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**Latent Derivative Bayesian Last Layer Networks**

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Bayesian neural networks (BNN) [43, 50] offer the possibility of combining the expressivity of neural networks with the principled uncertainty quantification and regularization derived from Bayesian methods. However, inference for priors over the weights is intractable, resulting in extensive study of learning such BNNs via approximate inference [50, 31, 19, 25, 38, 7]. Despite their many varieties, these approximate models can suffer from several drawbacks, such as unintuitive priors, expensive training procedures, inaccurate posteriors and/or large model parameter spaces. Moreover, these models are typically restricted to sampling from implicit predictive densities, and have been criticized for their inaccurate uncertainty quantification [24, 23, 53, 54, 81, 85]. In many risk-sensitive and safety-critical applications, such as in medical diagnosis [21] and model-based control [17], well-calibrated predictive uncertainty is essential when the model is used outside of the data distribution (OOD). In contrast to Bayesian neural networks, Gaussian processes (GP) offer exact nonlinear, non-parametric Bayesian modeling through linear regression of a rich (often infinite) feature space, specified by a derived kernel function [63]. While a powerful and popular tool for probabilistic modeling [29], exact inference computation does not scale gracefully for large datasets, and the model’s quality depends heavily on the choice of kernel given the data. Moreover, some kernels have been shown to suffer from the curse of dimensionality due to their use of distance metrics in the data space [4]. Despite sparse methods improving scalability [72, 75], parametric models still provide an attractive offer of

1 Introduction

Bayesian neural networks (BNN) [43, 50] offer the possibility of combining the expressivity of neural networks with the principled uncertainty quantification and regularization derived from Bayesian methods. However, inference for priors over the weights is intractable, resulting in extensive study of learning such BNNs via approximate inference [50, 31, 19, 25, 38, 7]. Despite their many varieties, these approximate models can suffer from several drawbacks, such as unintuitive priors, expensive training procedures, inaccurate posteriors and/or large model parameter spaces. Moreover, these models are typically restricted to sampling from implicit predictive densities, and have been criticized for their inaccurate uncertainty quantification [24, 23, 53, 54, 81, 85]. In many risk-sensitive and safety-critical applications, such as in medical diagnosis [21] and model-based control [17], well-calibrated predictive uncertainty is essential when the model is used outside of the data distribution (OOD). In contrast to Bayesian neural networks, Gaussian processes (GP) offer exact nonlinear, non-parametric Bayesian modeling through linear regression of a rich (often infinite) feature space, specified by a derived kernel function [63]. While a powerful and popular tool for probabilistic modeling [29], exact inference computation does not scale gracefully for large datasets, and the model’s quality depends heavily on the choice of kernel given the data. Moreover, some kernels have been shown to suffer from the curse of dimensionality due to their use of distance metrics in the data space [4]. Despite sparse methods improving scalability [72, 75], parametric models still provide an attractive offer of
Figure 2: A toy example depicting Bayesian modeling of a complex function from sparse data. The Gaussian process has well-calibrated uncertainty, but its fixed kernel structure can result in an undesirable function space for inference. Weight space mean-field variational inference (VI) exhibits poor in-between uncertainty and has an implicit predictive density. The Bayesian last layer has an explicit density and a deterministic feature space. However, these features overfit, resulting in poor epistemic uncertainty too. We introduce a 'latent derivative' prior to the BLL that encourages variance in the model’s Jacobian distribution, diversifying the feature space by capitalizing on the network’s large hypothesis space. This diversity results in increased predictive uncertainty, without sacrificing the fit.

flexible model specification with data-independent computation and parameterization.

The Bayesian last layer [39, 51] is an alternative BNN approach in the spirit of GPs, combining a Bayesian linear model with a learned, finite feature space, represented by a neural network. While the linear model ensures the analytical tractability of both inference and predictive distribution, the neural features provide the broad, adaptive hypothesis space offered by neural network architectures. Since Gaussian processes are just Bayesian linear models in an expressive feature space, can’t learned neural features perform as well as kernels? While the network can be trained easily using gradient descent on the negative marginal likelihood, there is a catch: The overparameterization leads to overfitting of the feature space [39, 63], resulting in a reduced hypothesis space of functions which severely limits the predictive uncertainty quantification (Figure 2).

