Discretely Relaxing Continuous Variables for tractable Variational Inference

Trefor W. Evans
University of Toronto
trefor.evans@mail.utoronto.ca

Prasanth B. Nair
University of Toronto
pbn@utias.utoronto.ca

Abstract

We explore a new research direction in Bayesian variational inference with discrete latent variable priors where we exploit Kronecker matrix algebra for efficient and exact computations of the evidence lower bound (ELBO). The proposed "DIRECT" approach has several advantages over its predecessors; (i) it can exactly compute ELBO gradients (i.e. unbiased, zero-variance gradient estimates), eliminating the need for high-variance stochastic gradient estimators and enabling the use of quasi-Newton optimization methods; (ii) its training complexity is independent of the number of training points, permitting inference on large datasets; and (iii) its posterior samples consist of sparse and low-precision quantized integers which permit fast inference on hardware limited devices. In addition, our DIRECT models can exactly compute statistical moments of the parameterized predictive posterior without relying on Monte Carlo sampling. The DIRECT approach is not practical for all likelihoods, however, we identify a popular model structure which is practical, and demonstrate accurate inference using latent variables discretized as extremely low-precision 4-bit quantized integers. While the ELBO computations considered in the numerical studies require over $10^{2352}$ log-likelihood evaluations, we train on datasets with over two-million points in just seconds.

1 Introduction

Hardware restrictions posed by mobile devices make Bayesian inference particularly ill-suited for on-board machine learning. This is unfortunate since the safety afforded by Bayesian statistics is extremely valuable in many prominent mobile applications. For example, the cost of erroneous decisions are very high in autonomous driving or mobile robotic control. The robustness and uncertainty quantification provided by Bayesian inference is therefore extremely valuable for these applications provided inference can be performed on-board in real-time [1, 2].

Outside of mobile applications, resource efficiency is still an important concern. For example, deployed models making billions of predictions per day can incur substantial energy costs, making energy efficiency an important consideration in modern machine learning architectures [3].

We approach the problem of efficient Bayesian inference by considering discrete latent variable models such that posterior samples of the variables will be quantized and sparse, leading to efficient inference computations with respect to energy, memory and computational requirements. Training a model with a discrete prior is typically very slow and expensive, requiring the use of high variance Monte Carlo gradient estimators to learn the variational distribution. The main contribution of this work is the development of a method to rapidly learn the variational distribution for such a model without the use of any stochastic estimators; the objective function will be computed exactly at each iteration. To our knowledge, such an approach has not been taken for variational inference of large-scale probabilistic models.
In this paper, we compare our work not only to competing stochastic variational inference (SVI) methods for discrete latent variables, but also to the more general SVI methods for continuous latent variables. We make this comparison with continuous variables by discretely relaxing continuous priors using a discrete prior with a finite support set that contains much of the structure and information as its continuous analogue. Using this discretized prior we show that we can make use of Kronecker matrix algebra for efficient and exact ELBO computations. We will call our technique DIRECT (DIscrete RElaxation of ConTinuous variables). We summarize our main contributions below:

- We efficiently and exactly compute the ELBO using a discrete prior even when this computation requires more likelihood evaluations than the number of atoms in the known universe. This achieves unbiased, zero-variance gradients which we show outperforms competing Monte Carlo sampling alternatives that give high-variance gradient estimates while learning.
- Complexity of our ELBO computations are independent of the quantity of training data using the DIRECT method, making the proposed approach amenable to big data applications.
- At inference time, we can exactly compute the statistical moments of the parameterized predictive posterior distribution, unlike competing techniques which rely on Monte Carlo sampling.
- Using a discrete prior, our models admit sparse posterior samples that can be represented as quantized integer values to enable efficient inference, particularly on hardware limited devices.
- We present the DIRECT approach for generalized linear models and deep Bayesian neural networks for regression, and discuss approximations that allow extensions to many other models.
- Our empirical studies demonstrate superior performance relative to competing SVI methods on problems with as many as 2 million training points.

The paper will proceed as follows; section 2 contains a background on variational inference and poses the learning problem to be addressed while section 3 outlines the central ideas of the DIRECT method, demonstrating the approach on several popular probabilistic models. Section 4 discusses limitations of the proposed approach and outlines some work-arounds, for instance, we discuss how to go beyond mean-field variational inference. We empirically demonstrate our approaches in section 5, and conclude in section 6. Our full code is available at https://github.com/treforevans/direct.

## 2 Variational Inference Background

We begin with a review of variational inference, a method for approximating probability densities in Bayesian statistics [4, 5, 6, 7, 8, 9]. We introduce a regression problem for motivation; given $X \in \mathbb{R}^{n \times d}$, $y \in \mathbb{R}^n$, a $d$-dimensional dataset of size $n$, we wish to evaluate $y_a$ at an untried point $x_a$ by constructing a statistical model that depends on the $b$ latent variables in the vector $w \in \mathbb{R}^b$. After specifying a prior over the latent variables, $Pr(w)$, and selecting a probabilistic model structure that admits the likelihood $Pr(y|w)$, we may proceed with Bayesian inference to determine the posterior $Pr(w|y)$ which generally requires analytically intractable computations.

Variational inference turns the task of computing a posterior into an optimization problem. By introducing a family of probability distributions $q_\theta(w)$ parameterized by $\theta$, we minimize the Kullback-Leibler divergence to the exact posterior [9]. This equates to maximization of the evidence lower bound (ELBO) which we can write as follows for a continuous or discrete prior, respectively.

\[
\text{ELBO}(\theta) = \int q_\theta(w) \left( \log Pr(y|w) + \log Pr(w) - \log q_\theta(w) \right) dw, \quad (1)
\]

\[
\text{ELBO}(\theta) = q^T \left( \log \ell + \log p - \log q \right), \quad (2)
\]

where $\log \ell = \{\log Pr(y|w_i)\}_{i=1}^m$, $\log p = \{\log Pr(w_i)\}_{i=1}^m$, $q = \{q_\theta(w_i)\}_{i=1}^m$, and $\{w_i\}_{i=1}^m = W \in \mathbb{R}^{b \times m}$ is the entire support set of the discrete prior.

It is immediately evident that computing the ELBO is challenging when $b$ is large, since in the continuous case eq. (1) is a $b$-dimensional integral, and in the discrete case the size of the sum in eq. (2) generally increases exponentially with respect to $b$. Typically, the ELBO is not explicitly computed and instead, a Monte Carlo estimate of the gradient of the ELBO with respect to the variational
parameters $\theta$ is found, allowing stochastic gradient descent to be performed. We will outline some existing techniques to estimate ELBO gradients with respect to the variational parameters, $\theta$.

For continuous priors, the reparameterization trick [10] can be used to perform variational inference. The technique uses Monte Carlo estimates of the gradient of the evidence lower bound (ELBO) which is maximized during the training procedure. While this approach has been employed successfully for many large-scale models, we find that discretely relaxing continuous latent variable priors can improve training and inference performance when using our proposed DIRECT technique which computes the ELBO (and its gradients) exactly.

When the latent variable priors are discrete, reparameterization cannot be applied, however, the REINFORCE [11] estimator may be used to provide an unbiased estimate of the ELBO during training (alternatively called the score function estimator [12], or likelihood ratio estimator [13]). Empirically, the REINFORCE gradient estimator is found to give a high-variance when compared with reparameterization, leading to a slow learning process. Unsurprisingly, we find that our proposed DIRECT technique trains significantly faster than a model trained using a REINFORCE estimator.

Recent work in variational inference with discrete latent variables has largely focused on continuous relaxations of discrete variables such that reparameterization can be applied to reduce gradient variance compared to REINFORCE. One example is CONCRETE [14, 15] and its extensions [16, 17]. We consider an opposing direction by identifying how the ELBO (eq. (2)) can be computed exactly for a class of discretely relaxed probabilistic models such that the discrete latent variable model can be trained more easily then its continuous counterpart. We outline this approach in the following section.

