Last Layer Marginal Likelihood for Invariance Learning

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Abstract

Data augmentation is often used to incorporate inductive biases into models. Traditionally, these are hand-crafted and tuned with cross validation. The Bayesian paradigm for model selection provides a path towards end-to-end learning of invariances using only the training data, by optimising the marginal likelihood. Computing the marginal likelihood is hard for neural networks, but success with tractable approaches that compute the marginal likelihood for the last layer only raises the question of whether this convenient approach might be employed for learning invariances. We show partial success on standard benchmarks, in the low-data regime and on a medical imaging dataset by designing a custom optimisation routine. Introducing a new lower bound to the marginal likelihood allows us to perform inference for a larger class of likelihood functions than before. On the other hand, we demonstrate failure modes on the CIFAR10 dataset, where the last layer approximation is not sufficient due to the increased complexity of our neural network. Our results indicate that once more sophisticated approximations become available the marginal likelihood is a promising approach for invariance learning in neural networks.

1 INTRODUCTION

Human learners generalise from example to category with seemingly little effort. Machine learning models

Figure 1: A non-invariant model M1 and its sign invariant (i.e. symmetric around $x=0$) counterpart M2. The non-invariant M1 has a better train MSE, but the invariant M2 has a better test MSE. The log marginal likelihood correctly identifies M2 as better.

aim to make accurate predictions on unseen data points based on finitely many examples.

This generalisation is enabled by inductive biases. In Steps toward Artificial Intelligence Marvin Minsky [1961] highlights the importance of invariance as an inductive bias: ‘One of the prime requirements of a good property is that it be invariant under the commonly encountered equivalence transformations. Thus for visual Pattern-Recognition we would usually want the object identification to be independent of uniform changes in size and position.’ In modern machine learning pipelines invariances are achieved through data augmentation. If we, for example, would like our neural network to be invariant with respect to rotation, we simply present it with rotated versions of the input data. Data augmentation schemes are almost always hand-crafted, based on assumptions and expert knowledge about the data, or found by cross-validation. We aim to learn invariances with backpropagation, to reduce the human intervention in the design of ML algorithms.

Learning invariances through gradients requires a suit-
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able loss function. Standard losses like negative log-likelihood or mean squared error solely measure how tightly we fit the training data. Good inductive biases (e.g., convolutions) constrain the expressiveness of a model, and therefore do not improve the fit on the training data. Thus, they can not be learned by minimising the training loss alone.

In Bayesian inference, this problem is known as model selection, and is commonly solved by using a different training objective: the marginal likelihood. For a model of data $y$, parametrised by weights $w$ and hyperparameters $\theta$ it is given by

$$ p(y|\theta) = \int p(y|w)p(w|\theta)dw . \tag{1} $$

As opposed to standard training losses, it correlates with generalisation, and thus provides a general way to select an inductive bias, independent of parameterisation (Williams and Rasmussen, 2006; Rasmussen and Ghahramani, 2001; MacKay, 2003). Van der Wilk et al. (2018) demonstrated that invariances can be learned by straightforward backpropagation using the marginal likelihood in Gaussian process (GP) models, where the marginal likelihood can be accurately approximated. Fig. [1] shows an invariant and a non-invariant GP; the invariant model has higher marginal likelihood as well as lower test mean squared error. Thus, the marginal likelihood correctly identifies invariance as a useful inductive bias.

Current GP models often lack predictive performance compared to their highly expressive neural network counterparts, hence applying this elegant principle to neural networks is attractive. The challenge is, however, that finding accurate and differentiable marginal likelihood approximations for neural networks is still an open problem. In this work we investigate a convenient short-cut: computing Bayesian quantities only in the last layer. This avoids difficulties of the marginal likelihood in the full network, and has already been shown helpful (Wilson et al., 2016a,b). Given the possible impact of invariance learning with the convenience of the last-layer approximation, it is important to investigate its potential. Our results provide a nuanced picture of this approach: there are situations where the last-layer approximation is sufficient, but others where it is not.

To provide these results, we

1. construct a deep neural network with a Bayesian last layer that incorporates invariance, based on invariant GPs (van der Wilk et al., 2018) and deep kernel learning (Wilson et al., 2016b),

2. overcome problems with the training implied by a straightforward combination of Van der Wilk et al. (2018) and Wilson et al. (2016b) via a new optimisation scheme, and a new variational bound that allows for non-Gaussian likelihoods,

3. investigate failure modes on more complex model architectures to show limitations of using the last-layer approximation for invariance learning.

2 RELATED WORK

Bayesian Deep Learning aims to provide principled uncertainty quantification for deep models. Exact computation for Bayesian deep models is intractable, so different approximations have been suggested. Variational strategies (e.g. Blundell et al., 2015) maximise the evidence lower bound (ELBO) to the marginal likelihood, thereby minimising the gap between approximate and true posteriors. To remain computationally feasible, approximations for Bayesian neural networks are often crude, and while weight posteriors are useful in practice, the marginal likelihood estimates are typically too imprecise for hyperparameter estimation (Blundell et al., 2015; Turner and Sahani, 2011). Hyperparameter estimation in deep GPs has achieved more success (Damianou and Lawrence, 2013; Dutordoir et al., 2020), but training deep GPs can be challenging. Some very recent works have shown initial promise in using the marginal likelihood for hyperparameter selection in Bayesian neural networks (Ober and Aitchison, 2020; Immer et al., 2021; Dutordoir et al., 2021). Instead of a Bayesian treatment of all weights using rough approximations, we follow a deep kernel learning approach, i.e. computing the marginal likelihood for the last layer only.

