, 2015], improve over naïve MC implementations of VI, but often approximate the posterior of a neural network with a Gaussian distribution. However, [Farquhar et al., 2020] shows that Gaussians are sensitive to hyperparameter choices, among other problems during training. For this reason, a non-Gaussian distribution can be used as an approximate posterior in the traditional VI setup, but its lack of a closed form solution for the KLterm ends up needing MC approximation. This is where our proposal offers value. Also, note some other issues that emerge in Deterministic Variational Inference and Probabilistic Backpropagation: (a) the methods need non-trivial modification of the network to perform a moment matching and (b) replacing the Gaussian assumption with another distribution requires new analytical solutions of closed forms. This is more complicated than a MC approximation.

Our work is distinct from other works that also target MC estimation in neural networks. For example, one may seek to derive new estimators with an explicit goal of variance reduction (e.g., [Miller et al., 2017]). Here, we do <u>not</u> obtain a new estimator replacing the MC procedure with a smaller variance procedure. Instead, we study a scheme that makes the computation graph mostly independent of the number of samples, and is applicable to ideas such as those in [Miller et al., 2017] as well.

```
optimizer.zero_grad()
for _ in range(n_mc_iter):
    output = model(inputs)
    loss = computeLoss(output, targets)
    loss.backward()
optimizer.step()
```

Figure 5: PyTorch implementation of "gradient accumulation" technique, a standard method to collect gradient from several different forward passes. Memory consumption is equivalent to 1 forward pass, but time complexity is proportional to number of forward passes.

Sampling: $W(\theta, \xi)$	Approximate Posterior p.d.f. q_{θ}		Prior p.d.f. $p(w)$		
Scaling property family: $W(\theta, \xi) = \theta \xi$ and related – Corollary 1	Exponential(θ) Rayleigh(θ)	Standard Wald(θ) Weibull(k, θ)	Exponential Dirichlet	Standard Wald Chi-squared	Rayleigh Pareto
	Erlang (k, θ)	$Gamma(k, \theta)$	Inverse-Gamma	Gamma	Erlang
and related coronary r	Error (a, θ, c)	$Log-Gamma(k, \theta)$	Log-normal	Error	Weibull
	Inverse-Gamma (k, θ)		Inverse-Gaussian	Normal	
Location-Scale family:	Normal(μ, σ)	Laplace (μ, σ)	Logistic	Exponential	Normal
$W(\theta, \xi) = \mu + \sigma \xi,$	Logistic(μ, σ)	Horseshoe(μ, σ)		Laplace	
$\theta = (\mu, \sigma)$	Radial(μ, σ)		Normal variations, e.g., Horseshoe, Radial		
Corollary 2	Log-Normal(μ, σ)		Dirichlet	Pareto	

Table 1: Summary list of approximate posterior distributions q_{θ} and priors p(w), which allows to define a parameterization tuple P for MC estimation, such that d_P is independent of M. For every cell in "Sampling: $W(\theta, \xi)$ " we can select any combination of q_{θ} and p(w). Reference: Radial [Farquhar et al., 2020], Horseshoe [Ghosh and Doshi-Velez, 2017].

3 COMPUTATION GRAPHS FOR MC ITERATIONS

Despite the ability to approximate the expectation in principle, the minimization in (1) via (3) is difficult for common architectures, and relies on gradient computations at each iterate. Standard implementations make use of automatic differentiation based on computation graphs [Griewank, 2012]. Computation graphs are directed acyclic graphs, where nodes are the inputs/outputs and edges are the operations. If there is a single input to an operation that requires a gradient, its output will also require a gradient. As noted in PyTorch manual (cf. Autograd mechanics), a backward computation is never performed for subgraphs where no nodes require gradients. This allows us to replace such a subgraph with one output node and to define the size of the computation graph as the minimal number of nodes necessary to perform backpropagation: the number of nodes which require gradients. Modern neural networks lead to graphs where the number of nodes range from a few hundred to millions. To define the size of a graph, accounting for the probabilistic nature of the MC approximation, we propose the following construction.

Definition 1 Consider w as sampled based on a parameter θ and an ancillary random variable ξ , i.e., $w = W(\theta, \xi)$. If there exist functions G, n, and t such that a function $F(w_1, \ldots, w_n)$ can be expressed as $G(n(\theta) \circ t(\xi_1, \ldots, \xi_n))$, then we say P := (G, n, t) is a **parameterization tuple** for the function F, where \circ is the Hadamard product. Let d_P be the dimension of $n \circ t$, corresponding to the number of nodes requiring gradients with respect to θ .