### 3 DIRECT: Efficient ELBO Computations with Kronecker Matrix Algebra

We outline the central ideas of the DIRECT method and illustrate its application on several probabilistic models. The DIRECT method allows us to efficiently and exactly compute the ELBO which has several advantages over existing SVI techniques for discrete latent variable models such as, zero-variance gradient estimates, the ability to use a super-linearly convergent quasi-Newton optimizer (since our objective is deterministic), and the per-iteration complexity is independent of training set size. We will also discuss advantages at inference time such as the ability to exactly compute predictive posterior statistical moments, and to exploit sparse and low-precision posterior samples.

To begin, we consider a discrete prior over our latent variables whose support set $W$ forms a Cartesian tensor product grid as most discrete priors do (e.g. any prior that factorizes between variables) so that we can write

$$W = \begin{pmatrix} \tilde{w}_1^T \otimes 1_m \otimes \cdots \otimes 1_m \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ 1_m \otimes 1_m \otimes \cdots \otimes \tilde{w}_b^T \end{pmatrix},$$

where $1_m \in \mathbb{R}^m$ denotes a vector of ones, $\tilde{w}_i \in \mathbb{R}^m$ contains the $m$ discrete values that the $i$th latent variable $w_i$ can take, $m = \tilde{m}^b$, and $\otimes$ denotes the Kronecker product [18]. Since the number of columns of $W \in \mathbb{R}^{b \times \tilde{m}^b}$ increases exponentially with respect to $b$, it is evident that computing the ELBO in eq. (2) is typically intractable when $b$ is large. For instance, forming and storing the matrices involved naively require exponential time and memory. We can alleviate this concern if $\mathbf{q}$, $\log p$, $\log \ell$, and $\log q$ can be written as a sum of Kronecker product vectors (i.e. $\sum_i \otimes_{j=1}^b f^{(i)}_j$, where $f^{(i)}_j \in \mathbb{R}^m$). If we find this structure, then we never need to explicitly compute or store a vector of length $m$. This is because eq. (2) would simply require multiple inner products between Kronecker product vectors which the following result demonstrates can be computed extremely efficiently.

**Proposition 1.** The inner product between two Kronecker product vectors $k = \otimes_{i=1}^b k^{(i)}$, and $a = \otimes_{i=1}^b a^{(i)}$ can be computed as follows [18],

$$a^T k = \prod_{i=1}^b a^{(i)} T k^{(i)},$$

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*The discrete values that the $i$th latent variable can take, $\tilde{w}_i$, may be chosen a priori or learned during ELBO maximization (may be helpful for coarse discretizations). For the sake of simplicity, we focus on the former.*
where \( a^{(i)} \in \mathbb{R}^m, a \in \mathbb{R}^m, k^{(i)} \in \mathbb{R}^m \), and \( k \in \mathbb{R}^m \).

This result enables substantial savings in the computation of the ELBO since each inner product computation is reduced from the naive exponential \( \mathcal{O}(m^b) \) cost to a linear \( \mathcal{O}(bn) \) cost.

We now discuss how the Kronecker product structure of the variables in eq. (2) can be achieved. Firstly, if the prior is chosen to factorize between latent variables, as it often is, (i.e. \( \Pr(w) = \prod_{i=1}^b \Pr(w_i) \)) then \( p = \bigotimes_{i=1}^b p_i \) admits a Kronecker product structure where \( p_{bi} = \{ \Pr(w_i = \bar{w}_{ji}) \}_{j=1}^m \in (0, 1)^m \). The following result demonstrates how this structure for \( p \) enables \( \log p \) to be written as a sum of \( b \) Kronecker product vectors.

**Proposition 2.** The element-wise logarithm of the Kronecker product vector \( k = \bigotimes_{i=1}^b k^{(i)} \) can be written as a sum of \( b \) Kronecker product vectors as follows,

\[
\log k = \bigoplus_{i=1}^b \log k^{(i)}, \tag{5}
\]

where \( k^{(i)} \in \mathbb{R}^m, k \in \mathbb{R}^m \) contain positive values, and \( \bigoplus \) is a generalization of the Kronecker sum \([19]\) for vectors which we define as follows

\[
\bigoplus_{i=1}^b \log k^{(i)} = \sum_{i=1}^b \left( \bigotimes_{j=1}^{i-1} 1_m \right) \log k^{(i)} \otimes \left( \bigotimes_{j=i+1}^b 1_m \right). \tag{6}
\]

The proof is trivial. We will first consider a mean-field variational distribution that factorizes over latent variables such that both \( q_i = \bigotimes_{j=1}^b q_{ij} \) and \( \log q_i = \bigotimes_{j=1}^b \log q_{ij} \) can be written as a sum of Kronecker product vectors, where \( q_{ij} = \{ \Pr(w_{ji} = \bar{w}_{ji}) \}_{j=1}^m \in (0, 1)^m \) are used as the variational parameters, \( \theta \), with the use of the softmax function. For the mean-field case we can rewrite eq. (2) as

\[
\text{ELBO}(\theta) = q^T \log \ell + \sum_{i=1}^b q_{i}^T \log p_i - \sum_{i=1}^b q_{i}^T \log q_i, \tag{7}
\]

where we use the fact that \( q_i \) defines a valid probability distribution for the \( i \)th latent variable such that \( q_{i}^T 1_m = 1 \). We extend results to unfactorized prior and variational distributions later in section 4.

The structure of \( \log \ell \) depends on the probabilistic model used; in the worst case, \( \log \ell \) can always be represented as a sum of \( m \) Kronecker product vectors. However, many models admit a far more compact structure where dramatic savings can be realized as we demonstrate in the following sections.

### 3.1 Generalized Linear Regression

We first focus on the popular class of Bayesian generalized linear models (GLMs) for regression. While the Bayesian integrals that arise in GLMs can be easily computed in the case of conjugate priors, for general priors inference is challenging.

This highly general model architecture has been applied in a vast array of application areas. Recently, Wilson et al. [20] used a scalable Bayesian generalized linear model with Gaussian priors on the output layer of deep neural network with notable empirical success. They also considered the ability to train the neural network simultaneously with the approximate Gaussian process which we also have the ability to do if a practitioner were to require such an architecture.

Consider the generalized linear regression model \( y = \Phi w + \epsilon \), where \( \epsilon \sim \mathcal{N}(0, \sigma^2 \mathbf{I}) \), and \( \Phi = \{ \phi_j(x_i) \}_{i,j} \in \mathbb{R}^{n \times b} \) contains the evaluations of the basis functions on the training data. The following result demonstrates how the ELBO can be exactly and efficiently computed, assuming the factorized prior and variational distributions over \( w \) discussed earlier. Note that we also consider a prior over \( \sigma^2 \).

**Theorem 1.** The ELBO can be exactly computed for a discretely relaxed regression GLM as follows

\[
\text{ELBO}(\theta) = -\frac{n}{2} q_{\sigma}^T \log \sigma^2 - \frac{1}{2} q_{\sigma}^T (\sigma^{-2}) \left( y^T y - 2 s^T (\Phi^T \Phi) s + \text{diag}(\Phi^T \Phi)^T s^2 + \sum_{j=1}^b q_{\phi_{ji}}^T \Phi_{ji} \right) + \sum_{i=1}^b (q_i^T \log p_i - q_i^T \log q_i) + q_{\sigma}^T \log p_{\sigma} - q_{\sigma}^T \log q_{\sigma}, \tag{8}
\]
where $q_\ell, P_\ell \in \mathbb{R}^m$ are factorized variational and prior distributions over the Gaussian noise variance $\sigma^2$ for which we consider the discrete positive values $\sigma^2 \in \mathbb{R}^m$, respectively. Also, we use the shorthand notation $H = \{w_\ell^T \sum_{j=1}^m \phi_j^T y \}^b_{j=1} \in \mathbb{R}^{m \times b}$, and $s = \{q_\ell^T w \}^b_{j=1} \in \mathbb{R}^b$.