Deep Kernel Learning (DKL; Hinton and Salakhutdinov, 2007; Calandra et al., 2016; Bradshaw et al., 2017) replaces the last layer of a neural network with a GP, where marginal likelihood estimation is accurate (Burt et al., 2020; Wilson et al., 2016a,b) had significant success achieving improved uncertainty estimates. Their results indicate that such a neural network-GP hybrid is promising for invariance learning. Ober et al. (2021) identify difficulties with overfitting in DKL models, but also show mechanisms by which such overfitting is mitigated. We find similar issues and adapt the standard DKL training procedure to avoid them when learning invariance hyperparameters. We will discuss these issues in more depth as we describe our training procedure in Sec. [5].

Data Augmentation is used to incorporate invariances into deep learning models. Where good invariance assumptions are available a priori (e.g. for natural images) this improves generalisation performance and is ubiquitous in deep learning pipelines. Instead of relying on assumptions and hand-crafting, recent approaches
learn data augmentation schemes. Cubuk et al. (2019, 2020) and Ho et al. (2019) train on the validation data, and use reinforcement learning and evolutionary search respectively to find parameters. Zhou et al. (2021); Lorraine et al. (2020) compute losses on validation sets for learning invariance parameters, and estimate gradients w.r.t. them in outer loops. Similar to our work, Benton et al. (2020) learn data augmentations on training data end-to-end, by adding a regularisation term to the negative log-likelihood loss that encourages invariance. They argue that tuning this regularisation term via cross-validation can be avoided, since the loss function is relatively flat. Yet, the method relies on explicit regularisation, and thus on an understanding of the parameters in question. Our method is based on a Bayesian view of data augmentation as incorporating an invariance on the functions in the prior distribution (van der Wilk et al., 2018; Nabarro et al., 2021). This allows the marginal likelihood to be used as an objective for learning invariances. This has many advantages, such as allowing backpropagation from training data, automatic and principled regularisation, and parameterisation independence (see Sec. 3). This makes the marginal likelihood objective a promising avenue for future work, which may want to incorporate invariances whose parameterisations are non-interpretable.

3 BACKGROUND

3.1 Variational Gaussian processes

A Gaussian process (GP) (Williams and Rasmussen, 2006) is a distribution on functions with the property that any vector of function values \( f = (f(x_1), \ldots, f(x_N)) \) is Gaussian distributed. We assume zero mean functions and real valued vector inputs.

Inference in GP models with general likelihoods and big datasets can be done with variational approximations (Titsias, 2009; Hensman et al., 2015). The approximate posterior is constructed by conditioning the prior on \( M \) inducing variables \( u \in \mathbb{R}^M \), and specifying their marginal distribution with \( q(u) = \mathcal{N}(m, S) \) (for overviews see Bui et al., 2017; van der Wilk et al., 2020). This results in a variational predictive distribution:

\[
q(f(x^*)) = \mathcal{N}(\alpha(x^*)^\top m, \alpha(x^*)^\top (K_{zz} - S) \alpha(x^*)),
\]

where \( z \in \mathbb{R}^{M \times d} \) are inducing inputs, \( K_{zz} \) is the matrix with entries \( k(z_i, z_j) \), \( \alpha(x^*) = K_z^{-1} k(z, x^*) \), and \( k \) is the chosen covariance function.

Variational inference (VI) selects an approximation by minimising the KL divergence of the approximation to the true posterior with respect to the variational parameters \( z, m, S \). This is done by maximising a lower bound to the marginal likelihood (the “evidence”), which has the KL divergence as its gap (Matthews et al., 2016). The resulting evidence lower bound (ELBO) is

\[
\log p(y) \geq \mathcal{L} = \sum_{n=1}^{N} \mathbb{E}_{q(f(x_n))} \left[ \log p(y_n | f(x_n)) \right] - \text{KL}[q(u) | | p(u)].
\]

In exact GPs, (kernel) hyperparameters are found by maximising the log marginal likelihood \( \log p(y) \) (Williams and Rasmussen, 2006). For our models of interest, the exact marginal likelihood is intractable. We use the ELBO as a surrogate. This results in an approximate inference procedure that maximises the ELBO with respect to both the variational parameters and the hyperparameters. Optimising the variational parameters improves the quality of the posterior approximation, and tightens the bound to the marginal likelihood. Optimising the hyperparameters hopefully improves the model, but the slack in the ELBO can lead to worse hyperparameter selection (Turner and Sahami, 2011).

3.2 Invariant Gaussian Processes

A function \( f : \mathcal{X} \to \mathcal{Y} \) is invariant to a transformation \( t : \mathcal{X} \to \mathcal{X} \) if \( f(x) = f(t(x)) \), \( \forall x \in \mathcal{X} \), \( \forall t \in T \). I.e., an invariant function will have the same output for a certain range of transformed inputs known as the orbit. A straightforward way to construct invariant functions is to simply average a function over the orbit (Kondor, 2008; Ginsbourger et al., 2012; 2013). We consider a similar construction where we average a function over a data augmentation distribution, which results in an approximately invariant function where \( f(x) = f(t(x)) \) (van der Wilk et al., 2018; Dao et al., 2019). Augmented data samples \( x_a \) are obtained by applying random transformations \( t \) to an input, \( x_a = t(x) \), leading to the distribution \( p(x_a | x) \).