To demonstrate the application of the Def. 1, as an example, consider the computation graph for the MC approximation of the function $g(w)=w^2$ in (3) and given one weight $W_{\theta} \sim N(\mu, \sigma^2)$. Applying the reparameterization trick: $W_{\theta} = \mu + \sigma \xi, \, \xi \sim N(0, 1)$, the Python form is,

```
for i in range(M):
    # sample 1 observation from N(0, 1)
    sample = sampler_normal.sample()
    w = mu * 1 + sigma * sample
    loss += w^2 / M
```

The computation graph, a function of both the parameters $\theta=(\mu,\sigma)$ and of the auxiliary samples $\xi_1,\ \xi_2,$ and $\xi_3,$ generated by PyTorch/AutoGrad for $\mathbf{M}=\mathbf{3}$ iterations of this loop is shown in Figure 4a. According to Def. 1, $d_P=9$ and

$$n(\theta) = (\mu^2, 2\sigma\mu, \sigma^2, \mu^2, 2\sigma\mu, \sigma^2, \mu^2, 2\sigma\mu, \sigma^2),$$

$$t(\xi) = (1, \xi_1, \xi_1^2, 1, \xi_2, \xi_2^2, 1, \xi_3, \xi_3^2),$$

$$G(n(\theta) \circ t(\xi)) = n_1(\theta)t_1(\xi) + \dots + n_9(\theta)t_9(\xi)$$

Naïvely, the graph size grows linearly O(M) with the number of MC iterations, as in the direct implementation (Fig. 4a). For Bayesian VI in DNNs, this is a problem. We need to perform MC approximations of KL terms at every layer. Also, [Miller et al., 2017] shows that iterating over a large number of samples M might be important for convergence. This constrains model sizes given limited hardware resources. One might suspect that a "for" loop is a poor way to evaluate this expectation and instead the expression should be vectorized. Indeed, creating a vector of size M and summing it will clearly help runtime. But the loop does not change the computation graph; all trainable parameters maintain the same corresponding connections to samples, and rapidly exhaust memory.

But graphs for the same function can be constructed differently (see Fig. 4b). For the right parameterization tuple P, we can achieve $d_P=3$. This leads us to,

Remark 1 For computation graph of MC approximation $\sum_{i=1}^{M} g(w_i)$ and specific g, there exists a parameterization tuple P = (G, n, t), such that d_P is independent of M.

For which class of distributions Q_{θ} and functions $g(\cdot)$ can we always construct reparameterizations of the MC estimation (3), maintaining the size of the computation graph d_P as **independent** of number of iterations M? We explore this in the next section.

4 MC REPARAMETERIZATION ENABLES FEASIBLE TRAINING

Our approach is partly inspired by a vast literature on known distributional families and their use within VI. For example, in VI, commonly one chooses distributions that fall within *exponential families* (e.g., Gaussian, Laplace, Horseshoe). With this assumption on the prior, we can express

$$p(w;\zeta) = h(w)\exp(\eta(\zeta)'T(w) - A(\zeta)) \tag{4}$$

where ζ is a parameter defining w. The sufficient statistics T(w) and natural parameters $\eta(\zeta)$ completely define a specific distribution.

Relevance of PKD theorem. While the foregoing discussion links our approach to well-known statistical concepts, it does not directly yield our proposed scheme. To see this, recall that the Pitman-Koopman-Darmois (PKD) theorem states that for exponential families in (4), there exist sufficient statistics such that the number of scalar components does not increase as the sample size increases. However, in approximating (2) with MC, we need to compute not only terms containing the sufficient statistics T(w) but also $\frac{1}{M} \sum_{i=1}^{M} \log h(w_i)$. Regardless, even though the PKD result cannot be applied directly in our case, it still suggests considering members of the exponential family as candidates for Q_{θ} . We derive technical results for the forms of $W(\theta, \xi)$ and $g(\cdot)$, where the graph size is not affected by MC sampling.