A proof is provided in appendix A of the supplementary material. We can pre-compute the terms $y^T y, \Phi^T \Phi, H, \text{and } \Phi \Phi$ before training begins (since these do not depend on the variational parameters) such that the final complexity of the proposed DIRECT method outlined in Theorem 1 is only $\mathcal{O}(bn_1 + b^2)$. This complexity is independent of the number of training points, making the proposed technique ideal for massive datasets. Also, each of the pre-computed terms can easily be updated as more data is observed making the techniques amenable to online learning applications.

**Predictive Posterior Computations** Typically, the predictive posterior distribution is found by sampling the variational distribution at a large number of points and running the model forward for each sample. To exactly compute the statistical moments, a model would have to be run forward at every point in the hypothesis space with is typically intractable, however, we can exploit Kronecker matrix algebra to efficiently compute these moments exactly. For example, the exact predictive posterior mean for our generalized linear regression model is computed as follows

$$
\mathbb{E}(y_*) = \sum_{i=1}^m q(w_i) \int y_* \Pr(y_* \mid w_i) dy_* = \Phi_* W q = \Phi_* s,
$$

where $s = \{q^T w \}^b_{j=1} \in \mathbb{R}^b$, and $\Phi_* j \in \mathbb{R}^{1 \times b}$ contains the basis functions evaluated at $x_*$. This computation is highly efficient, requiring just $\mathcal{O}(b)$ time per test point. It can be shown that a similar scheme can be derived to exactly compute higher order statistical moments, such as the predictive posterior variance, for generalized linear regression models and other DIRECT models.

We have shown how to exactly compute statistical moments, and now we show how to exploit our discrete prior to compute predictive posterior samples extremely efficiently. This sampling approach may be preferable to the exact computation of statistical moments on hardware limited devices where we need to perform inference with extreme memory, energy and computational efficiency. The latent variable posterior samples $\tilde{W} \in \mathbb{R}^{b \times \text{num}. \text{samples}}$ will of course be represented as a low-precision quantized integer array because of the discrete support of the prior which enables extremely compact storage in memory. Much work has been done elsewhere in the machine learning community to quantize variables for storage compression purposes since memory is a very restrictive constraint on mobile devices [21, 22, 23, 24]. However, we can go beyond this to additionally reduce computational and energy demands for the evaluation of $\Phi_* \tilde{W}$. One approach is to constrain the elements of $\tilde{w}$ to be 0 or a power of 2 so that multiplication operations simply become efficient bit-shift operations [25, 26, 27]. An even more efficient approach is to employ basis functions with discrete outputs so that $\Phi_*$ can also be represented as a low-precision quantized integer array. For example, a rounding operation could be applied to continuous basis functions. Provided that the quantization schemes are an affine mapping of integers to real numbers (i.e. the quantized values are evenly spaced), then inference can be conducted using extremely efficient integer arithmetic [28]. Either of these approaches enable extremely efficient on-device inference.

### 3.2 Deep Neural Networks for Regression

We consider the hierarchical model structure of a Bayesian deep neural network for regression. Considering a DIRECT approach for this architecture is not conceptually challenging so long as an appropriate neuron activation function is selected. We would like a non-linear activation that maintains a compact representation of the log-likelihood evaluated at every point in the hypothesis space, i.e. we would like $\log \ell$ to be represented as a sum of as few Kronecker product vectors as possible. Using a power function for the activation can maintain a compact representation; the natural choice being a quadratic activation function (i.e. output $x^2$ for input $x$).

It can be shown that the ELBO can be exactly computed in $\mathcal{O}(\ell b n_1 / \ell^4)$ for a deep Bayesian neural network with $\ell$ layers, where we assume a quadratic activation function and an equal distribution of discrete latent variables between network layers. This complexity evidently enables scalable Bayesian inference for models of moderate depth, and like we found for the regression GLM model of section 3.1, computational complexity is independent of the quantity of training data, making this approach ideal for large datasets. We outline this model and the computation of its ELBO in appendix D.
4 Limitations & Extensions

In generality, when the support of the prior is on a Cartesian grid, any prior, likelihood, or variational distribution can be expressed using the proposed Kronecker matrix representation, however, this representation will not always be compact enough to be practical. We can see this by viewing these probability distributions over the hypothesis space as high-dimensional tensors. In section 3, we exploited some popular models whose prior, likelihood and variational probability tensors and log-probability tensors all admit a low-rank structure, however, other models may not, in which case their representation will not be so compact. In the interest of generalizing the technique, we outline a likelihood, a prior, and a variational distribution that does not admit a compact representation of the ELBO and discuss several ways the DIRECT method can still be used to efficiently compute, or lower bound the ELBO. We hope that these extensions inspire future research directions in approximate Bayesian inference.

Generalized Linear Logistic Regression Logistic regression models do not easily admit a compact representation for exact ELBO computations, however, we will demonstrate that we can efficiently compute a lower-bound of the ELBO by leveraging developed algebraic techniques. To demonstrate, we will consider a generalized linear logistic regression model which is commonly employed for classification problems. Such a model could easily be extended to a deep architecture following Bradshaw et al. [2], if desired. All terms in the ELBO in eq. (7) can be computed exactly, and as a result, stochastic optimization techniques should be considered. Using this lower bound, the log-likelihood is accurately approximated for hypotheses that correctly classify the training data, however, this representation will not always be compact enough to be practical. In the interest of generalizing the technique, we outline a likelihood, a prior, and a variational distribution that does not admit a compact representation for exact ELBO computations, however, we will demonstrate that we can efficiently compute, or lower bound the ELBO. We hope that these extensions inspire future research directions in approximate Bayesian inference.

Theorem 2. For a generalized linear logistic regression model with classification training labels \( y \in \{0, 1\}^n \), the class-conditional probability \( \Pr(y_i=0|w) = (1 + \exp(-\Phi[i,:]w))^{-1} \), and with the assumption that training examples are sampled independently, the following inequality holds

\[
q^T \log \ell \geq -s^T (\Phi^T y) - \sum_{i=1}^{n} \left\{ \prod_{j=1}^{b} q_{ij}^T \exp(-\phi_{ij} \bar{w}_j) \right\} \tag{10}
\]

We prove this result in appendix B of the supplement. This computation can be performed in \( O(nmbn) \) time, where dependence on \( n \) is evident unlike in the case of the exact computations described in section 3. As a result, stochastic optimization techniques should be considered. Using this lower bound, the log-likelihood is accurately approximated for hypotheses that correctly classify the training data, however, hypotheses that confidently misclassify training labels may be over-penalized. In appendix B we further discuss the accuracy of this approximation and discuss a stable implementation.

Unfactorized Variational Distributions We now consider going beyond a mean-field distribution to account for correlations between latent variables. Considering a finite mixture of factorized categorical distributions as is used in latent structure analysis [29, 30], we can write \( q = \sum_{i=1}^{r} \alpha_i \otimes_{j=1}^{m} q_j^{(i)} \), where \( \alpha \in (0, 1)^r \) is a vector of mixture probabilities for \( r \) components, and \( q_j^{(i)} = \{ \Pr(w_j = \bar{w}_j[k]|i) \}^{m} \in (0, 1)^m \).

While \( q \) can evidently be expressed as a compact sum of Kronecker product vectors, \( \log q \) is more challenging to compute than in the mean-field case, however, the following result demonstrates how we can lower-bound the term involving \( \log q \) in the ELBO (eq. (7)).

Theorem 3. The following inequality holds when we consider a finite mixture of factorized categorical distributions for \( q_0(w) \),

\[
-q^T \log q \geq \max_{\{a_i\}_{i=1}^{r} \in (0, 1)^m \} \left( 1 - \sum_{j=1}^{r} \alpha_j \left( \sum_{i=1}^{b} q_{ij}^{(i)} T \log a_i + \alpha_j \prod_{i=1}^{b} q_{ij}^{(i)} T \frac{q_{ij}^{(i)}}{a_i} + 2 \sum_{k=j+1}^{r} \alpha_k \prod_{i=1}^{b} q_{ij}^{(i)} T \frac{q_{ij}^{(i)}}{a_i} \right) \right),
\]

where \( a = \otimes_{i=1}^{b} a_i \), \( a_i \in (0, 1)^m \) is the center of the Taylor series approximation of \( \log q \).