That is, an approximately invariant function \( f \) can be constructed from any non-invariant \( g \) as

\[
f(x) = \sum_{t \in T} g(t(x)), \quad \text{or} \quad f(x) = \int g(x_a)p(x_a | x)dx_a.
\]

Van der Wilk et al. (2018) exploit this construction to build a GP with continuously adjustable invariances. They place a GP prior on \( g \sim \mathcal{GP}(0, k_g(\cdot, \cdot)) \), and since Gaussians are closed under summations, \( f \) is a GP too. By construction \( f \) is invariant to the augmentation distribution \( p(x_a | \cdot) \) and its kernel is given by

\[
k_f(x, x') = \int k_g(x_a, x_a') p(x_a | x)p(x_a' | x')dx_adx_a'.
\]

Non-trivial \( p(x_a | x) \) densities present a problem for standard VI, as the kernel evaluations in eq. 2 become
intractable. This is solved by making the inducing variables observations of \( g \) rather than the usual \( f \). This ensures that \( K_{zz} \) is tractable, as it only requires evaluations of \( k_f \), which makes the KL divergence tractable. When the likelihood is Gaussian, it additionally provides a way to tackle the expected log likelihood:

\[
\mathbb{E}_q(f(x)) \log \mathcal{N}(y; f(x), \sigma^2) = \text{const} - \frac{(y - \mu)^2 + \tau}{2\sigma^2} \tag{6}
\]

where \( \mu, \tau \) are the mean and variance in (2). Only unbiased estimates of \( \mu, \mu^2 \) and \( \tau \) are needed for an unbiased estimate of the ELBO. These can be obtained from simple Monte Carlo estimates of \( k_f \) (5), and \( k(z, x) \).

### 3.3 Parameterising learnable invariances

The invariance of the GP in (5) is learned by adjusting the augmentation distribution. We parameterise the distribution and treat its parameters as kernel hyperparameters. We learn these by maximising the ELBO. As done in similar work (Benton et al., 2020; van der Wilk et al., 2018), we consider affine transformations. Our affine transformations are controlled by \( \phi = (\alpha, s^2, \delta, \phi_x, \phi_y) \), which describes rotation, scale, shearing and horizontal and vertical translation. We parameterise a family of augmentation distributions by specifying uniform ranges with \( \phi_{\min}, \phi_{\max} \in \mathbb{R}^7 \) that are to be applied to the input image. Different ranges that are learned on \( \phi_{\min}, \phi_{\max} \) correspond to different invariances in \( f(\cdot) \). For example, learning \( \phi_{\min/\max} = (\pm \pi, 0, 0, 0, 0, 0) \) corresponds to full rotational invariance (sampling any angle between \(-\pi \) and \( \pi \)) but no scaling, shearing or translations.

We sample from the resulting \( p(x_a|x, \phi_{\max}, \phi_{\min}) \) (we will write \( p(x_a|x, \phi) \) for brevity) by 1) sampling the parameters for a transformation from a uniform distribution, 2) generating a transformed coordinate grid, and 3) interpolating the image:

\[
x_a = t_\nu(x), \quad \nu \sim U(-\phi_{\min}, \phi_{\max}). \tag{7}
\]

Since transforming \( t_\nu(x) \) is differentiable, this procedure is reparameterisable w.r.t. \( \phi_{\max}, \phi_{\min} \) via \( \nu = \phi_{\min} + (\phi_{\max} - \phi_{\min}) \varepsilon, \varepsilon \sim U(0, 1) \). Straightforward automatic differentiation of the unbiased ELBO estimator described in the previous section provides the required gradients.

In summary, we learn \( \phi_{\min/\max} \) by maximising the ELBO, so the transformations and their magnitudes are learned based on the specific training set. Different invariances will be learned for different training sets, and different magnitudes will be learned for different training data. The next sections show how these principles have potential in neural network models, beyond the single layer GPs of (van der Wilk et al., 2018).

### Algorithm 1: InvDKGP forward pass

1. Draw \( S \) samples from the augmentation distribution \( x_a^i \sim p(x_a|x, \phi), \ i = 1...S \).
2. Pass the \( x_a^i \) through the neural net \( h_w \).
3. Map extracted features using the non-inv. \( g \).
4. Aggregate samples to obtain inv. \( f(x) \) by
   (i) using the unbiased estimators from Sec. 3.2 in the Gaussian case, or
   (ii) averaging predictions \( g(h_w(x_a^i))) \), \( i = 1, ..., S \) directly in the Softmax case, see (16).

### 4 MODEL

As discussed in Sec. 1, we aim to learn neural network (NN) invariances through backpropagation, in the same way as is possible for single-layer GPs. Since finding high-quality approximations to the marginal likelihood of a NN is an ongoing research problem, we investigate whether a simpler deep kernel approach is sufficient. This uses a GP as the last layer of a NN, and takes advantage of accurate marginal likelihood approximations for the GP last layer. Success with such a simple method would significantly help automatic adaptation of data augmentation in neural network models. We hypothesise that the last layer approximation is sufficient, since data augmentation influences predictions only in the last layer (in the sense that one can construct an invariant function \( f \) from an arbitrary non-invariant \( g \) by summing in the last layer, eq. 4). See Fig. 2 for a graphical representation and Algorithm 1 for forward pass computations.