To encourage diversity in the BLL’s neural features, without sacrificing the model’s attractive properties, we incorporate a novel functional prior into the model specification. We posit that well-calibrated uncertainty quantification may be effectively characterized by the distribution of the model’s Jacobian w.r.t. the network’s inputs (Figure 2), which is also a Gaussian process [63, 75]. Previous methods to improve BNN uncertainty quantification rely on additionally modeling the data distribution, requiring OOD samples to explicitly boost the predictive uncertainty [27]. The derivative prior works in- and outside the data distribution, influencing the model’s hypothesis space and therefore epistemic uncertainty directly. By incorporating this prior into the objective, using the functional KL divergence (fKL) [68, 16, 76], the smoothness and diversity of the feature space is influenced by the variance of the prior. As a result, this training procedure resembles functional variational inference (fVI) [76]. Due to the BLL’s deterministic features, the Jacobian may be computed directly using forward mode automatic differentiation (AD) [61], and the distribution over Jacobians can be obtained in closed-form thanks to the Bayesian last layer. However, during training the divergence to the functional prior must be approximated using samples. Moreover, we believe this prior is intuitive, and its functional nature should enable the model to remain suitably calibrated independent of model size, as Bayesian models should [62].

By combining the analytic convenience of the Bayesian last layer, forward mode automatic differentiation and the novel functional prior over the Jacobian, we present a practical, calibrated Bayesian neural network that offers comparable utility to Gaussian processes, across small and large tasks. This class of BNN makes the case, like GPs, that priors are not required over the feature parameters. This reduction of complexity is motivated for applied domains such as robotics, where fast, well-calibrated Bayesian models are needed for sample-efficient, safe and risk-averse settings such as model-based reinforcement learning [18]. In these domains, the balance between simplicity and performance is key for practical use, and ideally not dependent on the amount of data in the task. To evaluate the benefit of this prior, we compare against standard BLLs and other baselines for OOD regression, active learning and Bayesian optimization, where epistemic uncertainty

Note that the arcsine kernel is equivalent to an infinite hidden layer network with erf activations, therefore shares a similar hypothesis space to sigmoid and tanh networks.
is needed. We show that the latent derivative (LD) prior enhances the BLL’s predictive uncertainty, scaling to large, real-world datasets with high-dimensional inputs, achieving superior or comparable performance on key tasks. We also introduce two new benchmarks for OOD prediction and active learning that utilize existing datasets from real-world robotic systems.

2 The Bayesian Last Layer

Intuitively, Bayesian last layer networks can be viewed as Bayesian linear regression in a projected feature space, where the projection is learned by a neural network. Alternatively, they can be thought of as a neural network whose parameters of the last layer are integrated out via exact, analytical Bayesian inference. While the BLL can be easily deployed for multivariate integration, the following derivations (and later experiments) in this work focus on univariate targets for simplicity.

Let \( D = \{(x_i, y_i)\}_{i=1}^{n} \) be the observed data, \( x_i \in \mathbb{R}^k \), \( y_i \in \mathbb{R} \), such that \( X \in \mathbb{R}^{n \times k} \) and \( y \in \mathbb{R}^n \). Additionally, let \( \phi(\cdot; \theta) : \mathbb{R}^k \rightarrow \mathbb{R}^m \) be a feature space projection with parameters \( \theta \), \( \phi_i = \phi(x_i; \theta) \) and \( \Phi = [\phi_1, \ldots, \phi_n]^T \in \mathbb{R}^{n \times m} \), the matrix of vertically stacked row vectors. It is common to add constant or linear terms to \( \Phi \) to implicitly represent bias or identity terms. However, for notational clarity, we ignore these terms and denote the projected feature space as \( \mathbb{R}^m \).