We prove this result in appendix C and discuss a stable implementation. Note that if the mixture variational distribution \( q \) degenerates to a mean-field distribution equal to \( a \), then the ELBO will be computed exactly, and as \( q \) moves away from \( a \), the ELBO will be underestimated.
Unfactorized Prior Distributions  To consider an unfactorized prior, we assume a prior mixture distribution given by $p = \sum_{i=1}^{r} \alpha_i \otimes_{j=1}^{b} p_j^{(i)}$. When we use this mixture distribution for the prior, $p$ can evidently be expressed as a compact sum of Kronecker product vectors but $\log p$ cannot. The following result demonstrates how we can still lower-bound the term involving $\log p$ in the ELBO (eq. (2)). For simplicity, we assume that the variational distribution factorizes, however, the result could easily be extended to the case of a mixture variational distribution.

Theorem 4. The following inequality holds when we consider a finite mixture of factorized categorical distributions for $p_\theta(\mathbf{w})$.

$$q^T \log p \geq \sum_{i=1}^{r} \sum_{j=1}^{b} q_j^T \log p_j^{(i)}$$

The proof is trivial by Jensen’s inequality. Note that the equality only holds when the prior mixture degenerates to a factorized distribution with all mixture components equivalent.

Unbiased Stochastic Entropy and Prior Expectation Gradients  We previously showed how to lower bound the ELBO terms $q^T \log p$ and $-q^T \log q$ when the variational and/or prior distributions do not factor, however, optimizing this bound introduces bias and does not guarantee convergence to a local optimum of the true ELBO. Here we reintroduce REINFORCE to deliver unbiased gradient estimates for these terms. The REINFORCE estimator typically has high variance, however, since gradient estimates for these terms are so cheap, a massive number of samples can be used per stochastic gradient descent (SGD) iteration to decrease variance. Since we can still compute the expensive $q^T \log \ell$ term exactly when $q$ is an unfactorized mixture distribution, its gradient can be computed exactly. The unbiased gradient estimator of $q^T \log q$ is expressed as follows

$$\frac{\partial}{\partial \theta} q^T \log q = \frac{1}{2} q^T \left( \frac{\partial}{\partial \theta} \left( \log q + 1 \right)^2 \right) \approx \frac{\partial}{\partial \theta} \frac{1}{2t} \sum_{i=1}^{t} \left( \log q(s_i) + 1 \right)^2; \quad (11)$$

where $s_i \in \mathbb{R}^b$ is the $i$th of $t$ samples from the variational distribution used in the Monte Carlo gradient estimator. It is evident that this surrogate loss can be easily optimized using automatic differentiation, and the per-sample computations are extremely cheap.

5 Numerical Studies

5.1 Comparison with REINFORCE

As discussed in section 2, we cannot reparameterize because of the discrete latent variable priors considered, however, we can directly compare the optimization performance of the proposed techniques with the REINFORCE gradient estimator [11]. In fig. 1, we compare ELBO maximization performance between the proposed DIRECT, and the REINFORCE methods. For this study we generated a dataset from a random weighting of $b = 20$ random Fourier features of a squared exponential kernel [31] and corrupted by independent Gaussian noise. We use a generalized linear regression model as described in section 3.1 which uses the same features with $\tilde{m} = 3$. We consider a prior over $\sigma^2$, and a mean-field variational distribution giving $\tilde{m}(b + 1) = 63$ variational parameters which we initialize to be the same as the prior; a uniform categorical distribution. For DIRECT, a L-BFGS optimizer is used [32] and stochastic gradient descent is used for REINFORCE with a varying number of samples used for the Monte Carlo gradient estimator. Both methods use full batch training and are implemented using TensorFlow [33]. It can be seen that DIRECT greatly outperforms REINFORCE both in the number of iterations and computational time. As we move to a large $n$ or a larger $b$, the difference between the proposed DIRECT technique and REINFORCE becomes more profound. The superior scaling with respect to $n$ was expected since we had shown in section 3.1 that the DIRECT computational runtime is independent of $n$. However, the improved scaling with respect to $b$ is an interesting result and may be attributed to the fact that as the dimension of the variational parameter space increases, there is more value in having low (or zero) variance estimates of the gradient.

\[ ^2 \text{We used the identity } \left( \log q + 1 \right) \odot \frac{\partial \log q}{\partial \theta} = \frac{1}{2} \frac{\partial}{\partial \theta} \left( \log q + 1 \right)^2, \text{ where } \odot \text{ denotes an elementwise product.} \]
5.2 Relaxing Gaussian Priors on UCI Regression Datasets

In this section, we consider discretely relaxing a continuous Gaussian prior on the weights of a generalized linear regression model. This allows us to compare performance between a reparameterization gradient estimator for a continuous prior and our DIRECT method for a relaxed, discrete prior.

Considering regression datasets from the UCI repository, we report the mean and standard deviation of the root mean squared error (RMSE) from 10-fold cross validation. Also presented is the mean training time per fold on a machine with two E5-2680 v3 processors and 128Gb of RAM, and the expected sparsity (percentage of zeros) within a posterior sample. All models use $b = 2000$ basis functions. Further details of the experimental setup can be found in appendix E. In table 1, we see the results of our studies across several model-types. In the left column, the “REPARAM Mean-Field” model uses a (continuous) Gaussian prior, an uncorrelated Gaussian variational distribution and reparameterization gradients. The right two models use a discrete relaxation of a Gaussian prior (DIRECT) with support at 15 discrete values, allowing storage of each latent variable sample as a vector of 4-bit quantized integers. Therefore, each ELBO evaluation requires $15^{2000} > 10^{2352}$ log-likelihood evaluations, however, these computation can be done quickly by exploiting Kronecker matrix algebra. We compute the ELBO as described in section 3.1 for the “DIRECT Mean-Field” model, and use the low-variance, unbiased gradient estimator described in eq. (11) for the “DIRECT 5-Mixture SGD” model which uses a mixture distribution with $r = 5$ components, and $t = 3000$ Monte Carlo samples for the entropy gradient estimator.

The boldface entries indicate top performance on each dataset, where it is evident that the DIRECT method not only outperformed REPARAM on most datasets but also trained much faster, particularly on the large datasets due to the independence of dataset size on computational complexity. The DIRECT mean-field model contains $\tilde{m}b = 30,000$ variational parameters, however, training took just seconds on all datasets, including electric with over 2 million points. The DIRECT mixture model contains $\tilde{m}br = 150,000$ variational parameters, and since the gradient estimates are stochastic, average training times are on the order of hundreds of seconds across all datasets. While the time for precomputations does depend on dataset size, its contribution to the overall timings are negligible, being well under one second for the largest dataset, electric. Additionally, it is evident that posterior samples from the DIRECT model tend to be very sparse. For example, the DIRECT models on the gas dataset admit posterior samples that are over 84% sparse on average, meaning that over 1680 weights are expected to be zero in a posterior sample with $b = 2000$ elements. This would yield massive computational savings on hardware limited devices. Samples from the DIRECT models on the electric dataset are over 99.6% sparse.