**Deep Kernels** take advantage of covariance functions being closed under transformations of their input. That is, if \( k_g(\cdot, \cdot) \) is a covariance function on \( \mathbb{R}^D \times \mathbb{R}^D \), then \( k_g(h_w(\cdot), h_w(\cdot)) \) is a covariance function on \( \mathbb{R}^d \times \mathbb{R}^d \) for mappings \( h_w : \mathbb{R}^d \rightarrow \mathbb{R}^D \). In our case, \( h_w \) is a NN parametrised by weights \( w \), and hence \( w \) are viewed as hyperparameters of the kernel. The GP prior becomes

\[
p(g) = \mathcal{GP}(0, k_g(h_w(\cdot), h_w(\cdot))). \tag{8}
\]

The idea is to learn \( w \) along with the kernel hyperparameters. Importantly, this model remains a GP and so the inference described in Sec. 3 applies.

**Our invariant model** combines the flexibility of a NN \( h_w(\cdot) \) with a GP \( g \) in the last layer, while ensuring
overall invariance using the construction from (4):

$$f(x) = \int g(h_w(x_a))p(x_a|x, \phi)dx_a.$$  \hfill (9)

Thus, combining (5) and (9), $f$ is an invariant GP with a deep kernel given as

$$k_f(x, x') = \int k_g(h_w(x_a), h_w(x'_a))$$

$$p(x_a|x, \phi)p(x'_a|x', \phi)dx_adx'_a.$$ \hfill (10)

The model is trained to fit observations $y$ through the likelihood function $p(y|f(x))$, where we assume observations $y_i$ are independent conditioned on the marginals $f(x_i)$.

Initially, we investigate training a model by simply combining the invariant GP training objective for Gaussian likelihoods (van der Wilk et al., 2018) with standard deep kernel learning (Wilson et al., 2016a,b). However, as we will discuss, several issues prevent these training procedures from working. In following sections we investigate why, provide solutions, and introduce a new ELBO that is suitable for more general likelihoods which improves training behaviour. We refer to our model as the Invariant Deep Kernel GP (InvDKGP). An implementation can be found at https://github.com/polaschwoebel/InvDKGP.

5 DESIGNING A TRAINING SCHEME

The promise of deep kernel learning as presented by Wilson et al. (2016a,b) lies in training the NN and GP hyperparameters jointly, using the marginal likelihood as for standard GPs.\footnote{Given that this quantity is difficult to approximate, we verify experimentally that we indeed need it and cannot use a simple NN with max-likelihood (see Appendix).} However, prior works have noted shortcomings of this approach (Ober et al., 2021; Bradshaw et al., 2017; van Amersfoort et al., 2021): the DKL marginal likelihood correctly penalises complexity for the last layer only, while the NN hyperparameters can still overfit. In our setting, i.e. when trying to combine deep kernel learning with invariance learning, joint training produces overfit weights which results in simplistic features with little intra-class variation. In particular, all training points from the same class are mapped to very similar activations, independent of orientation. This causes a loss of signal for the invariance parameters (see Fig. 3).
Coordinate ascent training fixes this problem. We pre-train the NN using negative log-likelihood loss. Then, we replace the fully connected last layer with an invariant GP. The marginal likelihood is a good objective given fixed weights (we obtain a GP on transformed inputs), so we fix the NN weights. However, some adaptation of the NN to the transformed inputs is beneficial. We thus continue training by alternating between updating the NN, and the GP variational parameters and orbit parameters, hereby successfully learning invariances. (See Fig. 7 and 8: flat parts of the training curves indicate NN training where all kernel hyperparameters, including invariances, remain fixed. When to toggle between the GP and NN training phase is determined using validation data.)

Choosing an invariance parameterisation is simple with our method. Other invariance learning approaches, e.g. Benton et al. (2020) and Schwöbel et al. (2020) rely on explicitly regularising augmentation parameters to be large, and thus require interpretability of their parameters. The marginal likelihood objective is independent of parameterisation. To illustrate this we compare parameterising the range of angles by the angle in radians $\alpha$ and by its reciprocal $\xi = \frac{1}{\alpha}$. In the rotMNIST example (see Fig. 4) large invariances are needed. This corresponds to large $\alpha$ or small $\xi$ – our method obtains this in both parameterisations. In contrast, explicitly regularising invariance parameters to be large would fail for $\xi$. We wish to stress that generating the orbit distributions is not restricted to affine image transformation and parameterisation independence will be more important as more complicated, non-interpretable invariances are considered.

The Gaussian likelihood is chosen by Van der Wilk et al. (2018) due to its closed-form ELBO. For classification problems, this is a model misspecification. The penalty for not fitting the correct label value becomes large and we can therefore overfit the training data. To alleviate this problem, we fix likelihood and kernel variance (see Fig. 5). The fixed values were determined by trying out a handful candidates – this was sufficient to make invariance learning work. To remove this manual tuning, we will derive an ELBO that works with likelihoods like Softmax in Sec 6.