To approximate KL in (2), we need to compute MC estimation (3) for $g(w) = \log q_{\theta}(w)$ (or $\log p(w)$). Assume that the factorization form (4) of distributions $q_{\theta}^d(w)$ (and similarly $p^d(w)$) and recall that the weights of NN are parameterized as $w \sim W(\theta, \xi)$. Then, $\mathbb{E}_{\theta} \log q_{\theta}(w)$ is approximated as:

$$\frac{1}{M} \sum_{i=1} \left\{ \log h(w(\theta, \xi_i)) + \eta(\theta)' T(w(\theta, \xi_i)) \right\} - A(\theta)$$
 (5)

To keep the graph size agnostic of M, we must handle the initial two terms in (5). Checking distributions from Tab. 1, our work reduces to functions of the form w^k and $\log(w)$.

Denote S as the dimension of θ , i.e., number of parameters defining the distribution Q_{θ} . For example, for the Exponential(λ): S=1 and $\theta=(\lambda)$; for Gaussian(μ , σ): S=2 and $\theta=(\mu,\sigma)$. Denote k to be a positive integer.

Theorem 1 If $W(\theta, \xi) = \eta(\theta)T(\xi)$ (S = 1), then there exists a parameterization tuple P with $d_P = 1$ for the following functions g(w): w^k , $\log(w)$, and $\frac{1}{w^k}$.

Corollary 1 If $W(\theta, \xi)' = f(W(\theta, \xi))$ and $W(\theta, \xi) = \eta(\theta)T(\eta)$, then Theorem 1 applies to $W(\theta, \xi)'$ and $g(W(\theta, \xi)')$ if g(w'(w)) is: w^k , $\log(w)$, and $\frac{1}{w^k}$.

Theorem 2 If $W(\theta, \xi) = \sum_{s=1}^{S} \eta_s(\theta) T_s(\xi)$, and $g(w) = w^k$, then there exists a parameterization tuple P with

$$d_P = \binom{k+S-1}{S-1}. (6)$$

Remark 2 As long as $d_P < M$, it is possible to create a computation graph of a smaller size by reparameterization, compared to a direct implementation of the MC approximation. Note that for a small M it is still possible for a parameterization tuple to generate a graph larger than a naïve implementation. For example, consider $\sum_{i=1}^{M} (\mu + \sigma \xi)^2$. When M=1, the naïve construction would have $d_P=2$, $(n=(\mu,\sigma),t=(1,\xi)$, while a "nicer" tuple may have $d_P=3$ independent of M $(n=(\mu^2,2\mu\sigma,\sigma^2),t=(1,\xi,\xi^2))$.

Corollary 2 If $W(\theta, \xi)' = f(W(\theta, \xi))$ and $W(\theta, \xi) = \sum_{s=1}^{S} \eta_s(\theta) T_s(\eta)$, where $S \ge 2$, then Theorem 2 applies to $W(\theta, \xi)'$ and $g(W(\theta, \xi)')$ if $g(w'(w)) = w^k$.

Relevance of results: (1) Thm. 1 can be applied when $W(\theta,\xi)$ represents a distribution with scaling property: any positive real constant times a random variable having this distribution comes from the same distributional family. (2) Thm. 2 can be applied, when $W(\theta,\xi)$ is a member of the location-scale family. (3) Corollaries 1 and 2 are useful when random variables can be presented as a transformation of other distributions, e.g. $\operatorname{LogNormal}(\mu,\sigma^2)$ can be generated as $\exp(\operatorname{N}(\mu,\sigma^2))$. Table 1 summarizes the choice of q_θ and p for Bayesian VI, which lead to the computation graph size d_P being independent of M in MC estimation.

Although Theorem 2 does not suggest that there are no nice parameterization tuples for the case where $g(w) = \log w, 1/w^k$, empirically we did not find tuples that allow for d_P to be independent of M. But it is interesting to consider an approximation which does allow for this independence.

