Learning to Explore for Stochastic Gradient MCMC

SeungHyun Kim *1 Seohyeon Jung *2 Seonghyeon Kim 1 Juho Lee 1 3

Abstract

Bayesian Neural Networks (BNNs) with high-dimensional parameters pose a challenge for posterior inference due to the multi-modality of the posterior distributions. Stochastic Gradient Markov Chain Monte Carlo (SGMCMC) with cyclical learning rate scheduling is a promising solution, but it requires a large number of sampling steps to explore high-dimensional multi-modal posteriors, making it computationally expensive. In this paper, we propose a meta-learning strategy to build SGMCMC which can efficiently explore the multi-modal target distributions. Our algorithm allows the learned SGMCMC to quickly explore the high-density region of the posterior landscape. Also, we show that this exploration property is transferrable to various tasks, even for the ones unseen during a meta-training stage. Using popular image classification benchmarks and a variety of downstream tasks, we demonstrate that our method significantly improves the sampling efficiency, achieving better performance than vanilla SGMCMC without incurring significant computational overhead.

1. Introduction

Bayesian methods have received a lot of attention as powerful tools for improving the reliability of machine learning models. Bayesian methods are gaining prominence due to their ability to offer probability distributions over model parameters, thereby enabling the quantification of uncertainty in predictions. They find primary utility in safety-critical domains like autonomous driving, medical diagnosis, and finance, where the accurate modeling of prediction uncertainty often takes precedence over the predictions themselves. The integration of Bayesian modeling with (deep) neural networks, often referred to as Bayesian Neural Networks (BNNs), introduces exciting prospects for the development of secure and trustworthy decision-making systems.

However, there are significant problems for the successful application of BNNs in real-world scenarios. Bayesian inference in high-dimensional parameter space, especially for deep and large models employed for the applications mentioned above, is notoriously computationally expensive and often intractable due to the complexity of the posterior distribution. Moreover, posterior landscapes of BNNs frequently display multi-modality, where multiple high density regions exist, posing a significant challenge to efficient exploration and sampling. Due to this difficulty, the methods that are reported to work well for relatively small models, for instance, variational inference (Blei & McAuliffe, 2017) or Hamiltonian Monte Carlo (HMC) (Neal et al., 2011), can severely fail for deep neural networks trained with large amount of data, when applied without care.

Recently, Stochastic Gradient Markov Chain Monte Carlo (SGMCMC) methods (Welling & Teh, 2011; Chen et al., 2014; Ma et al., 2015) have emerged as powerful tools for enhancing the scalability of approximate Bayesian inference. This advancement has opened up the possibilities of applying Bayesian methods to large-scale machine learning tasks. SGMCMC offers a versatile array of methods for constructing Markov chains that converge towards the target posterior distributions. The simulation of these chains primarily relies on stochastic gradients, making them particularly suitable for BNNs trained on large-scale datasets. However, despite the notable successes of SGMCMC in some BNN applications (Welling & Teh, 2011; Chen et al., 2014; Ma et al., 2015; Zhang et al., 2020), there remains a notable challenge. Achieving optimal performance often demands extensive engineering efforts and hyperparameter tuning. This fine-tuning process typically involves human trial and error or resource-intensive cross-validation procedures. Furthermore, it’s worth noting that even with the use of SGMCMC methods, there remains room for improvement in efficiently exploring multi-modal posterior distributions.

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As a result, in practical applications, a trade-off between precision and computational resources often becomes necessary.

To address these challenges, we introduce a novel meta-learning framework tailored to promote the efficient exploration of SGMCMC algorithms. Traditional SGMCMC methods often rely on handcrafted design choices inspired by mathematical or physics principles. Recognizing the pivotal role these design components play in shaping the trade-off between exploration and exploitation within SGMCMC chains, we argue in favor of learning them directly from data rather than manually specifying them. To achieve this, we construct neural networks to serve as meta-models responsible for approximating the gradients of kinetic energy terms. These meta-models are trained using a diverse set of BNNs inference tasks, encompassing various datasets and architectural configurations. Our proposed approach, termed Learning to Explore (1.2E), exhibits several advantageous properties, including better mixing rates, improved prediction performance, and a reduced need for laborious hyperparameter tuning.

Our contributions can be summarized as follows:

- We introduce 1.2E, a novel meta-learning framework enhancing SGMCMC methods. In contrast to conventional hand-designed approaches and meta-learning approach (Gong et al., 2018), 1.2E learns the kinetic energy term directly, offering a more data-driven and adaptable solution.

- We present a multitask training pipeline equipped with a scalable gradient estimator for 1.2E. This framework allows the meta-learned SGMCMC techniques to generalize effectively across a wide range of tasks, extending their applicability beyond the scope of tasks encountered during meta-training.

- Using real-world image classification benchmarks, we demonstrate the remarkable performance of BNNs inferred using the SGMCMC algorithm discovered by 1.2E, both in terms of prediction accuracy and sampling efficiency.

2. Backgrounds

2.1. SGMCMC for Bayesian Neural Networks

Settings. In this paper, we focus on supervised learning problems with a training dataset \( D = \{(x_i, y_i)\}_{i=1}^n \) with \( x_i \) being observation and \( y_i \) being label. Given a neural network with a parameter \( \theta \in \mathbb{R}^d \), a likelihood \( p(y_i \mid x_i, \theta) \) and a prior \( p(\theta) \) are set up, together defining an energy function \( U(\theta) = -\sum_{i=1}^n \log p(y_i \mid x_i, \theta) - \log p(\theta) \). The goal is to infer the posterior distribution \( p(\theta \mid D) \propto \exp(-U(\theta)) \).