Comparing the DIRECT mean-field model to the mixture model, we observe gains in the RMSE performance on many datasets, as we would expect with the increased flexibility of the variational distribution. While we only showed the posterior mean in our results, we would expect an even larger disparity in the quality of the predictive uncertainty which was not analyzed. In table 2 of the supplement, we present results for a DIRECT mixture model that uses the ELBO lower bound presented in Theorem 3. This model does not perform as well as the DIRECT mixture model trained using an unbiased SGD approach, as would be expected, however, it does train faster since its...
Table 1: Mean and standard deviation of test error, average training time, and average expected sparsity of a posterior sample from 10-fold cross validation on UCI regression datasets.

| Dataset     | n | d | Continuous Prior | Discrete 4-bit Prior |
|-------------|---|---|------------------|----------------------|
|             |   |   | REPARAM Mean-Field | DIRECT Mean-Field | DIRECT 9-Mixture SGD |
|             |   |   | Time RMSE Sparsity | Time RMSE Sparsity | RMSE Sparsity       |
| challenger  | 23| 4 | 8.0515 ± 0.284% 0%| 1.0523 ± 0.248 17%| 0.525 ± 0.246 17%   |
| fertility   | 100| 9 | 0.361 ± 0.043% 0%| 0.159 ± 0.041 17%| 0.16 ± 0.041 17%    |
| automobile | 159| 25| 0.425 ± 0.2 0%   | 0.129 ± 0.063 51%| 0.122 ± 0.056 51%   |
| servo       | 167| 4 | 0.524 ± 0.184% 0%| 0.271 ± 0.08 35% | 0.274 ± 0.077 35%   |
| cancer      | 194| 33| 27.488 ± 5.4 19% | 22.954 ± 3.09 19%| 22.837 ± 3.15 19%   |
| hardware    | 209| 7 | 1.796 ± 1.537 0% | 1.401 ± 0.048 51%| 0.401 ± 0.046 51%   |
| yacht       | 308| 6 | 0.815 ± 0.17 0%  | 0.234 ± 0.07 96% | 0.225 ± 0.082 96%   |
| autotmp     | 392| 7 | 4.05 ± 0.739 0%  | 2.564 ± 0.306 31%| 2.543 ± 0.362 31%   |
| housing     | 506| 15| 3.014 ± 0.567 0% | 2.752 ± 0.405 40%| 2.699 ± 0.361 39%   |
| forest      | 517| 12| 1.378 ± 0.148 0% | 1.363 ± 0.15 17% | 1.357 ± 0.155 17%   |
| stock       | 536| 11| 0.751 ± 0.338 0% | 0.011 ± 0.003 98%| 0.008 ± 0.001 98%   |
| pendulum    | 630| 9 | 1.465 ± 0.26 0%  | 1.329 ± 0.282 68%| 1.312 ± 0.253 63%   |
| energy      | 768| 8 | 78.852 ± 21.73 0%| 3.272 ± 0.332 99%| 2.911 ± 0.309 99%   |
| concrete    | 1030| 8 | 10.347 ± 2.847 0%| 5.316 ± 0.716 82%| 5.477 ± 0.632 82%   |
| solar       | 1066| 10| 0.902 ± 0.171 0% | 0.787 ± 0.192 23%| 0.788 ± 0.189 23%   |
| airfoil     | 1503| 5 | 2.071 ± 0.271 0% | 2.175 ± 0.349 48%| 2.156 ± 0.316 45%   |
| wine        | 1599| 11| 0.939 ± 0.33 0%  | 0.472 ± 0.044 54%| 0.469 ± 0.042 54%   |
| gas         | 2565|128| 0.27 ± 0.052 0%  | 0.211 ± 0.058 84%| 0.184 ± 0.063 76%   |
| skillcraft  | 3338|19 | 0.273 ± 0.029 0% | 0.253 ± 0.016 97%| 0.253 ± 0.016 97%   |
| sml         | 4137|26 | 0.327 ± 0.013 0% | 0.677 ± 0.044 57%| 0.671 ± 0.047 57%   |
| parkinsons  | 5875|20 | 0.158 ± 0.009 0% | 0.651 ± 0.034 13%| 0.613 ± 0.083 13%   |
| poleteile   | 15000|26 | 12.487 ± 0.363 0%| 13.65 ± 0.348 16%| 13.369 ± 0.431 17%  |
| elevators   | 16599|18 | 0.247 ± 0.156 0% | 0.124 ± 0.003 99%| 0.124 ± 0.003 99%   |
| protein     | 45730|9 | 0.642 ± 0.006 0% | 0.619 ± 0.007 76%| 0.618 ± 0.007 60%   |
| kegg        | 48827|20 | 0.178 ± 0.012 0% | 0.222 ± 0.009 96%| 0.205 ± 0.004 95%   |
| ctslice     | 53500|385| 4.415 ± 0.113 0%| 6.063 ± 0.122 19%| 5.478 ± 0.137 42%   |
| kegg4       | 63608|27 | 0.122 ± 0.004 0% | 0.139 ± 0.004 87%| 0.136 ± 0.006 87%   |
| 5brood      | 434874|3 | 11.057 ± 0.091 0%| 10.493 ± 0.105 40%| 10.354 ± 0.077 33%  |
| song        | 515345|90 | 0.037 ± 0.002 0% | 0.503 ± 0.002 32%| 0.498 ± 0.002 29%   |
| buzz        | 583250|77 | 0.94 ± 0.006 0%  | 1.007 ± 0.007 82%| 0.959 ± 0.004 80%   |
| electric    | 2049280|11 | 9.26 ± 4.47 0% | 5.575 ± 0.055 99.6% | 5.575 ± 0.055 99.6% |

6 Conclusions

We have shown that by discretely relaxing continuous priors, variational inference can be performed accurately and efficiently using our DIRECT method. We have demonstrated that through the use of Kronecker matrix algebra, the ELBO of a discretely relaxed model can be efficiently and exactly computed even when this computation requires significantly more log-likelihood evaluations than the number of atoms in the known universe. Through this ability to exactly perform ELBO computations we achieve unbiased, zero-variance gradient estimates using automatic differentiation which we show significantly outperforms competing Monte Carlo alternatives that admit high-variance gradient estimates. We also demonstrate that the computational complexity of ELBO computations is independent of the quantity of training data using the DIRECT method, making the proposed approaches amenable to big data applications. At inference time, we show that we can again use Kronecker matrix algebra to exactly compute the statistical moments of the parameterized predictive posterior distribution, unlike competing techniques which rely on Monte Carlo sampling. Finally, we discuss and demonstrate how posterior samples can be sparse and can be represented as quantized integer values to enable efficient inference which is particularly powerful on hardware limited devices, or if energy efficiency is a major concern.

We illustrate the DIRECT approach on several popular models such as mean-field variational inference for generalized linear models and deep Bayesian neural networks for regression. We also discuss some models which do not admit a compact representation for exact ELBO computations. For these cases, we discuss and demonstrate novel extensions to the DIRECT method that allow efficient computation of a lower bound of the ELBO, and we demonstrate how an unfactorized variational distribution can be used by introducing a manageable level of stochasticity into the gradients. We hope that these new approaches for ELBO computations will inspire new model structures and research directions in approximate Bayesian inference.
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A  Proof of Theorem 1: ELBO Computation for a Regression GLM

For our generalized linear regression model with a prior over \( \sigma^2 \), we can re-write eq. (7) as follows

\[
\text{ELBO}(\theta) = (q_\sigma \otimes q)^T \log \ell + \sum_{i=1}^{b} q_i^T \log p_i - \sum_{i=1}^{b} q_i^T \log q_i + q_\sigma^T \log p_\sigma - q_\sigma^T \log q_\sigma,
\]

where we have simply expanded the factorized variational distribution to include \( \sigma^2 \), resulting in the two extra terms. To complete the ELBO in eq. (12), we need to take the inner product between the variational distribution and log-likelihood for each point in the hypothesis space, \( (q_\sigma \otimes q)^T \log \ell \).