4.1 TAYLOR APPROXIMATED MONTE CARLO

Our results extend to the generic polynomial case where $g(w) = p_K(w)$, an arbitrary polynomial of degree K:

Corollary 3 If $W = \sum_{s=1}^{S} \eta_s(\theta) T_s(\xi)$, and $g(w) = p_K(w)$, then there exists parameterization tuple P, such that for any M iterations

$$d_P = \binom{K+S}{S} - 1. \tag{7}$$

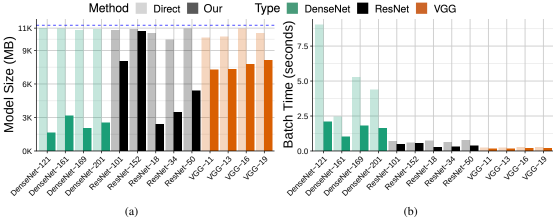


Figure 6: For maximum possible number of MC iterations for a given model via the direct MC method, we show: (a) Model size (dashed blue line indicates GPU capacity, 11GB), (b) Training time. For some networks, our method occupies less than 25% of memory and 5 times faster.

So, can we find a parameterization tuple for any g(w) that we can approximate via a polynomial Taylor expansion?

Theorem 3 Let $W = \sum_{s=1}^{S} \eta_s(\theta) T_s(X)$, $S \ge 2$. If an approximation of g(w) uses K Taylor terms, then Cor. 3 applies.

Practical implications. If one is limited to running a maximum number of MC iterations M_{max} , such an approximation of g(w) allows a tradeoff between accuracy of running just M_{max} iterations for the real g(w) versus approximating g(w) with $K(M_{max})$ terms and running $M\gg M_{max}$ iterations instead, since d_p is independent of M. This strategy may not work for approximating non-polynomial functions, and is a "fall-back" that could be used for arbitrary distributions.

Example 1. Let $W = \mu + \sigma \xi \implies S = 2$ and $g(w) = \log w$, then

$$\sum_{i=1}^{M} g(w_i) = \sum_{i=1}^{M} \log(w_i) \approx \sum_{i=1}^{M} \sum_{k=0}^{K} \frac{1}{k!} (\mu + \sigma \xi_i - 1)^k$$

where we take the Taylor expansion of $\log(w)$ around w=1. This is clearly a polynomial function of order K, and applying Corollary 3, we have $d_P=\frac{1}{2}(K+1)(K+2)-1$ interactions. For example, if one is able to run just 9 direct MC iterations, it is possible to approximate g(w) with K=3 terms, allowing any number of MC iterations M.

4.2 APPLYING REPARAMETERIZATION IN BAYESIAN NN

Recall that training a Bayesian NN via VI requires the approximation of both the KL term and expected value of log-likelihood in (2). While it is clear how MC reparameterization can be applied to approximate the KL term, what

can we say about the likelihood term? In general, this term cannot be handled by the ideas described so far although some practical strategies are possible.

Usually, estimating the expectation of the likelihood term is based on [Kingma and Welling, 2013, Kingma et al., 2015], where for every data item b in the minibatch (of size B), one MC sample is selected, which results in B different samples – in fact, [Kingma and Welling, 2013] suggests that the number of samples per data item can be set to one if the minibatch size is "large enough" which we will discuss more shortly. If a large B is feasible, then our scheme might not contribute substantially in estimating the likelihood term. However, if B is small, then our scheme can provide some empirical benefits, described next.

Let (x,y) be the observed data and (x_b,y_b) be the observed b-th data point. Let w correspond to the weights of NN with L layers. We can use $w(l,\cdot)$ to index the weights of layer l. Note that we can draw a unique sample of w for each data point b which we denote as w(l,b). When M samples are drawn for b, these will be indexed by $w_i(l,b)$ for $i=1,\cdots,M$. Notice that $w_1(l,b)$ is the same as w(l,b). In the forward pass, u_b^l is the output for the b-th data point and u_b^l is the output of the last layer for data point x_b .

Observation 1 (Likelihood form in BNN) Consider the following form for regression and classification tasks, **Regression:** Consider $y \sim N(u^L, \hat{\sigma})$, where $\hat{\sigma}$ is fixed. Then,

$$\log p(y_b \mid w, x_b) = \log \left(\frac{1}{\sqrt{2\pi}} \exp \left(-\frac{1}{2} (y_b - u_b^L)^2 \right) \right)$$
$$= \log \frac{1}{\sqrt{2\pi}} - \frac{1}{2} y_b^2 - y_b u_b^L + \frac{1}{2} (u_b^L)^2.$$

Classification: Consider a binary classification problem.