5.1 MNIST subsets – the low data regime

Having developed a successful training scheme we evaluate it on MNIST subsets. The generalisation problem is particularly difficult when training data is scarce. Inductive biases are especially important and usually parameter-rich neural networks rely on heavy data augmentation when applied to smaller datasets. We train on different subsets of MNIST (LeCun et al.). InvDKGP outperform both NNs and non-invariant deep kernel GPs. The margin is larger the smaller the training set — with only 1250 training examples we can nearly match the performance of a NN trained on full MNIST (Fig. 6). We conclude it is possible to learn useful invariances even from small data (see Fig. 7). This data efficiency is desirable since models trained on small datasets benefit crucially from augmentation.
6 CORRECTING MODEL MISSPECIFICATION

The key observation for inference under the Gaussian likelihood was the unbiasedness of the estimators. In this section, we introduce a controlled bias to allow for easy inference in a wide class of likelihoods. In the limit of infinite sampling, the bias disappears and the invariance does not add additional approximation error.

Recall that $f(x)$ constructed in (4) is intractable but can be estimated by Monte Carlo sampling

$$
\hat{f}(x) := \frac{1}{S_0} \sum_{i=1}^{S_0} g(x_a^i),
$$

where $x_a^i \sim p(x_a|x, \phi)$. Notice,

$$
f(x) = \mathbb{E}_{\prod_{i=1}^{S_0} p(x_a^i|x, \phi)} \left[ \hat{f}(x) \right] =: \mathbb{E} \left[ \hat{f}(x) \right],
$$

where $\prod_{i=1}^{S_0} p(x_a^i|x, \phi)$ is the product density over $S_0$ orbit densities. We remark that $f$ is deterministic in $x$ but stochastic in $g$, which is a GP. Thus, we can write

$$
\mathbb{E}_{q(f(x))} [\log p(y|f(x))] = \mathbb{E}_{q(g)} [\log p(y|\mathbb{E}[f(x)])]
$$

$$
= \mathbb{E}_{q(g)} \left[ \log p(y|\mathbb{E}\hat{f}(x)) \right]
$$

$$
\geq \mathbb{E}_{q(g)} \left[ \mathbb{E} \left[ \log p(y|\hat{f}(x)) \right] \right].
$$

Equality holds above when $\text{Var}(\hat{f}(x)) = 0$, i.e. the bound becomes tighter as $S_0$ increases (see also Burda et al., 2016). Hence aggressive sampling recovers accurate VI. The right-hand side of (15) can now, without additional bias, be estimated by

$$
\frac{1}{S_g} \sum_{k=1}^{S_g} \frac{1}{S_A} \sum_{j=1}^{S_A} \log p \left( y \mid \frac{1}{S_0} \sum_{i=1}^{S_0} g_k(x_a^{ji}) \right).
$$

Since extensive sampling is required to keep the bound above tight, it is important to do this efficiently. From a GP perspective this is handled with little effort by sampling the approximate posteriors $q(g)$ using Matheron’s rule (Wilson et al., 2020). Thus, sampling $S_g$ GPs is cheap compared to sampling from the orbit. $S_A$ denotes the number of $f$ samples, this can be fixed to 1 as long as $S_0$ is large.

Summarising, we have shown how we can infer through the marginal likelihood, for the wide class of log-concave likelihoods, by maximising the stochastic ELBO:

$$
\mathcal{L} = \frac{1}{S_g} \sum_{k=1}^{S_g} \frac{1}{S_A} \sum_{j=1}^{S_A} \log p(y) \left[ \frac{1}{S_0} \sum_{i=1}^{S_0} g_k(h_w(x_a^{ji})) \right] - \text{KL}[q(u)||p(u)],
$$

with $x_a^{ij} \sim p(x_a|x, \phi)$.

The benefits of our new sample based bound are three-fold: It broadens model specification, avoids hand-picking and fixing the artificial Gaussian likelihood variance, and doubles training speed.
Table 1: Test accuracies on rotated MNIST. Invariant models outperform non-invariant counterpart. So do deep kernels contra shallow ones. The invariant deep kernel GPs perform best, outperforming state-of-the-art of 0.989 for learned invariance [Benton et al., 2020].

| Model                          | Likelihood       | Test acc. |
|-------------------------------|------------------|-----------|
| M1   NN                        | Softmax          | 0.9433    |
| M2   Non-inv. Shallow GP       | Gaussian         | 0.8357    |
| M3   Non-inv. Shallow. GP      | Softmax          | 0.7918    |
| M4   Inv. Shallow GP           | Gaussian         | 0.9316    |
| M5   Inv. Shallow. GP          | Softmax          | 0.9316    |
| M6   Non-inv. Deep Kernel GP   | Gaussian         | 0.9387    |
| M7   Non-inv. Deep Kernel GP   | Softmax          | 0.9351    |
| M8   Inv. Deep Kernel GP       | Gaussian         | 0.8996    |
| M9   Inv. Deep Kernel GP       | Softmax          | 0.9867    |

Table 2: PCam results. InvDKGP performs best.

| Model                           | Test acc. |
|---------------------------------|-----------|
| NN                              | 0.7905    |
| Deep Kernel GP + no inv.        | 0.8018    |
| NN + small inv.                 | 0.7420    |
| Deep Kernel GP + small inv.     | 0.8115    |
| Deep Kernel GP + learned inv.   | 0.8171    |