We can write this relation as follows for our generalized linear regression model, (see e.g. [34])

\[
(q_\sigma \otimes q)^T \log \ell = -\frac{n}{2} q_\sigma^T \log \sigma^2 - \frac{1}{2} (q_\sigma \sigma^2) \left( q^T \{(y - \Phi w_i)^T(y - \Phi w_i)\}_{i=1}^m \right),
\]

whose computation would be prohibitively expensive when \( m = \tilde{m}b \) is large. We will now focus on computing the inner product involving the variational distribution over the \( w \) variables, \( q \), which we can break into three terms as follows,

\[
q^T \{(y - \Phi w_i)^T(y - \Phi w_i)\}_{i=1}^m = y^T y - 2q^T \{y^T \Phi w_i\}_{i=1}^m + q^T \{w_i^T \Phi^T \Phi w_i\}_{i=1}^m,
\]

for which the first term is trivial to compute as written since it does not depend on \( w \). We now demonstrate how the second and third terms can be computed, recalling that we have assumed that \( q \) is a mean-field variational distribution. Firstly, define \( Z = (\Phi W)^T = \{\phi_{ij} \}_{j=1}^b \in \mathbb{R}^{m \times n} \) whose columns contain the model prediction for a single training point at every possible set of latent variable values in the hypothesis space. Observe that each column is represented as a sum of \( b \) Kronecker product vectors. We can then write the second term of eq. (14) as

\[
q^T \{y^T \Phi w_i\}_{i=1}^m = \sum_{i=1}^{n} y_i q_i^T z_i = \sum_{i=1}^{n} y_i \sum_{j=1}^{b} \phi_{ij} q_j^T w_j = \sum_{j=1}^{b} q_j^T w_j \left( \sum_{i=1}^{n} y_i \phi_{ij} \right) = s^T \Phi^T y,
\]

where \( s = \{q_j^T w_j\}_{j=1}^b \in \mathbb{R}^b \). Finally, considering the third term of eq. (14), observe that we can write \( \{w_i^T \Phi^T \Phi w_i\}_{i=1}^m = \sum_{i=1}^{m} z_i^2 \), and since each \( z_i \) is a sum of \( b \) Kronecker product vectors, \( z_i \) a sum of \( b + b^2 \) Kronecker product vectors. We can then write the third term of eq. (14) as follows,

\[
q^T \{w_i^T \Phi^T \Phi w_i\}_{i=1}^m = \sum_{i=1}^{n} \sum_{j=1}^{b} q_j^T \bar{w}_j^2 \phi_{ij}^2 + \sum_{k=j+1}^{b} \phi_{ij} \phi_{ik} \left( q_j^T \bar{w}_j \right) \left( q_k^T \bar{w}_k \right),
\]

where we have used the short-hand notation \( H = \{\bar{w}_j^2 \sum_{i=1}^{n} \phi_{ij} \}_{j=1}^b \in \mathbb{R}^{b \times b} \). Substituting eq. (15) and eq. (18) into eq. (14), we can re-write the inner product between the variational distribution and the log-likelihood in eq. (13) as follows,

\[
(q_\sigma \otimes q)^T \log \ell = -\frac{n}{2} q_\sigma^T \log \sigma^2 - \frac{1}{2} (q_\sigma \sigma^2) \left( y^T y - 2s^T \Phi^T y \right) + s^T \Phi^T \Phi s - \text{diag}(s^T \Phi^T \Phi s) =
\]

\[
\text{diag}(s^T \Phi^T \Phi s) + \sum_{j=1}^{b} q_j^T h_j,
\]

and substituting this into eq. (12) completes the proof.

\( \square \)

B  Proof of Theorem 2: Logistic Regression ELBO Lower Bound

For the generalized linear logistic regression model considered, we can write the log likelihood as follows (see e.g. [34])

\[
\log \ell = \sum_{i=1}^{n} -y_i z_i - \log (1 + \exp(-z_i)),
\]
where \( \mathbf{Z} = (\mathbf{\Phi W})^T = \{\oplus_{j=1}^b \phi_{ij} \mathbf{w}_j\}_{i=1}^n \in \mathbb{R}^{m \times n} \) is a matrix whose columns contain the log values for a single training point at every possible set of latent variables in the hypothesis space. It is evident that the first term is identical to that discussed in eq. (15), however, computation of the second term requires more development. We can write

\[
\mathbf{q}^T \log \ell = -s^T (\mathbf{\Phi}^T \mathbf{y}) - \sum_{i=1}^n \mathbf{q}^T \log (1 + \exp(-\mathbf{z}_i)).
\]  

(21)

Since \( \mathbf{z}_i = \oplus_{j=1}^b \phi_{ij} \mathbf{w}_j \in \mathbb{R}^m \) is a sum of \( b \) Kronecker product vectors, each with one unique sub-matrix that is not unity, \( \exp(-\mathbf{z}_i) \) is a single Kronecker product vector. This follows from Proposition 2. We can then take a Taylor series explanation of \( \log (1 + \exp(-\mathbf{z}_i)) \) as follows

\[
\log(1 + \exp(-\mathbf{z}_i)) = -\sum_{k=1}^\infty \frac{(-1)^k \exp(-k\mathbf{z}_i)}{k} \quad \text{for } \exp(-\mathbf{z}_i) < 1 \rightarrow \mathbf{z}_i > 0,
\]

(22)

\[
\log(1 + \exp(-\mathbf{z}_i)) = -\mathbf{z}_i - \sum_{k=1}^\infty \frac{(-1)^k \exp(k\mathbf{z}_i)}{k} \quad \text{for } \exp(-\mathbf{z}_i) > 1 \rightarrow \mathbf{z}_i < 0,
\]

(23)

and although the use of either choice would result in an ELBO lower bound, we choose the approximation based on the training label as follows; if \( y_i = 0 \) or \( y_i = 1 \) then we would choose the \((\mathbf{z}_i > 0)\) or \((\mathbf{z}_i < 0)\) approximation, respectively. We choose this because \( z_i > 0 \) gives a higher class conditional probability to class 0 than class 1 so this approximation would yield a tight lower bound when the training examples are correctly classified. These approximations are plotted in fig. 2 with a first-order expansion where it is evident that the computation lower-bounds the exact computation. Using this first-order Taylor series approximation, we can write our lower bound for the inner product between the variational distribution and the log-likelihood as follows which completes the proof,

\[
\mathbf{q}^T \log \ell \geq -s^T (\mathbf{\Phi}^T \mathbf{y}) - \sum_{i=1}^n \left\{ \begin{array}{ll}
\mathbf{q}^T \exp(-\mathbf{z}_i) & \text{if } y_i = 0 \\
\mathbf{q}^T \exp(\mathbf{z}_i) - \mathbf{q}^T \mathbf{z}_i & \text{if } y_i = 1
\end{array} \right.
\]

(24)

\[
= -s^T (\mathbf{\Phi}^T \mathbf{y}) - \sum_{i=1}^n \left\{ \begin{array}{ll}
\prod_{j=1}^b \mathbf{q}^T \exp(-\phi_{ij} \mathbf{w}_j) & \text{if } y_i = 0 \\
\prod_{j=1}^b \mathbf{q}^T \exp(\phi_{ij} \mathbf{w}_j) - \sum_{j=1}^b \mathbf{q}_i^T \phi_{ij} \mathbf{w}_j & \text{if } y_i = 1
\end{array} \right.
\]

(25)

\[\square\]

Remark We expect these Taylor series approximations to admit a tight bound within and just outside of their logit domains as we can see in fig. 2. Equivalently, the log-likelihood approximation is accurately computed for hypotheses that correctly classify the training data when we use this lower bound, however, hypotheses that confidently misclassify training labels may be over-penalized. This can be seen by observing how the approximations in fig. 2 significantly underestimate the exact solution when they are far outside of the approximations domain.

Remark In practice, the products over \( b \) terms in Theorem 2 may result in overflow or loss of precision, however, computations can be performed in a stable manner in logit space and the LogSumExp trick [35] can be used to avoid precision loss for the sum over \( n \).