A latent function \( f \) is modeled using Bayesian linear regression with weights \( \beta \) and additive, zero-mean, Gaussian noise \( \epsilon \) with variance \( \sigma^2 \), where

\[
y_i = f(x_i; \theta) = \phi_i^T \beta + \epsilon_i.
\] (1)

Placing a conjugate Gaussian prior \( \mathcal{N}(\mu_0, \Lambda_0^{-1}) \) over \( \beta \) results in a Gaussian posterior \( \mathcal{N}(\mu_n, \Lambda_n^{-1}) \) with an explicit Gaussian predictive distribution for query \( x \),

\[
y \mid x, D, \theta \sim \mathcal{N}(\mu_n, \sigma^2 + \phi_x^T \Lambda_n^{-1} \phi_x),
\] (2)

where \( \mu_n \) and \( \Lambda_n \) are the mean vector and precision matrix of the posterior weight distribution.

The observation noise \( \sigma^2 \), prior weight parameters \( \mu_0 \) and \( \Lambda_0 \), and \( \theta \) can be either set to constants or optimized jointly by maximizing the log-marginal likelihood. With \( \mu_0 = 0 \), this model is equivalent to a Gaussian process with kernel \( k(x, x'; \theta) = \phi(x; \theta)^T \Lambda_0^{-1} \phi(x'; \theta) \) [63]. For a more Bayesian treatment, an inverse gamma prior can be placed on \( \sigma^2 \), inducing a Student-t weight posterior and predictive density (Section A). We use ‘GBLL’ and ‘TBLL’ to differentiate these two approaches.

3 Latent Derivative Priors

For the Bayesian last layer, the neural features \( \phi \) are optimized using type-II maximum likelihood on the marginal likelihood (Equation (18)). While type-II ML can be effective at tuning hyperparameters, e.g. the lengthscale of a kernel, optimizing too many parameters runs the risk of overfitting [63]. For the BLL, this manifests as the feature space converging about the mean function. While this results in adequate uncertainty far away from the data, predictions are overconfident between datapoints (Figure 2).

To improve the diversity of the feature space, we are motivated to augment the marginal likelihood objective to leverage the expressiveness of the neural network without sacrificing fit. In this work, we build on the intuition that the distribution of the model’s derivatives influences the epistemic uncertainty OOD (Figure 3). Given that the derivative of a Gaussian process is also a Gaussian process [66], computing the feature Jacobian \( J_{\phi} \), using forward mode AD allows us to reason about the predictive Jacobian in closed-form, which for 1D regression is a vector-valued, probabilistic function, i.e. a Gaussian process, which we denote \( z \),

\[
\frac{\partial f}{\partial x}(x) := z(x) = J_{\phi_{x}}^\top \beta, \quad z \sim p(\cdot \mid x, D, \theta),
\] (3)

\[
p(z \mid x, D, \theta) = \mathcal{GP}(z | J_{\phi_{x}}^\top \mu_n, J_{\phi_{x}}^\top \Lambda_n^{-1} J_{\phi_{x}}). \quad (4)
\]

In typical regression, \( z \) is unobserved and therefore a quantity we wish to remain uncertain about. Moreover, with expressive function approximators we should be free to shape the uncertainty of \( z \) without interfering with the fit of \( f \). In the Bayesian framework, we can shape this uncertainty by placing a functional prior \( \pi \) on \( z \in \mathbb{R}^k \)

\[
\min_{\theta} D_{\text{KL}}(\pi(z \mid x) \parallel p(z \mid x, D, \theta)),
\] (5)

enforced through a functional KL divergence (fKL).
Combining the conventional marginal likelihood with this fKL, we propose a novel joint objective

$$\max_{\theta} \log p(D \mid \theta) - D_{KL}(\pi(z \mid x) \mid\mid p(z \mid x, D, \theta)), \quad (6)$$

which can be interpreted from two perspectives.

**Maximum Entropy Regularization** Since Bayesian linear regression typically considers a fixed feature space or kernel, jointly learning the features can be viewed as an inverse problem [17]. Inverse problems, especially in probabilistic settings, are commonly regularized using the principle of maximum entropy [34, 71]. Choosing the features that are the most unstructured, or the least committed to a specific model, offers not just robustness but ideally translates to calibrated epistemic uncertainty in the predictions, as illustrated how controlling the distribution of $z$ does not influence the likelihood of the observations, as $\delta = 0$ for the training data. Also, our fKL uses the forward KL (M-projection) instead of the reverse KL (I-projection), which the ELBO uses. The forward KL encourages the distribution to cover as much probability mass as possible which translates to a flat distribution with higher variance, i.e. higher entropy, whereas the reverse KL prefers to seek an individual mode which typically results in lower variance and potential overfitting [17]. We discuss this topic in more detail in Section C of the Appendix.