When the size of the dataset \( n \) is large, evaluating the energy function \( U(\theta) \) or its gradient \( \nabla_{\theta} U(\theta) \) may be undesirably costly as they require a pass through the entire dataset \( D \). For such scenarios, SGMCMC (Welling & Teh, 2011; Chen et al., 2014; Ma et al., 2015) is a standard choice, where the gradients of the energy function \( \nabla_{\theta} U(\theta) \) are approximated by a stochastic gradient computed from mini-batches. That is, given a mini-batch \( B \subset \{1, \ldots, n\} \) where \(|B| \ll n\), an unbiased estimator of the full gradient \( \nabla_{\theta} U(\theta) \) with \( B \) can be computed as

\[
\nabla_{\theta} \hat{U}(\theta) = -\frac{n}{|B|} \sum_{i \in B} \nabla_{\theta} \log p(y_i \mid x_i, \theta) - \nabla_{\theta} \log p(\theta)
\]

A complete recipe. There may be several ways to build a Markov chain leading to the target posterior distribution. Ma et al. (2015) presented a generic recipe that includes all the convergent SGMCMC algorithms as special cases, constituting a complete framework. In this recipe, a parameter \( \theta \) of interest is augmented with an auxiliary momentum variable \( r \), and an Stochastic Differential Equation (SDE) of the following form is defined for a joint variable \( z = (\theta, r) \in \mathbb{R}^{2d} \) as follows.

\[
\begin{align*}
dz &= -[(D(z) + Q(z))\nabla_z H(z) + \Gamma(z)]dt \\
&+ \sqrt{2D(z)}d\omega_t \\
H(z) &= U(\theta) + g(\theta, r) \\
\Gamma(z) &= \sum_{j=1}^{2d} \frac{\partial}{\partial z_j}(D_{ij}(z) + Q_{ij}(z)),
\end{align*}
\]

where \( g(\theta, r) \) is the conditional energy function of the momentum \( r \) such that \( p(z) \propto \exp(-H(z)) \) and \( \omega_t \) is a \( 2d \)-dimensional Brownian motion. Here, \( D(z) \in \mathbb{R}^{2d \times 2d} \) and \( Q(z) \in \mathbb{R}^{2d \times 2d} \) are restricted to be positive semi-definite and skew-symmetric, respectively. Given this SDE, one can obtain a SGMCMC algorithm by first substituting the full gradient \( \nabla_z H(z) \) with a mini-batch gradient \( \nabla_z \hat{H}(z) = \nabla_z (\hat{U}(\theta) + g(\theta, r)) \) and then discretizing it via a numerical solver such as symplectic Euler method. A notable example would be Stochastic Gradient Hamiltonian Monte Carlo (SGHMC) (Chen et al., 2014), where \( \theta(\theta, r) = \frac{1}{2} r^T M^{-1} r, \quad D(z) = \begin{bmatrix} 0 & 0 \\ 0 & C \end{bmatrix}, \quad \text{and} \quad Q(z) = \begin{bmatrix} 0 & -I \\ I & 0 \end{bmatrix} \) for some positive semi-definite matrices \( M \) and \( C \), leading to an algorithm when discretized with symplectic Euler method as follows.

\[
\begin{align*}
r_{t+1} &= r_t - \epsilon_t \nabla \hat{U}(\theta_t) - \epsilon_t C M^{-1} r_t + \xi_t \\
\theta_{t+1} &= \theta_t + \epsilon_t M^{-1} r_{t+1},
\end{align*}
\]

where \( \xi_t \sim \mathcal{N}(0, 2C\epsilon_t) \) and \( \epsilon_t \) is a step-size.

The complete recipe includes interesting special cases that introduce adaptive preconditioners to improve the mixing
of SGMCMC (Girolami & Calderhead, 2011; Li et al., 2016; Wenzel et al., 2020). For instance, Li et al. (2016) proposed Preconditioned Stochastic Gradient Langevin Dynamics (pSGLD), which includes RMSprop (Tieleman & Hinton, 2012)-like preconditioning matrix in the updates:

\[
\begin{align*}
\theta_{t+1} &= \theta_t - \epsilon_t [G(\theta_t) \nabla \hat{U}(\theta) + \Gamma(\theta_t)] + \xi_t \\
V(\theta_{t+1}) &= \alpha V(\theta_t) + (1 - \alpha) \frac{\nabla \hat{U}(\theta_t)}{n} \odot \frac{\nabla \hat{U}(\theta_t)}{n} \\
G(\theta_{t+1}) &= \text{diag}(1 \odot (\lambda I + \sqrt{V(\theta_{t+1})}))
\end{align*}
\]  

where \(\xi_t \sim \mathcal{N}(0, 2G(\theta_t)\epsilon_t)\) and \(\odot, \odot\) denotes elementwise division and multiplication, respectively. pSGLD exploits recent gradient information to adaptively adjust the scale of energy gradients and noise. However, this heuristic adjustment is still insufficient to efficiently explore the complex posteriors of BNNs (Zhang et al., 2020).

Recently, Zhang et al. (2020) introduced cyclic learning rate schedule for efficient exploration of multi-modal distribution. The key idea is using the spike of learning rate induced by cyclic learning rate to escape from a single mode and move to other modes. However, in our experiment, we find that SGMCMC with cyclical learning rate does not necessarily capture multi-modality and it also requires a large amount of update steps to move to other modes in practice.