C Proof of Theorem 3: Mixture Distribution Entropy Lower Bound

We begin by taking a Taylor series approximation of \( \log \mathbf{q} \) about \( \mathbf{a} = \oplus_{j=1}^b \mathbf{a}_i, \mathbf{a}_i \in (0,1)^m \) as follows,

\[
\log \mathbf{q} = \log \mathbf{a} + \sum_{k=1}^\infty \frac{(-1)^{(k+1)} (\mathbf{q} - \mathbf{a})^k}{k \mathbf{a}^k},
\]

(26)

which can be represented as a sum of Kronecker product vectors once the exponents are computed explicitly. However, the number of terms in this sum will grow quickly with respect to the order of the Taylor series approximation. When a first order Taylor series expansion is considered, the approximation will give a strict lower bound of \(- \log \mathbf{q}\) and consequently a lower bound of the ELBO (eq. (7)) will be achieved. The approximation for a linear Taylor series expansion is plotted in
The following image denotes the structure of a neuron that we will use in our neural network. In order to demonstrate DIRECT computation of the log-likelihood for a Bayesian neural network we will first perform a forward-pass through the neural network from top to bottom, however, unlike a layer but grows as we observe deeper layers. Using this nomenclature it is evident that we can move from one layer to the next. 

**Nomenclature and Neuron Structure** At all points in the forward-pass we can represent the internal (or final) state of the neural network with a special structure which is a sum of Kronecker product vectors as follows for \( i = 1, \ldots \), (number of neurons in the layer), and \( l = 1, \ldots, n \),

\[
\mathbf{u}_l^{(i)} = \sum_{j=1}^{h} c_{jl} \bigotimes_{k=1}^{b} g_{jk}^{(i)} ,
\]

where \( \mathbf{U}^{(i)} = (\mathbf{u}_l^{(i)})_{l=1}^{n} \in \mathbb{R}^{m_{b} \times n} \), \( \mathbf{u}_l^{(i)} \in \mathbb{R}^{m_{b}} \) denotes the internal state of the \( i \)th neuron of the current layer, and both \( \mathbf{G}^{(i)} = (\{\mathbf{g}_{jk}^{(i)}\})_{j=1}^{h} \bigotimes_{k=1}^{b} \in \mathbb{R}^{h \times b \times m_{b}} \), \( \mathbf{g}_{jk}^{(i)} \in \mathbb{R}^{m_{b}} \) and \( \mathbf{C} \in \mathbb{R}^{h \times n} \) change as we move from one layer to the next. \( h \) depends on the network architecture and it is constant throughout a layer but grows as we observe deeper layers. Using this nomenclature it is evident that we can compactly represent the internal state of any location within the neural network while we compute our forward pass.

The following image denotes the structure of a neuron that we will use in our neural network.

**Remark** The products over \( b \) terms might seem problematic, however, we do not expect the final results to be too large to be an overflow concern. To avoid precision loss, we compute the log of the products, which can be done stably, and then exponentiate.

**D DIRECT Bayesian Neural Networks for Regression**

In order to demonstrate DIRECT computation of the log-likelihood for a Bayesian neural network we will first perform a forward-pass through the neural network from top to bottom, however, unlike a conventional forward-pass is conventionally conducted in literature where the network is fixed at a specific location in the hypothesis space, we will simultaneously evaluate the neural network at all locations in entire hypothesis space. Consequently, a forward-pass through the neural network with our training set will give us \( m_{b}^{b} \times n \) values.

The products over \( b \) terms might seem problematic, however, we do not expect the final results to be too large to be an overflow concern. To avoid precision loss, we compute the log of the products, which can be done stably, and then exponentiate.
For clarity of illustration, we will not discuss the bias term although this can be easily added by associating a latent variable with a layer input that is fixed to unity. In our discussion, we will break the computation of the neuron into two stages; the first will involve multiplication of the layer inputs with the latent variables as well as the summation, and the second stage will involve passing this summation through a non-linear activation function.

**Multiplication with Latent Variables and Summation** Our computational neurons begin by multiplying the layer inputs with a specific latent variable and then summing up these values. Assuming we are conducting a forward-pass moving deeper into the network and are currently at the “layer inputs” location in our computational neuron figure, the internal state for the $i$th input is denoted by $U^{(i)}$, whose structure is defined in eq. (28). We must multiply this state by all possible values of the corresponding latent variable, which we will assume is indexed as the $p$th of our $b$ latent variables. We can easily perform this multiplication as follows for $l = 1, \ldots, n$

$$u_l^{(i)} = \left( \sum_{j=1}^{b} C_{jl} \bigotimes g_{jk}^{(i)} \right) \odot W[p,:)^T,$$

$$= \sum_{j=1}^{b} C_{jl} \left( \bigotimes_{k=1}^{p-1} g_{jk}^{(i)} \right) \odot \left( g_{jp}^{(i)} \odot W_p \right) \odot \left( \bigotimes_{k=p+1}^{b} g_{jk}^{(i)} \right) = \sum_{j=1}^{b} C_{jl} \bigotimes g_{jk}^{(i)},$$

where $\odot$ denotes element-wise multiplication, and we have taken advantage of the Kronecker product structure of the rows of $W$ as depicted in eq. (3). Finally, the summing operation is straightforward for our computational neuron. It simply involves summing the multiplied inputs from each layer input as follows,

$$\sum_{i=1}^{\text{num. inputs}} \sum_{j=1}^{b} C_{jl} \bigotimes g_{jk}^{(i)}.$$  

At this point we would update $h, G$ and $C$ to convert this double summation into a single summation before passing through the non-linear activation function, as we will discuss next.

**Quadratic Activation** We will use a quadratic activation function for our neural network. Any other non-linear activation could be used, however, we choose the quadratic since it allows a more compact representation of internal state of the network to be maintained, i.e. allows for a small $h$ versus other non-linear activations. Again assuming that the current state at the $i$th neuron is defined by $U^{(i)}$, the output for the activation function for the $i$th neuron is as follows for $l = 1, \ldots, n$

$$u_l^{(i)} = u_l^{(i)} \odot u_l^{(i)} = \left( \sum_{j=1}^{b} C_{jl} \bigotimes g_{jk}^{(i)} \right) \odot \left( \sum_{j=1}^{b} C_{jl} \bigotimes g_{jk}^{(i)} \right),$$

$$= \sum_{j=1}^{b} C_{jl}^2 \bigotimes g_{jk}^{(i)} \odot g_{jk}^{(i)} + 2 \sum_{j=1}^{b} C_{jl} g_{jk}^{(i)} \bigotimes g_{jk}^{(i)},$$

and at this point we would update $h, G$ and $C$ to convert this double summation into a single summation to represent the internal state compactly before moving deeper.
We now demonstrate how the second and third terms can be computed, recalling we assume
where we assume that we have already conducted algorithm forward_pass
Algorithm for which the first term is trivial to compute as written since it does not depend on the latent variables.

For stability, all matrices could be represented by storing both the sign and logarithm of all entries. For deep networks, this could be advantageous to avoid precision loss. Nonetheless, we will proceed with the algorithm as presented, for purposes of clarity.

Forward-Pass Algorithm Using the previously defined operations, we can summarize the forward-pass procedure in algorithm forward_pass. Note that algorithm forward_pass is simplified for clarity of presentation. The computations involved could be performed far more efficiently and in a more stable manner. For example, the vast majority of entries in the $G$ matrices are unity, so identifying this could massively decrease storage and computational requirements. Additionally, $\hat{C}$ evidently has a Kronecker product structure which could be carefully exploited to yield benefits for very wide neural networks. For stability, all matrices could be represented by storing both the sign and logarithm of all entries. For deep networks, this could be advantageous to avoid precision loss.