6.1 Rotated MNIST

The rotated MNIST dataset was generated from the original MNIST dataset by randomly rotating the images of hand-written digits between 0 and 2π radians. It consists of a training set of 12,000 images along with 50,000 images for testing. We pretrain the neural network from Sec. 5.1 on rotated MNIST (Table 1: M1) and proceed as outlined in Sec. 5. As discussed in Sec. 3, we do not have guarantees that the ELBO acts as a good model selector for the neural network hyperparameters. We thus use a validation set (3000 of the 12000 training points) to find hyperparameters for the NN updates. Once a good training setting is found we re-train on the entire training set (see Appendix for settings). Fig. 8 shows the learned invariances (we use the full φ parameterisation but only plot rotation and translation for brevity). Both Gaussian and Softmax models learn to be rotation-invariant close the full 2π rotations present in the data. Table 1 contains test accuracies. Deep kernel GPs outperform their shallow counterparts by large margins (differences in test accuracy of ≥ 10 percent points). The same is true for invariant compared to non-invariant models (≥ 3 percent points). While both likelihoods achieve similar test accuracies, we observe a 2.3× speedup per iteration, Gaussian + sample bound: 1.32 sec./iter., Softmax + sample bound: 1.13 sec./iter. All runs are executed on 12 GB Nvidia Titan X/Xp GPUs.

6.2 PatchCamelyon

The PatchCamelyon (PCam, CC0 License, Veeling et al. (2018)) dataset consists of histopathology scans of lymph nodes measuring 96 × 96 × 3 pixels. Labels indicate whether the centre patch contains tumor pixels. Veeling et al. (2018) improve test performance from 0.876 to 0.898 by using a NN which is invariant to (hard-coded) 90° rotations of the input. Such discrete, non-differentiable augmentations are not compatible with our backprop-based method, so we instead use continuously sampled rotations (a special case of the transformations described in Sec. 3.3 with φ = α = α min = −α max). This, contrary to Veeling et al. (2018)’s approach, introduces the need for padding and interpolation (see Fig. 9 left), effectively changing the data distribution. We thus apply small rotations as a preprocessing step (π/10 radians, ‘small inv.’ in Table 2). This lowers performance for a NN alone, i.e. when pre-training. The invariant models counterbalance this performance drop, and the learned invariances produce the best results in our experiments; however, they remain subpar to Veeling et al. (2018). This is due to the limitation to differentiable transformations, as well as our simpler NN (see Appendix). We highlight that our task is fundamentally different: instead of hard-coding invariances we learn those during optimisation.

7 EXPLORING LIMITATIONS

Rotated MNIST and PCAM are relatively simple datasets that can be modelled using small NNs. To investigate whether our approach can be used on more complex datasets, we turn to CIFAR-10 (Krizhevsky 2009), which is usually trained with larger models and data augmentation. Unfortunately, we found that we were unable to learn invariances for CIFAR-10.
To understand why, we designed a simple experiment. We first pretrain ResNet-18-based \cite{He2016} networks with different levels \( \nu \) of invariance transformations (see the Appendix for a definition of \( \nu \)). We then train sparse GP regression (SGPR; \cite{Titsias2009}) models on an augmented training set created by propagating ten points sampled from the augmentation distribution through these neural networks. The samples are generated at different levels of invariance \( \nu \), not necessarily matching the levels of the pretrained NNs. We plot the results in Fig. 10: when the network is trained at a small invariance level \( \epsilon \), the performance of the SGPR model is highest at an invariance level of 0.01, and rapidly drops off for larger invariances (note the logarithmic x scale). We see a similar result for the network trained at a level of 0.1. Finally, when the network is trained at the same level that the orbit points for the SGPR model are sampled at (‘adapted’), we see that added invariance helps the accuracy, with no steep drop off in accuracy for larger invariances. Therefore, adding invariance does help, but only when the network has already been adapted to that invariance. Currently, in our method this coadaptation is prevented by the current need for coordinate ascent training (Sec. 5).

This experiment indicates that for datasets requiring larger neural networks, we are in a difficult position. We need to adapt the feature extractor jointly with the invariances. However, this approach leads to pathologies as the neural network parameters are not protected from overfitting \cite{Ober2021}, which we previously mitigated with coordinate ascent. Therefore, relying on the marginal likelihood to learn invariances with a large feature extractor can easily lead to unwanted behavior – this behavior prevents us from learning these invariances as easily as the marginal likelihood promises. We believe that ongoing research in Bayesian deep learning will alleviate this problem. Bayesian neural networks with methods for marginalising over lower layers too, thus protecting them against overfitting, will render our approach more easily applicable. Such advances will allow us to learn invariances more easily and on more complex tasks than we did for the MNIST and PCAM datasets.

8 CONCLUSION

Neural networks depend on good inductive biases in order to generalise well. Practitioners usually – successfully – handcraft inductive biases, but the idea of learning them from data is appealing. Might we automate the modelling pipeline, moving from handcrafted models to data driven models; much like we replaced hand-crafted features with learned features in deep neural networks? This work proposes one step in this direction. Inspired by Bayesian model selection we employ the marginal likelihood for learning inductive biases. We avoid the intractability of the marginal likelihood for neural networks by using Deep Kernel Learning. This enables us to leverage previous work on invariance learning in GPs for learning data augmentation in neural networks. We learn useful invariances and improve performance, but encounter challenges when optimising our models. We introduce a new sampling-based bound to the ELBO allowing for inference for the Softmax likelihood, the natural choice for classification tasks, hereby alleviating some of the optimisation difficulties. Others we identify as fundamental limitations of the Bayesian last layer approach.