**Prediction via Bayesian model averaging.** After inferring the posterior \(p(\theta | D)\), for a test input \(x_\ast\), the posterior predictive is computed as

\[
p(y_\ast | x_\ast, D) = \int_{\mathbb{R}^d} p(y_\ast | x_\ast, \theta)p(\theta | D)d\theta, \tag{4}
\]

which is also referred to as Bayesian Model Averaging (BMA). In our setting, having collected from the posterior samples \(\theta_1, \ldots, \theta_K\) from a convergent chain simulated from SGMCMC procedure, the predictive distribution is approximated with a Monte-Carlo estimator,

\[
p(y_\ast | x_\ast, D) \approx \frac{1}{K} \sum_{k=1}^{K} p(y_\ast | x_\ast, \theta_k). \tag{5}
\]

As one can easily guess, the quality of this approximation depends heavily on the quality of the samples drawn from Markov Chain Monte Carlo (MCMC) procedure. For over-parameterized deep neural networks that we are interested in, the target posterior \(p(\theta | D)\) is typically highly multi-modal, so simple SGMCMC methods suffer from poor mixing; that is, the posterior samples collected from those methods are not widely spread throughout the parameter space, so it takes exponentially many samples to achieve desired level of accuracy for the approximation. Hence, a good SGMCMC algorithm should be equipped with the ability to efficiently explore the parameter space, while still be able to stay sufficiently long in high-density regions. That is, it should have a right balance between exploration-exploitation.

**Meta Learning** Meta-learning, or learning to learn, refers to the algorithm that learns the useful general knowledge from source tasks that can transfer to the unseen tasks. Most meta-learning algorithms involve two levels of learning: an inner-loop and outer-loop (Metz et al., 2018). Inner-loop usually contains the training procedure of particular task. In our work, inner-loop for our meta-training is iteratively update the model parameter \(\theta\) by running SGMCMC with learnable transition kernel. Outer-loop refers to the training procedure of meta-parameter \(\phi\), which is done by minimizing meta-objective \(L(\phi)\).

**Meta Learning and MCMC** There exists line of work that parameterize the transition kernel of MCMC with trainable function for various purposes. Levy et al. (2017) used learnable invertible operator to automatically design the transition kernel of HMC for good mixing. For SGMCMC, Gong et al. (2018) parameterized the curl matrix and acceleration matrix using neural networks under the framework of Ma et al. (2015). Although Gong et al. (2018) is the closest work for our method, this work did not verified its task generalization, i.e., evaluated only on the dataset that was used for training. Moreover, the scale of the experiments in Gong et al. (2018) are limited to small-sized network architectures. We further demonstrate the limitation of Gong et al. (2018) in simulating large scale multimodal BNNs posterior, along with detailed discussions in Appendix E. To the best of our knowledge, our work firstly proposes the method to meta-learn SGMCMC that can generalize to unseen datasets and scale to large-scale BNNs.

**Limitation of Meta-SGMCMC** (Gong et al., 2018) Meta-SGMCMC aims to learn the diffusion matrix \(D(z)\) and the curl matrix \(Q(z)\) in Equation 1 to build a SGMCMC algorithm that can quickly converge to a target distribution. For this purpose, neural network \(D_f(z)\) and \(Q_f(z)\) are employed to model \(D(z)\) and \(Q(z)\). We point out that this parameterization has notable limitations, especially in terms of efficient exploration of high-dimensional multi-modal target distribution.

- According to the recipe in Equation 1, the diffusion and the curl that are dependent on \(z\) involves the additional correction term \(\Gamma(z)\). This hurts the scalability of the algorithm, as computing \(\Gamma(z)\) involves computing the gradient of \(D_f(z)\) and \(Q_f(z)\) with respect to \(z\). This amounts to a significant computational burden as the dimension of \(z\) increases and involves finite-difference approximation. While Meta-SGMCMC attempted to address this issue of computational cost
through finite difference, but it still introduces additional computation with time complexity $O(HD)$ where $H$ is number of hidden units of neural network in meta-sampler and $D$ is dimension of $\theta$. Also, it is important to note that adding such approximation can negatively impact the convergence of the sampler.

- The objective function of Meta-SGMCMC is the KL divergence between the target distribution and the distribution of the samples obtained from the Meta-SGMCMC chain. This objective is not computed in closed-from and nor its gradient. Gong et al. (2018) proposes to use Stein gradient estimator (Li & Turner, 2018) which is generally hard to tune. Moreover, Gong et al. (2018) uses Truncated BackPropagation Through Time (TBPTT) for the gradient estimation, which is a biased estimator of the true gradient, and limited in the length of the inner step for meta-learning due to the memory consumption.

3. Main contribution: Learning to Explore

3.1. Overcome the limitations of Meta-SGMCMC

To avoid aforementioned limitations of Meta-SGMCMC, we newly design the meta-learning method for SGMCMC as follows:

- We parameterize the gradients of the kinetic energy function and keep $D_f(z)$ and $Q_f(z)$ independent of $z$. We can avoid the additional computational cost of computing $\Gamma(z)$ without sacrificing the flexibility of the sampler.

- We use a new meta-objective called BMA meta-loss which is the Monte-Carlo estimator of the predictive distribution to enhance the exploration and performance of the learned sampler. Both of our meta-objective and its gradient can be computed in closed-form. Also, we employ Evolution Strategy (ES) (Salimans et al., 2017; Metz et al., 2019) for gradient estimation, which is an unbiased estimator of the true meta-gradient. This also consumes significantly less memory compared to analytic gradient methods, allowing us to keep the length of the inner-loop much longer during meta-learning.

3.2. Meta-learning framework for SGMCMC

Instead of using a hand-designed recipe for SGMCMC, we aim to learn the proper SGMCMC update steps through meta-learning. The existing works, both the methods using hand-designed choices or meta-learning (Gong et al., 2018), try to determine the forms of the matrices $D(z)$ and $Q(z)$ while keeping the kinetic energy $g(\theta, r)$ as simple Gaussian energy function, that is, $g(\theta, r) = r^\top M^{-1} r/2$. This choice indeed is theoretically grounded, which can be shown to be optimal when the target distribution is Gaussian (Betancourt, 2017), but may not be optimal for the complex multimodal posteriors of BNNs. We instead choose to learn $g(\theta, r)$ while keeping $D(z)$ and $Q(z)$ as simple as possible. We argue that the meta-learning approach based on this alternative parameterization is more effective in learning versatile SGMCMC procedure that scales to large BNNs.