Then, $y \sim Bern(p)$, where $p = \frac{1}{1 + \exp(-u^L)}$. Thus,

$$\log p(y_b \mid w, x_b) = \log \left(p^{y_b} (1 - p)^{1 - y_b} \right)$$

$$= -\log \left(1 + e^{-u_b^L} \right) - (1 - y_b) u_b^L$$

$$= -\log(2) + \frac{u_b^L}{2} - \frac{\left(u_b^L \right)^2}{4} + O\left(\left(u_b^L \right)^3 \right)$$

$$- (1 - y_b) u_b^L.$$

Based on the above description, let us assume that the final layer output u^L corresponds to a convolution or a fully connected layer with no activation function. Then, the log-likelihood term in a regression and classification setup can be expressed as

$$\log p(y_b|w, x_b) = \text{polynomial}(u_b^{L-1}w(L)).$$

SGVB Estimator. Following [Kingma and Welling, 2013], the $\mathbb{E}_{q_{\theta}}$ [log p(y|w,x)] term for the minibatch (of size B) can be written as

$$S_1 := \frac{1}{B} \sum_{b=1}^{B} \mathbb{E}_{q_{\theta}} \left[\log p(y_b|w, x_b) \right].$$

To approximate the expectation, we use 1 sample $w(\cdot,b)$ for each data point (x_b,y_b) , which results in $S_1 = \frac{1}{B} \sum_{b=1}^{B} \log p(y_b|w(\cdot,b),x_b)$. Substituting in polynomial $(u_b^{L-1}w(L,b))$ into $\log p(y_b|w(\cdot,b),x_b)$ leads to the following form for variance $V(S_1)$,

$$\frac{1}{B} \left(V\left(w(L,b) \right) \mathbb{E} \left[\left(u_b^{L-1} \right)^2 \right] + V(u_b^{L-1}) \mathbb{E}^2 \left[w(L,b) \right] \right), \tag{8}$$

plus higher order terms which decreases as B grows. By efficiently evaluating the KL term, we can utilize the memory savings to increase the batch size B and thus, to decrease the variance of S_1 .

MC Reparameterization estimator of likelihood. The above strategy is practically sufficient. However, if B is limited by hardware, we can use the memory savings for more MC samples (higher M) for improving the estimate of the log likelihood term. This reduces the variance of first term in (8) by a factor of M, but the scheme described is restricted to the last layer.

5 EXPERIMENTS: BAYESIAN DENSENET, U-NET, AND OTHER NETWORKS

We perform experiments on Bayesian forms of several architectures and show that training is feasible. While we expect some drop in overall accuracy compared to a deterministic version of the network, these experiments shed light on the benefits/limitations of increasing MC iterations. Since

model uncertainty is important in scientific applications, we also study the feasibility of training such models for classifying high-resolution brain images from a public dataset.

Setup. For deterministic comparisons, we run several variations of PreActResNet [He et al., 2016] and Densenet [Huang et al., 2017] (9 in total) on CIFAR10. For brain images, we use a simple modification of 3D U-Net [Ronneberger et al., 2015]. Since our method is most relevant when a closed form for KL is unavailable, we select the approximate posterior to be a Radial distribution, where samples can be generated as: $\mu + \sigma * \frac{\xi}{||\xi||} * |r|$, where $\xi \sim MVN(0, I), r \sim N(0, 1)$ and the prior of our weights is a Normal distribution. This satisfies the conditions of Thm. 2, allowing us to find a parameterization tuple that does not grow with respect to M: we can run 1000 + MCiterations with almost no additional GPU memory cost compared to 1 MC iteration. Another reason for choosing the Radial distribution as our approximate posterior q_{θ} is because Gaussian-approximate posteriors do not perform well in high-dimensional settings [Farquhar et al., 2020]. Empirically, we find this to be the case as well; we were not able to train any models with a standard Gaussian assumption without any ad-hoc fixes such as pretraining, burn-in, or *KL*-reweighting (common in many implementations).

Parameter settings/hardware. All experiments used Nvidia 2080 TIs. The code was implemented in PyTorch, using the Adam optimizer [Kingma and Ba, 2014] for all models, with training data augmented via standard transformations: normalization, random re-cropping, and random flipping. All models were run for 100 epochs.

5.1 TIME AND SPACE CONSIDERATIONS

We first examine whether our MC-reparameterization leads to meaningful benefits in model size or runtime. We should expect a competitive advantage in model size as the number of MC iterations grows, which may come at the cost of significantly increased runtime. To allow ease of comparison, we fix the batch size for all models to be 32. We determine the maximum number of MC iterations able to run on a single GPU for a given model via the classical direct method. For DenseNet-121, we are able to run 89 MC iterations, while for VGG-16 we are only able to run 5 MC iterations.