Societal Impact: This work is situated within basic research in probabilistic ML and, as such, bears all the risks of automation itself: harmful redistribution of wealth to those with access to compute resources and data, loss of jobs, and the environmental impact of such technologies. In fact, our model is more computationally heavy than a standard neural network with hand-tuned data augmentation. However, in the long term, automatic model selection has the potential to reduce the need for hyperparameter tuning, which usually dramatically exceeds the resources needed for training the final model.

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J. van Amersfoort, L. Smith, A. Jesson, O. Key, and Y. Gal. On feature collapse and deep kernel learning for single forward pass uncertainty. *arXiv preprint arXiv:2102.11409*, 2021.
A  IS THE MARGINAL LIKELIHOOD NECESSARY?

Sec. 1 motivated the marginal likelihood for invariance learning. Given that this loss function is notoriously difficult to evaluate, we verify experimentally that using it is indeed necessary, i.e. that the standard maximum likelihood loss is insufficient. Fig. 11 shows invariances learned on rotated MNIST (rotMNIST, see Sec. 6.1 for a description of the dataset) by using a neural network with maximum likelihood loss for two initialisations (blue, green). They collapse as suggested by the theory. The marginal likelihood solution (red) instead identifies appropriate invariances.

![Image of Figure 11](image.png)

Figure 11: Max. likelihood (green, blue, collapsing) and marg. likelihood (red, useful) invariances.

B  EXPERIMENTAL DETAILS

Exploiting the ideas from Sec. 5, we start by training convolutional neural networks (CNNs, see below for architecture details). After pre-training the CNN, we replace the last fully connected layer with a GP and continue training. In the non-invariant case we train all parameters jointly from here. When learning invariances, we iterate between updating the GP variational- and hyperparameters, and the neural network weights.

B.1 MNIST variations

We here summarise the training setups for the experiments on MNIST variations, i.e. MNIST subsets (Sec. 5.1) and rotated MNIST (Sec. 6.1). We start by outlining the shared neural network architecture and will then list the hyperparameter settings for MNIST and rotMNIST, respectively.

The CNN architecture used in the (rot)MNIST experiments is depicted in Table 3. For rotated MNIST, we train the model for 200 epochs with the Adam optimiser (default parameters). For the MNIST subsets, we train for 60k iterations which corresponds to 200 epochs for the full dataset and respectively more epochs for smaller subsets. The remaining parameters are the same in all experiments: batch size 200, learning rate 0.001, no weight decay, other regularisation or data augmentation. In the pre-training phase we minimise negative log-likelihood, for updates during coordinate ascent we use the ELBO as a loss-function.

Hyperparameter initialisation for the MNIST subset experiments are lengthscale 10, likelihood variance 0.05, kernel variance 1 (fixed likelihood and kernel variance for the invariant model, see Sec. 5.1), posterior variance 0.01. We use 1200 inducing points which we initialise by first passing the images through the neural
Table 3: Neural network architecture for MNIST variations. After pre-training, the last fully connected layer (below dashed line) is replaced with a GP layer for the deep kernel models.

| Layer            | Specifications                                      |
|------------------|----------------------------------------------------|
| Convolution      | filters=20, kernel size=(5, 5), padding=same, activation=ReLU |
| Max pooling      | pool size=(2, 2), stride=2                          |
| Convolution      | filters=50, kernel size=(5, 5), padding=same, activation=ReLU |
| Max pooling      | pool size=(2, 2), stride=2                          |
| Fully connected  | neurons=500, activation=ReLU                        |
| Fully connected  | neurons=50, activation=ReLU                         |
| Fully connected  | neurons=10, activation=Softmax                      |

network, then using the ‘greedy variance’ method [Burt et al. 2020] on the extracted features. For the smallest dataset MNIST312 we use 312 inducing points only. The batch size is 200 and we choose learning rate 0.001 for the Adam optimiser. For the invariant models, the orbit size is 120 and affine parameters are initialised at \( \phi_{\text{min}} = \phi_{\text{max}} = 0.02 \), i.e. we initialise with a small invariance. Without this initialisation we encountered occasional numerical instabilities (Cholesky errors) on the small dataset runs. During coordinate ascent (InvDKGP models) we toggle between training GP and CNN after 25k steps.

**Hyperparameter initialisation for the rotMNIST experiment** as follows: For all models, we initialise kernel variance 1 (fixed at 1 for M9, see Sec. 5.1) and posterior variance 0.01. We use 1200 inducing points. For the invariant models, the orbit size is 120 and affine parameters are initialised at \( \phi_{\text{min}} = \phi_{\text{max}} = 0 \), i.e. invariances are learned from scratch. When using coordinate ascent (InvDKGP models, M8 & M9) we toggle between training GP and CNN after 30k steps. We train different models for a different number of iterations, all until the ELBO has roughly converged. Batch size 200 is used for all models. The remaining initialisations differ between models and are summarised in Table 4.

| Model     | Lengthsc. | Lik. var. | LR (decay) |
|-----------|-----------|-----------|------------|
| M2        | 10        | 0.02      | 0.001      |
| M3        | 10        | -         | 0.001      |
| M4        | 10        | 0.05      | 0.001      |
| M5        | 10        | -         | 0.001      |
| M6        | 10        | 0.05      | 0.001      |
| M7        | 20        | -         | 0.001      |
| M8        | 50        | 0.05 (F)  | 0.003 (steps / cyclic) |
| M9        | 9         | -         | 0.003 / 0.0003 (s / c) |

Table 4: Training settings for rotMNIST models: Kernel lengthscale and likelihood variance initialisations (‘F’ indicates a fixed likelihood variance, see Sec. 5.1). The learning rate column (‘LR’) also indicates whether the learning rate was decayed in the GP/CNN update phases of coordinate ascent. For the ‘steps’(s) decay, we divide by 10 after 50% and again 75% of iterations, for the ‘cyclic’(c) decay, learning rates are: [LR/100, LR/10, LR, LR/10, LR/100]. These training hyperparameters are determined using a validation set (see Sec. 6.1).