More specifically, we parameterize the gradients of the kinetic energy function $\nabla_\theta g(\theta, r)$ and $\nabla_r g(\theta, r)$ with neural networks $\alpha_\theta(\theta, r)$ and $\beta_\theta(\theta, r)$ respectively, and set $D(z)$ and $Q(z)$ as in SGMCMC. The update step of SGMCMC, when discretized with symplectic Euler method is,

$$r_{t+1} = r_t - \epsilon_t [\nabla_\theta \tilde{U}(\theta_t) + \alpha_\theta(\theta_t, r_t) + C \beta_\theta(\theta_t, r_t)] + \xi_t$$

$$\theta_{t+1} = \theta_t + \epsilon_t \beta_\theta(\theta_t, r_{t+1}).$$

(6)

where $\xi_t \sim \mathcal{N}(0, 2C\epsilon_t)$. Since we parameterize $\nabla g(\theta, r)$ and do not explicitly define the form of $g(\theta, r)$, we make the following assumptions about the underlying function $g(\theta, r)$.

**Assumption 3.1.** There exists an energy function $g(\theta, r)$ whose gradients with respect to $\theta$ and $r$ are $\alpha_\theta(\theta, r)$ and $\beta_\theta(\theta, r)$ respectively, and $\int \exp(-g(\theta, r))dr$ is constant.

We also present another version of T2E which does not require additional assumptions in Appendix D. The neural networks $\alpha_\theta$ and $\beta_\theta$ are parameterized as two-layer Multi-Layer Perceptrons (MLPs) with 32 hidden units. Specifically, $\alpha_\theta$ and $\beta_\theta$ are applied to each dimension of parameter and momentum independently, similar to the commonly used learned optimizers (Andrychowicz et al., 2016; Metz et al., 2019). Again, following the common literature in learned optimizers (Metz et al., 2019), for each dimension of the parameter and momentum, we feed the corresponding parameter and momentum values, the stochastic gradients of energy functions for that element, and running average of the gradient at various time scales, as they are reported to encode the sufficient information about the loss surface geometry. See Appendix I for implementation details of $\alpha_\theta$ and $\beta_\theta$. By leveraging this information, we expect our meta-learned SGMCMC procedure to capture the multi-modal structures of the target posteriors of BNNs, and thus yielding a better mixing method.

3.3. Meta-Objective and Optimization

**Objective functions for meta-learning.** Meta-objective should reflect the meta-knowledge one wants to learn. We design the meta-objective based on the hope that samples collected through SGMCMC should be good at approximating the posterior predictive $p(y_x|x, D)$. In order to achieve this goal, we propose the meta-objective called BMA meta-loss. After the sufficient number of inner-updates, we col-
Algorithm 1 Meta training procedure

1: **Input:** Task distribution $P(T)$, inner iterations $N_{inner}$, outer iterations $N_{outer}$, step size $\epsilon$, noise scale $\sigma^2$, initial meta-parameter $\phi_0$.
2: **Output:** Meta parameter $\phi$.
3: for $j = 1, \ldots, N_{outer}$ do
4: Sample task $T_i \sim P(T)$
5: Initialize model parameter $\theta_0$ for $T_i$
6: Sample $\eta_i \sim \mathcal{N}(0, \sigma^2 I)$
7: $L(\phi + \eta_i) \leftarrow \text{InnerLoop}(\theta_0, \phi + \eta_i, \epsilon, N_{inner})$
8: $L(\phi - \eta_i) \leftarrow \text{InnerLoop}(\theta_0, \phi - \eta_i, \epsilon, N_{inner})$
9: $\nabla_\phi L \leftarrow \frac{1}{2\sigma^2} \eta_i (L(\phi + \eta_i) - L(\phi - \eta_i))$
10: $\phi \leftarrow \phi - \gamma \nabla_\phi L(\phi)$
11: end for

let $K$ parameter samples with some interval between them (thinning). Let $\theta_k(\phi)$ be the $k^{th}$ collected parameter, and we compute the Monte-Carlo estimator of the predictive distribution and use it as a meta-objective function (note the dependency of $\theta_k$ on the meta-parameter $\phi$, as it is a consequence of learning SGCMC with the meta-parameter $\phi$),

$$L(\phi) = -\log \frac{1}{K} \sum_{k=1}^{K} p(y_s | x_s, \theta_k(\phi)), \quad (7)$$

where $(x_s, y_s)$ is a validation data point.

Gradient estimation for meta-objective. Estimating the meta-gradient $\nabla_\phi L(\phi)$ is highly non-trivial (Metz et al., 2018; 2019), especially when the number of inner update steps is large. For instance, a naive method such as back-propagation through time would require memory grows linearly with the number of inner-steps, so become easily infeasible for even moderate sized models. One might consider using the truncation approximation, but that would result in a biased gradient estimator. Instead, we adapt ES (Salimans et al., 2017) with antithetic sampling scheme, which has been widely used in recent literature of training learned optimizer. Metz et al. (2019) showed that unrolled optimization with many inner-steps can lead to chaotic meta-loss surface and ES is capable of relieving this pathology by employing smooth log.