We now discuss specific aspects of the LDBLL.

**The Latent Derivative Objective** Although the BLL’s derivative distribution can be represented in closed-form, it is a stochastic process rather than a weight distribution. As a result, its KL divergence to a prior process $\pi$ manifests as a functional KL, which is not a well-defined quantity when the prior does not share the same feature space. In contrast to a regular KL divergence between finite-dimensional probability distributions, a functional KL between stochastic processes requires the evaluation of an infinite-dimensional integral, which is intractable due to the lack of an infinite-dimensional Lebesgue measure [10]. However, it is possible to use a finite index set $T$ to estimate the otherwise intractable fKL because the fKL between a prior and posterior conditional Gaussian process is equal to the divergence at observations conditioned on $T$ [10].

$$D_{KL}(p(f) \mid\mid p(f \mid T)) = D_{KL}(p(f_T) \mid\mid p(f_T \mid T)). \quad (9)$$

While we could evaluate the divergence at the training data, i.e. $T = D$, to account for OOD prediction, we add some noise by defining $T = \{s_j \sim N(\cdot \mid x_j, \gamma I)\}^n_{j=1}$ and estimate the LD fKL as

$$\frac{1}{|T|} \sum_{s_j \in T} D_{KL}(\pi(z \mid s_j) \mid\mid p(z \mid s_j, D, \theta)). \quad (10)$$

The index set $T$ should ideally represent the true data distribution. Since this data distribution is typically unknown, we create index sets by sampling near the observed training data as a proxy. This is not necessarily the optimal sampling strategy, as this would depend on both the data distribution and task. However, we believe it balances staying within and outside the data distribution, and therefore should be a robust strategy across settings.

**Prior Specification** We choose the latent derivative prior $\pi$ as a Gaussian process with a mean function $\mu_\pi$.
Table 1: Means and standard errors of test metrics for different BNNs with leaky relu (LR) and tanh (TA) activations for nonlinear regression of gap (Cartpole, CO2, Sarcos, WAM) and standard (UCI) datasets.

(a) Log-Likelihood

<table>
<thead>
<tr>
<th>MODEL</th>
<th>GAP</th>
<th>CO2</th>
<th>SARCOS</th>
<th>WAM</th>
<th>BOSTON</th>
<th>CONCRETE</th>
<th>POWER</th>
<th>YACHT</th>
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(b) RMSE

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<th>WAM</th>
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<td>0.39</td>
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</tr>
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</table>

and covariance function \( \Sigma_x \).

\[
\pi(z \mid x) = GP(z \mid \mu_x(x), \Sigma_x(x)).
\] (11)

In practice, we set the prior to be constant, with \( \mu_x(x) = 0 \) and \( \Sigma_x(x) = I \) in whitened data space. The zero mean derivative prior is motivated by the zero mean weight prior. The derivative covariance is harder to specify. While a constant covariance may not be the optimal LD prior for a given task, from a practical perspective it is straightforward to specify, analogous to Gaussian weight priors used for BNNs. Domain knowledge (such as a physics model) could be used to define a more complex derivative prior process, which would combine the benefits of task-specific knowledge and black-box function approximation.

With the Bayesian last layer, it may appear that a LD prior ‘overdefines’ the BLL and that the two probabilistic treatments conflict. However, as the LD prior seeks to leverage the expressive feature space of the neural network, the universal approximation capability of the neural features should be capable of satisfying both the BLL likelihood and derivative prior. However, due to the construction of the model there are two constraints that can inform our choice of LD prior based on the weight prior. One is that as zero mean weight prior suggests a zero mean derivative prior, due to the linearity of Equation [3]. The other is that as \( \sigma^2 \rightarrow 0 \), \( \nabla [z \rightarrow 0 \) due to the weight posterior (defined in Equation (10)) in Equation (4).