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Algorithm forward_pass Perform a forward pass for through the neural network using the entire training set and simultaneously computing the outputs for all $m = \tilde{m}^h$ points in the hypothesis space. $\text{mult}_\text{var}$ multiplies the current state with the appropriate latent variable as is done in eq. (30), $\text{neuron}_\text{sum}$ computes the neuron sum as is done in eq. (31), and $\text{activation}$ computes the non-linear activation function as is done in eq. (33). All the pseudo-functions defined take $G$ and/or $C$ and perform the necessary computations with those inputs. We omit latent-variable indexing values for clarity of presentation.

\begin{align*}
\text{Input: } X & \in \mathbb{R}^{n \times d} \\
\text{Output: } C & \in \mathbb{R}^{h \times n} \text{ \& } G \in \mathbb{R}^{h \times b \times \tilde{m}} \text{ which define state } U \in \mathbb{R}^{\tilde{m}^h \times n} \text{ in eq. (28)} \\
& C = X^T, \; G^{(i)} = \text{ones}(1 \times b \times \tilde{m}), \; i = 1, \ldots, d \\
\text{for each layer do} & \\
& \text{for } j = 1 \text{ to num. neurons in layer do} \\
& \text{for } i = 1 \text{ to num. inputs to layer do} \\
& G^{(i)} = \text{\text{mult}_\text{var}}(G^{(i)}) \text{ \quad multiplication with the appropriate row of } W \\
\text{end for}
\end{align*}

\begin{align*}
\text{end for} & \\
& \tilde{G}^{(j)} = \text{\text{neuron}_\text{sum}}(G^{(1)}, \ldots, G^{(\text{num. inputs})}) \text{ \quad sum operation for the current (jth) neuron} \\
& \text{if not last layer then} \\
& \tilde{\gamma}^{(j)} \sim \tilde{C} = \text{\text{activation}}(\tilde{G}^{(j)}, \tilde{C}) \\
\text{end if} & \\
\text{end for}
\end{align*}

\begin{align*}
& C = \tilde{C}, \; G^{(j)} = \tilde{\gamma}^{(j)}, \; j = 1, \ldots, \text{num. neurons in layer} \quad \text{update variables} \\
\text{end for} & \\
& G = G^{(1)} \quad \text{only one neuron in the last (output) layer, so remove indexing}
\end{align*}

ELBO Computation Computation of the ELBO will proceed similarly to the GLM regression model in section 3.1, however, there are several differences since we no longer have constant basis functions so our state representation is more complicated. We will again assume a Gaussian noise model for the observed responses and will again place a prior over the Gaussian variance. We can then modify eq. (13) which focuses on the ELBO term related to the log-likelihood as follows

\begin{align*}
(q_\sigma \otimes q)^T \log \ell &= -\frac{n}{2} q_\sigma^T \log \sigma^2 - \frac{1}{2} (q_\sigma^T \sigma^{-2}) (q^T \{ y - U[i,:)T \}^T (y - U[i,:)T)_{i=1}^m), \quad (34)
\end{align*}

where we assume that we have already conducted algorithm forward_pass such that the state $U$ represents the output of the neural network. We will now focus on computing the inner product involving the variational distribution over the $w$ variables, $q$, which we can break into three terms as follows,

\begin{align*}
q^T \{ y - U[i,:]T \}^T (y - U[i,:)T)_{i=1}^m &= \\
& y^T y - 2q^T \{ y^T U[i,:]T \}_{i=1}^m + q^T \{ U[i,:]U[i,:]T \}_{i=1}^m, \quad (35)
\end{align*}

for which the first term is trivial to compute as written since it does not depend on the latent variables. We now demonstrate how the second and third terms can be computed, recalling we assume $q$ is a mean-field variational distribution (although we can extend beyond mean-field using the techniques
discussed in section 4). Considering the second term in eq. (35), we can write

\[ q^T \{ y^T U[i,:] \}_{i=1}^m = q^T \left( \sum_{k=1}^n y_k \sum_{j=1}^h c_{jk} \otimes g_{ij} \right) = \sum_{j=1}^h \left( \sum_{k=1}^n y_k c_{jk} \right) \prod_{i=1}^b q_i^T g_{ij}, \]

where we have used the short-hand notation \( p = \{ \sum_{k=1}^n y_k c_{jk} \}_j \in \mathbb{R}^h \). Finally, considering the third term in eq. (35), we can write

\[ q^T \{ U[i, :]U[i, :]^T \}_{i=1}^m = q^T \left( \sum_{i=1}^n u_i \otimes u_i \right) = q^T \left( \sum_{i=1}^n \sum_{j=1}^h c_{ji} c_{ki} \otimes (g_{jl} \otimes g_{kl}) \right), \]

where we define \( V = \{ \sum_{i=1}^n c_{ji} c_{ki} \}_{j,k} \in \mathbb{R}^{h \times h} \). Substituting eq. (36) and eq. (39) into eq. (35), we can now compute the inner product between the variational distribution and the log-likelihood in eq. (34). The other terms required to compute the ELBO can be seen in eq. (12), and the computation of these other terms do not differ from the case of the generalized linear regression model. So we can now tractably compute the ELBO for our DIRECT Bayesian neural network.

We can pre-compute the terms \( y^T y \), \( p \), and \( V \) before training begins (since these do not depend on the variational parameters) such that the final complexity of the DIRECT method is independent of the number of training points, making the proposed techniques ideal for massive datasets. Also, it is evident that each of these pre-computed terms can easily be updated as more data is observed making the techniques amenable to online learning applications. If we assume a neural network with \( l \) hidden layers and an equal distribution of latent variables between layers, the computational complexity of the ELBO computations are \( \mathcal{O}(\ell m (b/\ell)^{2l}) \). This can be seen by observing eq. (39) and noting that \( h = \mathcal{O}(l (b/\ell)^{2l}) \), and that only \( \mathcal{O}(\ell) \) of the vectors in \{\( g_{jl} \}_{l=1}^b \} are not unity for any value of \( j = 1, \ldots, h \), allowing computations to be saved.

### E UCI Regression Studies Setup & Additional Results

Considering regression datasets from the UCI repository, we report the mean and standard deviation of the root mean squared error (RMSE) from 10-fold cross validation\(^4\). Also presented is the mean training time per fold on a machine with two E5-2680 v3 processors and 128Gb of RAM, and the expected sparsity (percentage of zeros) within a posterior sample. Using a generalized linear model, we consider \( b = 2000 \) random Fourier features of a squared-exponential kernel with automatic relevance determination \([31]\). Before generating the features, we initialize the kernel hyperparameters including the prior variance \( \sigma_w^2 \) and the Gaussian noise variance \( \sigma^2 \) by maximizing the marginal likelihood of an exact Gaussian process constructed on \( \min(n, 1000) \) points randomly selected from the dataset \([36]\). All discretely relaxed models (containing “DIRECT”), only have support at \( w \in \text{linrange}(-3\sigma_w, 3\sigma_w, \bar{m}=15) \), allowing \( w \) to be stored as 4-bit quantized integers.

For REPARAM we perform doubly stochastic optimization using a mini-batch size of 100 and using 10 Monte Carlo samples for the gradient estimates at each iteration. For datasets with \( n < 3000 \) we optimize for 1000 iterations and we optimize for 10000 iterations for all larger datasets. This model was implemented in Edward \([37]\). For the DIRECT mean-field model we use an L-BFGS optimizer \([32]\) and run until convergence, or 1000 iterations are reached. For the DIRECT 5-mixture model we perform stochastic gradient descent using \( t = 3000 \) Monte Carlo samples for the entropy gradient estimator eq. (11).

\(^4\)90% train, 10% test per fold. We use folds from https://people.orie.cornell.edu/andrew/code
Table 2: Using a mixture variational distribution along with the ELBO lower bound presented in Theorem 3, we present the mean and standard deviation of test error, average training time, and average expected sparsity of a posterior sample from 10-fold cross validation on UCI regression datasets.

For the DIRECT mean-field model we initialize the variational distribution to the prior. For the DIRECT mixture models, we first run the mean-field model and then initialize each mixture component to be randomly perturbed from the mean-field solution, and we initialize \( \alpha \) to the mean-field solution. We initialize the mixture probabilities to be constant.

For predictive posterior mean computations, we use the exact computation presented in eq. (9) for both the DIRECT and mixture models. For REPARAM, we approximate the mean by sampling the variational distribution using 1000 samples.

In table 2 we consider again an unfactorized mixture variational distribution, however, we maximize the ELBO lower bound derived in Theorem 3. Since the ELBO gradients are deterministic, we again use an L-BFGS optimizer for training. In addition to the 150,000 variational parameters used by the DIRECT 5-Mixture SGD model in table 1, computing the ELBO lower bound involves the simultaneous optimization of \( \alpha \), adding 30,000 additional optimization parameters.