$$L(\phi) = \mathbb{E}_{\phi \sim \mathcal{N}(\phi, \sigma^2 I)} \left[ L(\phi) \right] \quad (8)$$

where $\sigma^2$ determines the degree of smoothing. Also, antithetic sampling is usually applied to reduce the estimation variance of $\nabla_\phi L(\phi)$. Through log-derivative trick, we can get unbiased estimator of (8),

$$\hat{g} = \frac{1}{N} \sum_{i=1}^{N} L(\phi + \eta_i) \frac{\eta_i}{\sigma^2} \quad (9)$$

where $\eta_i \overset{i.i.d.}{\sim} \mathcal{N}(0, \sigma^2 I)$. In addition, we can get another unbiased estimator $\hat{g}^{-1} = -\frac{1}{N} \sum_{i=1}^{N} L(\phi - \eta_i) \frac{\eta_i}{\sigma^2}$ by reusing the negative of $\eta_i$. By taking the average of two estimators, we can obtain the following gradient estimator.

$$\hat{\nabla}_\phi L(\phi) = \frac{1}{N} \sum_{i=1}^{N} \left[ \frac{L(\phi + \eta_i) - L(\phi - \eta_i)}{2\sigma^2} \right] \eta_i \quad (10)$$

The estimator is also amenable to parallelization, improving the efficiency of gradient computation.

3.4. Meta training procedure

**Generic pipeline.** General process of meta-training is as follows. First, for each inner-loop, we sample a task from the pre-determined task distribution. An inner-loop starts from an randomly initialized parameter and iteratively apply update step (6) to run a single chain of SGCMC. Please refer to Algorithm 2 for detailed description. In the initial stage of meta-training, the chains from these inner loops show poor convergence, but the performance improves as training progresses. Similar to general Bayesian inference, we consider the early part of the inner loop as a burn-in period and collect samples from the end of the inner-loop at regular intervals when evaluating the meta-objective. This training process naturally integrates the meta-learning and Bayesian inference in that mimicking the actual inference procedure of Bayesian methods in realistic supervised learning tasks. In Figure 1 we show that $t.2$E achieve desired level of accuracy for the approximation of posterior predictive with relatively small number of samples. This result indicates that $t.2$E has successfully acquired the desired properties through meta-training.

**Multitask training for better generalization.** In meta-learning, diversifying the task distribution is commonly known to enhance generalization. We include various neural network architectures and datasets in the task distribution to ensure that $t.2$E has sufficient generalization capacity. Also, we evaluate how the task distribution diversity affects the performance of $t.2$E in Table 11.

4. Experiments

In this section, we will evaluate the performance of $t.2$E in various aspects. Through extensive experiments, we would like to demonstrate followings:

- $t.2$E shows excellent performance on real-world image classification tasks and seamlessly generalizes to the tasks not seen during meta-training.
- $t.2$E produces similar predictive distribution to HMC as well as good mixing of BNNS posterior distribution in weight space.
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(a) fashion-MNIST (seen task)
(b) CIFAR-10
(c) CIFAR-100
(d) Tiny-ImageNet

Figure 1. Predictive performance trend of each method as the number of samples for BMA increases. L2E exhibits superior predictive accuracy compared to other baseline methods. Note that only the fashion-MNIST dataset is included in meta-training task distribution. We also plot the performance of HMC reference samples from Izmailov et al. (2021b) as dashed line for CIFAR-10 and CIFAR-100. For meta-training details of Meta-SGMCMC, please refer to Appendix I.4.

- L2E can effectively explore and sample from BNN posterior distribution, collecting diverse set of parameters both in weight space and function space.

Experimental details In this paragraph, we explain our experimental settings. We evaluate L2E and baseline methods on 4 datasets: fashion-MNIST (Xiao et al., 2017), CIFAR-10, CIFAR-100 (Krizhevsky et al., 2009) and Tiny-ImageNet (Le & Yang, 2015). Please refer to Appendix K.1 for full experimental setup and details. We report the mean and standard deviation of results over three different trials. Code is available at https://github.com/ciao-seohyeon/l2e.

Comparing L2E with HMC HMC is popular MCMC method since it can efficiently explore the target distribution and asymptotically converge to the target distribution, making it useful for inference in high-dimensional spaces. Due to its asymptotic convergence property, HMC often considered as golden standard in Bayesian Inference. However, its computational burden harms the applicability of HMC on modern machine learning tasks. Recently, Izmailov et al. (2021b) ran HMC on CIFAR-10 and CIFAR-100 datasets and released logits of HMC as reference. For deeper investigation of mixing of L2E, we compare L2E with HMC.

In order to compare the results with reference HMC samples from Izmailov et al. (2021b), we replicate the experimental setup of Izmailov et al. (2021b) including neural network architecture and dataset processing. The distinctive aspects of this setup are that they excluded data augmentation and replaced batch normalization layer in ResNet architecture to Filter Response Normalization (FRN) (Singh & Krishnan, 2020) to remove stochasticity in evaluating posterior distribution. We use the ResNet20-FRN architecture with Swish activations (Ramachandran et al., 2017) and we use 40960 samples for training all methods. Also, we compare L2E to HMC 1-D synthetic regression experiments in Appendix G. We also provide experimental results with data augmentation in Appendix H with other discussions.

Task distribution for meta-training We construct set of meta-training tasks using various datasets and model architectures. We use these tasks to meta-train meta-SGMCMC and L2E. Specifically, we use MNIST, fashion-MNIST, EMNIST (Cohen et al., 2017) and MedMNIST (Yang et al., 2021) as meta-training datasets. For model architecture, we fix the general structure with several convolution layers followed by readout MLP layer. For each outer training iteration, we randomly choose dataset and sample the configuration of architecture including number of channels, depth of the convolution layers and whether to use skip connections. See Appendix I for detailed configuration of task distribution. For evaluation, we use same meta-parameter of L2E and meta-SGMCMC for all experiments to check the generalization ability of each method.