In light of this second aspect, we found that scaling the LD prior with the aleatoric uncertainty \( \sigma^2 \) improved the prior specification and reduced underfitting in the nonlinear regression tasks. However, the fixed LD prior was beneficial for tasks requiring greater uncertainty quantification, such as active learning. While this scaling can be viewed as a form of empirical Bayes (EB), its limited application suggests better EB approaches may exist. For example, \( \mu_x \) would benefit from adapting to linear trends in the data, and \( \Sigma_x \) could be improved by adapting to the relative smoothness w.r.t. each input.
4 Experiments

We evaluated the LD prior on several tasks that require predictive uncertainty, namely nonlinear regression, active learning and Bayesian optimization, to verify that our proposed functional latent derivative prior improves the BLL in terms of adequate epistemic uncertainty in the absence of observed data. More detailed discussions and visualizations of all involved datasets can be found in Section I.

4.1 Nonlinear Regression

For the nonlinear regression benchmarks, we compare our LDBLL to the standard BLL and several other baselines: the nonparametric Gaussian process, a regularized network (MAP) and popular BNN approaches. These include mean-field variational inference (MFVI) [7], Monte Carlo dropout [25], ensembles [38] and stochastic weight averaging (SWAG) [44]. All regression problems involve real-world data, however, inspired by previous work based on in-between uncertainty [24], we distinguish between four novel ‘gap’ tasks, namely Cartpole, CO2, Sarcos and WAM, and ‘standard’ tasks from the popular UCI benchmark. Our goal is to show that the LDBLL improves the BLL significantly in terms of combating overconfidence during OOD prediction, which shall be demonstrated by the gap tasks, while maintaining competitive performance on the standard benchmarks. Due to the abundance of data, the Gaussian BLL backbone without Bayesian treatment of the observation noise was used for nonlinear regression.

CO2 The Mauna Loa atmospheric carbon dioxide dataset contains CO2 measurements over several decades [69]. To encode the periodicity without using specialized models, we augment the time input with sinusoidal features with an annual frequency. The gap region for testing considers central and edge portions.

Cartpole Here, telemetry is recorded from a Quanser cartpole system performing a swing-up maneuver. We use the dynamic state (position, velocity and acceleration) of the cart and pole for inverse dynamics modeling of the drive torque. The gap region is about the hanging position, where \( \theta < 45^\circ \), as depicted by Figure 11 in the Appendix.

Sarcos This dataset [79] contains the telemetry from a 7 DOF manipulator. It is used as a regression benchmark for inverse dynamics modeling, regressing the 21-dimensional state to a drive torque. In the central portion of the data, the robot’s pose induces a bias torque (likely due to gravity) in one of the upper motor drives. Therefore, modeling the inverse dynamics on this torque requires OOD prediction. Forecasting unseen aspects of dynamics from limited data represents a key challenge in MBRL for robotics.

WAM This dataset is also derived from a robotic manipulator, the cable-driven 4 DOF Barrett WAM. However, here the distribution shift is generated by demanding the same complex motion at different velocities. By training on a slower motion and evaluating the inverse dynamics model for data collected at a faster speed, the prediction considers the same trajectory but now with higher variance in the values of the state and input due to the larger accelerations at play.

UCI These datasets consists of several disparate regression problems that vary in size and dimension, and are a common benchmark for probabilistic nonlinear regression.

Flight Delay The flight delay dataset is a large-scale regression task of 700k datapoints used to demonstrate scalability [29]. While Bayesian methods are generally less useful for large datasets, as uncertainty should be minimal assuming no distribution shift, models should be able to scale adequately. We detail a batch method for training the BLL using a variational approximation, which aids the model in scaling to large datasets at the cost of non-exact inference during training. We describe this method in Section I and the results in Section I.

More details about the novel gap tasks are discussed in Section I.

Empirical results, displayed in Table 1 show that, in terms of the gap tasks, the LDBLL outperforms the standard BLL significantly in terms of test log-likelihood, which captures the adequacy of the ratio between goodness of fit (RMSE) and predicted uncertainty (entropy). This indicates the LD prior influences a better feature space for predictive uncertainty OOD. For standard regression, results were comparable, which makes sense as OOD uncertainty is not useful in this setting.