Figure 6 shows a comparison of computational performance between our method and the direct approach. (a) With our construction, we significantly **reduce model size** on the GPU (Fig. 6a). For smaller models like DenseNet, for the same number of MC iterations our method uses less than 25% of GPU memory, which allows for a significant **increase in batch size**. Since the size of the computation graph in our construction is independent of M, for the memory used in Fig. 6a we are able to run for M=1000 or more. (b) The significant reduction of model size on the

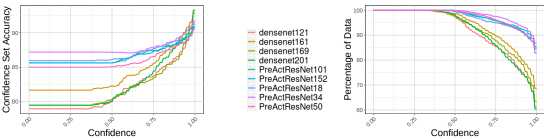


Figure 7: Confidence Set Accuracy and Confidence Sets on CIFAR-10 for a variety of ResNet and DenseNet models with 100 MC iterations (not previously possible). Both ResNet and Densenet achieve accuracy of more than 90% with 100% confidence, but ResNet is 100% confident on almost 90% of the data.

GPU results in a reduction of training time per batch, up to $5 \times$ (Figure 6b); the generated computation graph has fewer parameters (nodes on the path) during backpropagation.

5.2 PREDICTION CONFIDENCE/ACCURACY AND HOW MANY MC ITERATIONS?

For our next set of experiments, we run a Bayesian version of PreActResNet and DenseNet with 100 MC iterations, which is feasible.

- (a) We evaluate the accuracy concurrently with the confidence of the prediction, offered directly by the model. We expect that the model has a higher accuracy for those samples where it highly confident. This is indeed the case Figure 7 shows the accuracy for varying levels of confidence over the entire validation set for a number of models. At high confidence levels, all models perform well, competing strongly with state of the art results. Additionally, we observe the proportion of data for which the model is confident is large (Figure 7 right). We can see that Bayesian model is at least 75% confident on 85%–95% of data.
- (b) One issue in Bayesian networks is evaluating the expected drop in accuracy (compared to its deterministic versions), a behavior common in both shallow and deep models [Wenzel et al., 2020]. Figure 7 (left) reassures us that the drop in performance for a number of widely used architectures is not that significant even when the model is not confident.
- (c) To understand the effect of increasing the number of MC iterations, we run replications of experiment on ResNet-50 for 3 different number of MC iterations, Figure 8(left): 1 iteration (black), 17 iterations maximum possible on GPU with the traditional method (blue), and 100 iterations (red) possible to run due to our method. In all cases, as the threshold increases, model confidence increases and as expected, the accuracy does as well. However, we see that training with 100 MC iterations, consistently provides higher accuracy for the entire range of confidences. In contrast, with 1 MC iteration, accuracy has higher variance for the non-confident set.

5.3 NEUROIMAGING: PREDICTIVE UNCERTAINTY IN BRAIN IMAGING ANALYSIS

While we demonstrated advantages of our reparameterization in traditional image classification settings and benchmarks – mostly as a proof of feasibility – a real need for BNNs is in scientific/biomedical domains: where high confidence and accurate predictions may inform diagnosis/treatment. To evaluate applicability, we focus on a learning task with brain imaging data.

Data. Data used in our experiments were obtained from the Alzheimer's Disease Neuroimaging Initiative (ADNI). As such, the investigators within the ADNI contributed to the design and implementation of ADNI and/or provided data but did not participate in analysis or writing of this report. A complete listing of ADNI investigators can be found in [ADNI, 2020a]. The primary goal of ADNI has been to test whether serial magnetic resonance imaging (MRI), positron emission tomography (PET), other biological markers, and clinical and neuropsychological assessment can be combined to measure the progression of mild cognitive impairment (MCI) and early Alzheimer's disease (AD).

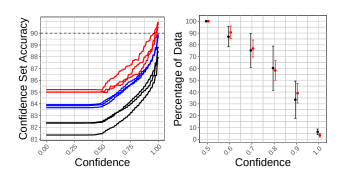


Figure 8: (left) Replicated Confidence Set Accuracy on CIFAR-10 for Resnet-50 with different number of MC iterations: 1 (black), 17 (blue, maximum allowed on GPU with standard method) and 100 (red). With M=100 the accuracy is higher for any confidence. (right) Distributions of Confidence Set Size for a number of replications, with 1 MC iteration (black) and 100 MC iterations (red). With 100 MC iterations variance is smaller.