4.1. Image classification results

Figure 1 illustrates the trend of predictive performance of each method as the number of samples for BMA increases. We confirm that L2E outperforms other baselines in terms of predictive accuracy on completely unseen datasets and architectures during meta-training. Among
datasets for evaluation, fashion-MNIST is the only dataset included in our task distribution. Despite not having seen other datasets during meta-training, L2E consistently outperforms other tuned baseline methods, showing that L2E can scale and generalize well to unseen larger tasks. Specifically, only DE shows comparable predictive accuracy compared to L2E in fashion-MNIST dataset. In general, L2E shows rapid performance gain and outperforms other methods with relatively small number of samples. According to Figure 2, L2E collects diverse weights and functions and this diversity presumably affect the huge performance gain from BMA.

On the other hand, Meta-SGMCMC exhibits poor predictive accuracy over all experiments. It converges very slowly compared to other methods and sticks in the low density region. This tendency gets worse when the evaluation task is significantly different from the distribution of meta-training tasks. In fashion-MNIST task, the performance of Meta-SGMCMC is still worse but relatively close to other methods than other tasks. Since learned sampler has seen fashion-MNIST dataset during meta-training, it adapts to this task to a moderate extent. However, it cannot produce even reasonable performance on other large scale tasks. We also evaluate L2E in the same experimental setup of Gong et al. (2018) and compare to the reported performance of Meta-SGMCMC in Appendix E. Without having seen CIFAR-10 dataset during meta-training, L2E significantly outperforms Meta-SGMCMC also in the experimental setup of Gong et al. (2018).

4.2. Similarity to the HMC samples

As mentioned above, we compare L2E to HMC samples from Izmailov et al. (2021b) to objectively evaluate and investigate whether L2E has converged to target distribution. We measure the agreement between predictive distribution of HMC and L2E, and the difference of probability vector of two methods following Izmailov et al. (2021b). Please refer to Appendix K for details of metrics. In Table 1, we confirm that L2E makes higher fidelity approximation to predictive distribution of HMC than other baselines both in CIFAR-10 and CIFAR-100. This implies that L2E shows good mixing in function space by effectively capturing multi-modality of complex BNNs posterior distribution. This high fidelity in predictive distribution can be attributed to the use of BMA-loss as meta-objective.

Also, we conduct 1-D synthetic regression task to visually check whether L2E can capture the epistemic uncertainty similar to HMC. Figure 8 shows that L2E better captures “in-between” uncertainty than DE. Although HMC captures epistemic uncertainty more effectively than L2E, we observe that L2E captures epistemic uncertainty in a similar manner to HMC. Please refer to Appendix G for details.

4.3. Capturing multi-modality

In Figure 2c and Figure 2b, we observe the behavior of DE, SGMCMC, and L2E in function space. In Figure 2b, we display the test error surface using a 2-dimensional subspace spanned by the first three collected parameters for each method following Garipov et al. (2018). Parameters of DE clearly located on multiple distinct modes as expected.
Table 2. AUROC for evaluating OOD. For in-distribution dataset, we use CIFAR-10 and CIFAR-100, while CIFAR-10, CIFAR-100, SVHN, and Tiny-ImageNet serve as OOD dataset.

<table>
<thead>
<tr>
<th>Metric</th>
<th>Dataset</th>
<th>CSGMCMC</th>
<th>Meta-SGMCMC</th>
<th>L2e</th>
<th>HMC</th>
</tr>
</thead>
<tbody>
<tr>
<td>ESS/s</td>
<td>fashion MNIST</td>
<td>21.95±0.64</td>
<td>33.54±4.42</td>
<td>136.31±0.42</td>
<td></td>
</tr>
<tr>
<td>DE</td>
<td>CIFAR-10</td>
<td>56.61±0.51</td>
<td>17.31±6.31</td>
<td>82.97±0.57</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CIFAR-100</td>
<td>41.14±1.51</td>
<td>48.29±7.43</td>
<td>63.81±0.00</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tiny-ImageNet</td>
<td>2.43±0.01</td>
<td>0.91±0.19</td>
<td>2.62±0.00</td>
<td></td>
</tr>
</tbody>
</table>

4.5. Robustness under covariate shift

Next, we consider CIFAR-10-C (Hendrycks & Dietterich, 2019) for evaluating robustness to covariate shift. In Table 3, DE outperforms L2e and CSGMCMC for all intensity levels. L2e shows competitive performance under mild corruption, but as the corruption intensifies, the performance of L2e significantly drops and shows worst accuracy over all methods. HMC also shows this trend, showing worst performance in general. Since L2e makes similar predictive distribution with HMC in CIFAR-10, this aligns with the result from (Izmailov et al., 2021a;b) that HMC and methods having high fidelity to HMC suffer greatly from the covariate shift. Although L2e is not robust at covariate shift, it is understandable considering the similarity of L2e to HMC in function space.

4.6. Convergence analysis

We also evaluate sampling efficiency and degree of mixing of L2e using ESS and $cR^2_0$ (Sommer et al., 2024). Please refer to Appendix L for details of metrics. In Table 4, L2e is the only method that consistently demonstrates decent performance both in terms of ESS and $cR^2_0$ across experiments. On the other hand, other methods show poor mixing, indicating that they hardly explore multi-modal BNNS posterior distributions.

5. Conclusion

In this work, we introduced a novel meta-learning framework called L2e to improve SGMCMC methods. Unlike conventional SGMCMC methods that heavily rely on manually designed components inspired by mathematical or physics principles, we aim to learn critical design components of SGMCMC directly from data. Through experiments, we show numerous advantages of L2e over existing
Learning to Explore for Stochastic Gradient MCMC

SGMCMC methods, including better mixing, improved prediction performance. Our approach would be a promising direction to solve several challenges that SGMCMC methods face in BNNs.