With respect to the baselines, the GP, MC dropout and ensembles performed better across gap and standard regression tasks. In fact, the GBLL performance was typically closer to the MAP model than the BNN, and the LD prior did not improve this performance enough to be deemed a competitive alternative. This could be due to capacity (GPs and ensembles have more parameters) and or a superior prior (e.g. the RBF kernel, the Bernoulli weight prior of MC dropout). Superior performance is also characterized by ‘underfitting’ on the training data (see the tables in Section I), sug-
Gesturing the LD prior is not regularizing enough during training. While the LD prior is interpretable due to the function space setting, it is not straightforward to assign values to when setting priors for a given task. As empirical Bayes would only tune the prior towards overfitting, it remains an open question how to design the LD prior to provide appropriate regularization for regression tasks.

4.2 Active Learning

Active learning [14] is the setting where a probabilistic model takes an active role in data acquisition, choosing points to optimize learning w.r.t. a utility measure. It is useful in domains such as system identification, where sampling data can be an expensive process. We use the Cartpole dataset introduced in Section 4.1 which contains a dynamical system performing a ‘swing-up’ control task. The (much smaller) swing-up portion is highly informative while the remaining samples from stabilization are generally redundant. Therefore, information-theoretic data acquisition offers a significant improvement over a random strategy. Section I.2 describes the experiment in detail and Figure 4 shows the results on a held out test set. In this experiment we also compare to a GP, which excels at uncertainty quantification under small datasets. The LDTBLL matches the GP in terms of RMSE and final LLH, however its predictions appear slightly overconfident during learning in comparison. Moreover, the LD prior evidently improves performance significantly on the standard BLL, which is miscalibrated with considerable variance. MFVI struggles to perform the task due to its lack cluster uncertainty quantification, which is evident from its relatively small predictive entropy on the test set. As a result, its selected data will likely be collected from uninformative regions and thus essentially random. This would explain its slow progress in both RMSE and LLH improvement.

4.3 Bayesian Optimization

Bayesian optimization (BO) [55, 73] is a sample-efficient black-box global optimization method. By constructing a Bayesian model of the objective, an uncertainty-derived utility function can be used to decide optimal function evaluations. Again, we compare to a GP, which is preferred for BO over BNNs. We performed BO on two tasks (Figure 5), chosen to highlight both the strengths and weaknesses of the LDBLL for BO. They are described in Section I.3.

Sinc in a Haystack This toy example is designed to demonstrate the utility of the LDBLL in optimizing high frequency functions, where the optima may be highly local. The function, \( f(x) = \text{sinc}(6(x - 1)) \), is challenging to optimize despite being smooth, as it requires large epistemic uncertainty to avoid suboptimal convergence. While all models demonstrate high variance in performance, the standard TBLL typically fails to achieve any improvement, whereas the LDTBLL is evidently superior. Its ‘maximum entropy’ nature translates to a powerful exploration strategy.

Hartmann6 This is a standard BO benchmark, with a six-dimensional state and six local minima. Figure 5 shows that the GP is vastly superior at this task, converging rapidly and consistently. This is due in part to the function’s smoothness combined with the smoothness assumption of the GP kernel. While the LDBLL converges faster than the BLL, indicating that the LD prior scales to higher dimensions, both converge to a similar suboptimal value compared to the GP. This suggests that either the LDBLL fails to capture the finer grained epistemic uncertainty required to fully converge, or that the specific BO optimizer used here benefits from the GP’s smoothness and is less suited to optimizing the BLL due to its increased roughness.
5 Related Work