**B.2 PCam**

The CNN architecture is a VGG-like convolutional neural network described in Table 5. The model is trained for 5 epochs using the Adam optimiser with batch size 64. We use learning rate 0.001 which we divide by 10 after 50% and again 75% of training iterations. In the fully connected block we use dropout with 50% probability when pre-training. Dropout is disabled when training the deep kernel models.

**Hyperparameters** for the deep kernel GP experiments on PCam are: lengthscale 10 (1 for the learned invariance model), kernel variance 1, posterior variance 0.01. We use 750 inducing points which we initialise as in the previous experiments. The batch size is 32. For PCAm we use coordinate ascent for all models since

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\[^{7}\text{We closely follow https://geertlitjens.nl/post/getting-started-with-camelyon/}\]
Layer Specifications

| Layer       | Specifications                          |
|-------------|----------------------------------------|
| Convolution | filters=16, kernel size=(3, 3), padding=valid, activation=ReLU |
| Convolution | filters=16, kernel size=(3, 3), padding=valid, activation=ReLU |
| Max Pooling | pool size=(2, 2), strides=2             |
| Convolution | filters=32, kernel size=(3, 3), padding=valid, activation=ReLU |
| Max Pooling | pool size=(2, 2), stride=2              |
| Convolution | filters=64, kernel size=(3, 3), padding=valid, activation=ReLU |
| Max Pooling | pool size=(2, 2), stride=2              |
| Fully Connected | neurons=256, activation=ReLU           |
| Dropout     | probability=0.5                         |
| Fully Connected | neurons=50, activation=None            |
| Dropout     | probability=0.5                         |
| Fully connected | neurons=2, activation=Softmax        |

Table 5: Neural network architecture for PCAM. After pre-training, the last fully connected layer (below dashed line) is replaced with a GP layer for the deep kernel models and dropout is disabled.

this improves training stability. Learning rates are 0.001 for the GP update steps and 0.0001 for the CNN updates, no LR decay. We toggle between the two coordinate ascent phases after 50k and 75k iterations in the non-invariant and invariant case, respectively. For the invariant models the orbit size is 20 and we initialise the rotation invariance with φ\text{min/max} = α\text{min/max} = ±π/10.

### B.3 CIFAR-10

Throughout our experiments, we train on a subset of 45,000 points from the full CIFAR-10 (Krizhevsky, 2009) training set and report results on the remaining 5,000 points, as a validation set.

The model we use is a sparse GP regression (SGPR; Titsias (2009)) model with a sum kernel corresponding to a Monte Carlo estimate of the kernel of Eq. 5 using an automatic relevance determination (ARD) squared exponential kernel as a base kernel. We achieve this by sampling 10 points from the full orbit for each data point, and propagating the points through the pretrained feature extractor. For the feature extractor, we choose a ReLU ResNet-18 architecture (He et al., 2016) with an output dimension of 50, using the post-ReLU features. Therefore, for our training set, we end up with a set of $45,000 \times 10 \times 50$ datapoints, where we sum over the 10 orbit samples.

Hyperparameters were chosen as follows. We pretrain the ResNet-18 by adding an additional fully-connected layer with softmax activations. We train for 160 epochs with a batch size of 100 and the Adam optimizer (Kingma and Ba, 2015), starting with a learning rate of 0.001, which we step down by a factor of 10 at epochs 80 and 120. We train the network without weight decay. The SGPR model was subsequently trained for a maximum of 1000 steps, using the Scipy optimizer provided in GPflow (Matthews et al., 2017). During training, we initially set the jitter to 1e-6, which we increased by a factor of ten if the Cholesky decomposition failed. For the SGPR model, we use 1000 inducing points, initialised as above. We found empirically that the likelihood variance did not have a significant impact on the results; we therefore fixed it to 0.01. Recalling that \( \phi = (\alpha, s^x, s^y, p^x, p^y, t^x, t^y) \), we parameterize the transformation by considering the “transformation level” \( \nu \) such that

\[
\phi_{\text{max}} = (0, 1, 1, 0, 0, 0, 0, 0, 0, 0) + \nu \times (\pi, 1, 1, 1, 1, 1),
\]

\[
\phi_{\text{min}} = (0, 1, 1, 0, 0, 0, 0, 0, 0, 0) - \nu \times (\pi, 1, 1, 1, 1, 1).
\]

For the “\( \epsilon \)” setting of the transformation level, we assign \( \nu = 0.01 \). We chose a non-zero value to ensure that any reduction in performance would be due to a different value of \( \nu \), and not because of the lack of presence of the image interpolator in the pretraining (see Sec. 6.2).