Ethics and Reproducibility statement Please refer to Appendix K for full experimental detail including datasets, models, and evaluation metrics. We have read and adhered to the ethical guideline of International Conference on Machine Learning in the course of conducting this research.

Acknowledgement

The authors gratefully acknowledge Giung Nam for helping experiments and constructive discussions. This work was partly supported by Institute of Information & communications Technology Planning & Evaluation (IITP) grant funded by the Korea government (MSIT) (No.RS-2019-II190075, Artificial Intelligence Graduate School Program (KAIST), No.2022-0-00184, Development and Study of AI Technologies to Inexpensively Conform to Evolving Policy on Ethics, and No.2022-0-00713, Meta-learning Applicable to Real-world Problems) and the National Research Foundation of Korea (NRF) grant funded by the Korea government (MSIT) (No. 2022R1A5A708390812). This research was supported with Cloud TPUs from Google’s TPU Research Cloud (TRC).

Impact Statement

This paper presents work whose goal is to advance the field of Machine Learning. There are many potential societal consequences of our work, none which we feel must be specifically highlighted here.

References


Learning to Explore for Stochastic Gradient MCMC


Yang, J., Shi, R., and Ni, B. Medmnist classification decathlon: A lightweight automl benchmark for medical

A. Analysis of exploration property

In this section, we analyze how $L_2E$ can collect diverse set of parameters with a single MCMC chain. In Figure 3, we analyze the behavior of $L_2E$ in downstream tasks (CIFAR-10,100) and in Figure 4, we visualize the scale of outputs of $\alpha_\phi$ and $\beta_\phi$ on the regular grid of input values following Gong et al. (2018). Firstly, we plot $l_2$ norm of $\Delta \theta = \theta_{t+1} - \theta_t$ at time $t$ and training cross-entropy loss (Negative Log-Likelihood (NLL)) for 200 epochs and comparing it with DE and CSGMCMC.

Recall the update rule of $L_2E$,

$$
    r_{t+1} = r_t - \epsilon_t \left[ \nabla_\theta \tilde{U}(\theta_t) + \alpha_\phi(\theta_t, r_t) + C\beta_\phi(\theta_t, r_t) \right] + \xi_t \\
    \theta_{t+1} = \theta_t + \epsilon_t \beta_\phi(\theta_t, r_{t+1}).
$$

(11)

where $\xi_t \sim \mathcal{N}(0, 2C\epsilon_t)$. According to the equation above, $\beta_\phi$ is responsible for updating $\theta$, so tracking $l_2$ norm of $\Delta \theta$ is same as tracking $l_2$ norm of $\beta_\phi$ since $||\beta_\phi||^2 = \frac{||\Delta \theta||^2}{\epsilon_t}$.

Figure 3. Plots of $||\Delta \theta||^2$ and train NLL during training of $L_2E$, CSGMCMC, DE on CIFAR-10. Unlike other methods, $L_2E$ actively updates $\theta$ in the local minima while maintaining training NLL as nearly constant.

Figure 4. Contour plots of absolute value of outputs of $\beta_\phi$(top) and $\alpha_\phi$(bottom) on the grid. $\beta_\phi$ produces large magnitude of output when $\nabla_\theta \tilde{U}(\theta)$ is high. When $\nabla_\theta \tilde{U}(\theta)$ gets smaller, the overall magnitude decreases as expected, but even when $\nabla_\theta \tilde{U}(\theta)$ is nearly zero, $\beta_\phi$ can still allow the sampler to move around posterior distribution when integrated with high momentum value. The regions marked with red dashed boxes can be beneficial for exploration in high density regions. $\alpha_\phi$ is proportional to $\nabla_\theta \tilde{U}(\theta)$ in general, which helps the sampler fastly converge to the high density region.
In Figure 4, we plot absolute value of outputs of $\alpha$ gradients learned the general knowledge to explore high density regions among different modes. (e.g., see Figure 2b) support that knowledge of posterior information across tasks for efficient exploration in low loss regions. Various experimental results $L$ constant. This trend is consistently observed in both CIFAR-10 and CIFAR-100, implying that small at local minima. Additionally, we notice that $L_2E$ actively updates $\theta$ MCMC. This tendency is different from other gradient-based optimizer or $L_2$.

In Figures 3 and 5, we find that $L_{\text{smooth}}$ and running average of $\nabla \theta^k U$ as inputs, analyzing the function itself is a complex problem. Therefore, to simplify the analysis, we follow the approach of Gong et al. (2018), where we fix other inputs except for the statistics we are interested in. For $\beta_\phi$, we choose three different fixed values of $\theta$ and plot the results, while for $\alpha_\phi$, we fix the momentum. We assume that there is no running average of $\nabla \theta^k U(\theta)$ so that running average term is fixed to $\nabla \theta^k U(\theta)$.

Since the value of $\theta$ itself does not encode the information about the posterior landscape, there is no clear distinction among contour plots in top row with different $\theta$ values. In general, the scale of outputs of $\alpha_\phi$ and $\beta_\phi$ is proportional to $\nabla \theta^k U(\theta)$ which is desirable for fast convergence to high density regions of posterior distribution. For $\beta_\phi$, when $\nabla \theta^k U(\theta)$ gets smaller, the overall magnitude of output decreases as expected, but even when $\nabla \theta^k U(\theta)$ is nearly zero, $\beta_\phi$ can still make large magnitude of update of $\theta$ when integrated with high momentum value. Also, when momentum is nearly zero, it still allows sampler to explore posterior distribution when $\nabla \theta^k U(\theta)$ is far from zero. This complementary relationship between two statistics is strength of $L_2E$ that the movement of sampler is not solely depend on a single statistic so that it can exploit more complex information about loss geometry unlike standard SGHMC. $\alpha_\phi$ produces an output proportional to the scale of the gradient regardless of the momentum values. This implies that $\alpha_\phi$ helps the acceleration of sampler in low-density regions as it is added to the energy gradient in Equation 11.