Bayesian Neural Networks BNNs have existed since the 1980s as a means of both utilizing neural networks as statistical models \[43\] and for general regularization \[31\]. The early work of Neal \[50\] provided several key contributions, namely the insight that the limit of an infinite hidden layer neural network is a Gaussian process under certain conditions, and the use of Hamiltonian Monte Carlo for approximate inference. Despite the statistical elegance of MCMC training methods \[1\] and their advancements \[33, 12\], they are expensive to deploy and scale poorly with larger models. These shortcomings have motivated a focus on variational inference methods \[31, 59, 26\] for BNNs, including unbiased gradient estimation \[7\] and other advanced techniques \[76, 20, 84, 28\]. There is a large family of alternative approximate methods: Including the Laplace approximation \[43, 19, 66\], ensembles \[38, 53, 3, 57\], expectation propagation \[30\], Monte Carlo dropout \[25\], variational dropout \[86\], and a range of gradient-based approaches \[40, 44\]. The Bayesian last layer (also referred to as adaptive basis function and neural linear) model was introduced as a ‘marginalized neural network’ (MNN) \[39\] as a neural equivalent to sparse GPs. To mitigate feature overfitting, the MNN uses an ensemble of feature networks. BLLs have previously been applied to Bayesian optimization methods \[74\], bandit problems \[80, 65\], active learning \[60\], reinforcement learning \[52\] and regression \[51\], but there appears a lack of work on improving their general performance. Inference networks \[70\] are similar to fVI, but take a functional mirror-descent interpretation and incrementally fit the GP prior, enabling minibatch training. Prior Networks \[45\] use the neural network to parameterize a marginalized distribution, therefore directly predicting Dirichlet distributions for classification.

Gaussian Processes Beyond Neal’s infinite limit, there is a rich body of research on the intersection of GPs and NNs. The arcsine (or MLP) \[82\] and arccosine \[13\] covariance functions represent the kernel of an infinite single hidden layer network with erf and ReLU activations respectively. The manifold GP \[11\] uses a neural network to learn an intermediate feature space so that the covariance function performs better on non-smooth functions. Deep kernels \[83\] define closed-form kernels using neural network components for more expressive covariance functions that are able to incorporate inductive biases such as convolutional operators. Deep Gaussian processes \[15, 10, 67\] stack GPs to learn a hierarchical representation of intermediate latent variables to build sophisticated statistical models. Moreover, the Student-\(t\) process \[69\] is a Gaussian process with an inverse gamma / Wishart prior over the aleatoric uncertainty.

Functional Priors As Gaussian processes are exact distributions over functions, sparse GPs may be viewed as approximate inference over functions \[16\], minimizing the fKL from its exact posterior via inducing points. The functional variational BNN (fBNN) \[76\] uses the fKL to use explicit or implicit stochastic processes as functional priors. They use a GP trained on the data as a prior, which can be viewed as an elaborate form of empirical Bayes. While this prior improves the performance of the variational BNN compared to other methods, it is not evident when and to what extent improvement is made over the GP prior. The noise contrastive prior (NCP) \[27\] is a similar idea where the training data is perturbed by random noise to serve as a ‘data prior’ for a BNN, in order to increase uncertainty estimation OOD. While effective empirically, the data prior is again akin to empirical Bayes as the prior is defined by the data. Related work has also considered transforming the BNN weight prior into the prior of a GP \[22\]. The practice of combining kernels in GPs has been translated to BNN architectures and
activation functions, producing periodic and mixing phenomena in the network’s feature space for more expressive models [58]. Variational implicit priors [41] use variational inference to work with functional priors you can only sample from, e.g. simulators, which provides the flexibility of a broad range of complex processes to be adopted as priors.

6 Conclusion

We introduced the latent derivative prior, a novel functional prior for the Bayesian last layer which improves epistemic uncertainty by promoting feature diversity. This model has several attractive properties over weight space BNNs, namely explicit predictive distributions and an intuitive prior that directly enhances functional uncertainty. The LDBLL further demonstrates that, like GPs, linear Bayesian models can be sufficient for many problems if the underlying feature space is adequately expressive. We have shown through a suite of tasks that the LD objective significantly improves the uncertainty quantification of the BLL, such that the model is a viable parametric alternative to GPs for downstream tasks like active learning. The LD prior would be further improved by adequate specification for a given task or dataset. Using the notion of derivatives, this prior could provide a way of incorporating domain knowledge (i.e. from physics) into the model to improve performance over pure black box models. Moreover, the application of the BLL and LDBLL to multivariate prediction tasks such as model-based control and classification is an open avenue, in particular how the notion of predictive derivative uncertainty applies to classification.

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