	Confidence						
	0.5	0.6	0.7	0.8	0.9	1	
m=1	63.07 ± 1.47	62.14 ± 1.59	63.01 ± 0.64	64.13 ± 4.15	59.59 ± 4.40	60.71 ± 15.15	
m = 100	64.39 ± 4.59	66.23 ± 4.19	66.05 ± 1.00	67.77 ± 4.00	66.82 ± 1.24	87.50 ± 17.68	
Δ	1.33	4.09	3.04	3.64	7.23	26.79	

Table 2: Average validation accuracy per model confidence for 2 values of MC iterations. $\Delta = A_{100} - A_1$, where A_i is validation accuracy for i MC iterations. With 100 MC iterations we got on average much better results, especially when prediction is highly confident.

For up-to-date information, see [ADNI, 2020b]. Classifying healthy and diseased individuals via their MR images, similar to ADNI, is common in the literature However, overfitting when using deep models remains an issue for two reasons: small dataset size and a large feature space. Here, we look at a specific setting where we have 388 individuals with pre-processed MR images of size $105 \times 127 \times 105$. Preprocessing. All MR images were registered to MNI space using SPM12 with default settings.

Network. We use a slightly modified version of the encoder from an off-the-shelf 3D U-Net architecture [Ronneberger et al., 2015], demonstrated in Figure 9, to learn a classifier for cognitively normal (CN) and Alzheimer's Disease (AD) subjects. We note that while this architecture is not competitive with those which achieve state-of-the-art classification accuracy on ADNI, our aim here is to demonstrate feasibility of training deep Bayesian models in this setting and evaluate the value of accurate confidence estimation.

We train the model on 300 individuals, and validate on the remaining 88. Additional experimental details can be found in the appendix. Since the input to the network is a mini-batch of high dimensional images, when we take into account the memory already needed by a deterministic model, we already reach the limits of the GPU memory. While we cannot perform more than 1 MC iteration with the standard method, we can successfully perform more than 100 with our scheme. We evaluate the consistency of performance with several runs of training when we are allowed to use 1 versus 100 MC iterations. (a) Table 2 shows the average validation accuracy for the choice of MC iterations and their

```
Conv3D_Block(1, 16)

MaxPool3d((3,3,3))

Conv3D_Block(16, 32, stride=1)

MaxPool3d((2,2,2))

Conv3D_Block(32, 64, stride=1)

MaxPool3d((2,2,2))

Conv3D_Block(64, 128, stride=3),

MaxPool3d((2,2,2))

Conv3D_Block(128, 256, stride=3)

Linear(256, 2)
```

Figure 9: Structure of the model we used for ADNI classification.

difference. We see that for every confidence threshold, training with 100 MC iterations provides higher accuracy on average. This is especially noticeable on a high confident set, where the difference approaches 26.7%. (b) In addition to accuracy, it is important to understand how consistent the estimation is. Figure 8 (right) demonstrates the distribution of the size of confident set. While on average, the size of the "confident set" of the two models is similar, the variance is significantly smaller when we use a larger number of MC iterations, consistent with our hypothesis in §1. In cases where this confidence needs to be measured as accurately as possible, one obtains benefits over a single MC iteration.

6 CONCLUSIONS

While a broad variety of neural network architectures are used in vision and medical imaging, successfully training them in a Bayesian setting poses challenges. Part of the reason has to do with distributional assumptions. Moving to a broader class of distributions involves MC estimations but direct implementations pose serious demands on memory and run-time. In this work, we identify that different computation graphs can be constructed for different parameterizations of the target function. Specifically when one is attempting a Monte Carlo approximation, these graphs can grow linearly with the number of MC iterations needed, which is undesirable. By directly characterizing the parameterizations that lead to different graphs, we analyze situations where it is possible for graphs to be constructed independent of this sampling rate (number of MC iterations). Evaluating our parameterization empirically, we find that it is feasible to run a large number of MC iterations for large networks in vision, with a nominal drop in accuracy (compared to deterministic versions). The code is available at https: //github.com/vsingh-group/mcrepar.

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