One limitation of our analysis is that we focus on analyzing the magnitude of the function output rather than its direction. Although the magnitude of $\alpha_\phi$ and $\beta_\phi$ are closely connected to the exploration property, delving deeper into how $L_2E$ traverses complex and multi-modal BNNs posterior landscape would be an interesting direction of future research.

### B. Bayesian Model Average(BMA) vs Cross-Entropy(CE) meta-loss

In this section, we will explain why BMA meta-loss enhances the exploration of the sampler, by comparing it with CE meta-loss. Average CE loss of individual models along the optimizer’s trajectory on meta-training tasks has been used as a meta-objective to train learned optimizers (Andrychowicz et al., 2016; Metz et al., 2019; 2022b) on classification tasks. Two different objectives have following forms:

$$L^{\text{CE}}(\phi) = -\frac{1}{K} \sum_{k=1}^{K} \log p(y^*_k | x^*_k, \theta_k(\phi))$$

$$L^{\text{BMA}}(\phi) = -\log \frac{1}{K} \sum_{k=1}^{K} p(y^*_k | x^*_k, \theta_k(\phi))$$

(12)
where \((x_*, y_*)\) is a validation data point. While BMA meta-loss is similar to CE meta-loss, BMA meta-loss differs significantly from CE meta-loss. BMA meta-loss is Monte Carlo approximation of the posterior predictive distribution for validation data points. Actually, we gather models along the sampler’s trajectory and minimize CE-loss of the average probability of collected models. BMA meta-loss not only encourages the sampler to minimize the average CE-loss of individual models but also promotes increased functional diversity among collected models. According to Wood et al. (2023, Equation 7), this loss of the ensemble model is decomposed as “ensemble loss = average individual loss - ambiguity”. Ambiguity refers to the difference among the ensemble models and individual models, and the larger it is, the more BMA meta-loss decreases. In Figure 6, we observe that L2E trained with BMA meta-loss exhibits a much greater loss barrier between collected parameters than L2E trained with CE meta-loss, indicating the higher exploration and larger functional diversity among samples. Also, our extensive downstream experiments and visualization demonstrates that L2E with BMA meta-loss actually increases functional diversity among collected models.

\[
L_{\text{diversity}} = \text{average individual loss} - \text{ambiguity}
\]

![Figure 6. Ablation study of BMA vs CE meta-loss on CIFAR-10.](image)

C. L2E on Text Dataset

In order to check whether L2E can adapt well to the other modalities (e.g. text dataset), we additionally conduct text classification on IMDB dataset with CNN-LSTM architecture following Wenzel et al. (2020) and Izmailov et al. (2021b).

For HMC, we use checkpoints of 1200 samples from Izmailov et al. (2021b). For other MCMC methods, we collect 50 samples with 50 epochs of thinning interval with 100 burnin epochs. Table 6 demonstrates that L2E shows competitive performance on the text dataset, meaning that L2E can still work well on unseen modalities. This transfer of knowledge from image datasets to text dataset supports that the generalization capacity of L2E is strong compared to Gong et al. (2018), which is only proven to generalize well to similar datasets with meta-training dataset.

D. Results on alternative parameterization

Although parameterizing \(\nabla g(\theta, r)\) allows learned sampler to be expressive and efficient, we should make assumptions on the underlying \(g(\theta, r)\). To avoid introducing additional assumptions, we propose another version of L2E that directly parameterizes kinetic energy function \(g(\theta, r)\) rather than its gradients and evaluate whether it can achieve comparable performance comparable to L2E. Specifically, we fix \(p(r|\theta)\) as normal distribution and parameterize its mean with \(f_\phi(\theta)\) that takes \(\theta, \nabla_\theta U(\theta)\) as inputs. This approach eliminates the need for assumptions regarding the existence and integrability of unnormalized probability density function. Since we set \(p(r|\theta) \sim N(f_\phi(\theta), I)\), kinetic energy function of \(p(r|\theta)\) is \(g(\theta, r) = (r - f_\phi(\theta))^T(r - f_\phi(\theta))\). Therefore, we can get \(\nabla_\theta g(\theta, r) = f_\phi(\theta)\nabla_\theta f_\phi(\theta)\) and \(\nabla_r g(\theta, r) = r - f_\phi(\theta)\). Then, update rule for this parameterization is as follows

\[
\begin{align*}
    r_{t+1} &= r_t - \epsilon_t (\nabla_\theta U(\theta_t) + f_\phi(\theta)\nabla_\theta f_\phi(\theta) + \alpha (r_t - f_\phi(\theta))) + \xi_t \\
    \theta_{t+1} &= \theta_t + \epsilon_t (r_t - f_\phi(\theta))
\end{align*}
\]

where \(\xi_t \sim N(0, 2\epsilon_t \alpha)\). We will refer to this version of sampler as Kinetic-L2E. In Table 5, Kinetic-L2E shows comparable performance to L2E across all image classification experiments. Also, Figure 7 demonstrates the exploration capacity of Kinetic-L2E. These results indicate that both versions can achieve similar performance in terms of predictive accuracy and exploration.
Both versions of \( L^2E \) have their own strength and weakness. \( L^2E \) requires assumptions on the underlying kinetic energy function to ensure the convergence guarantee by Ma et al. (2015), but it is computationally efficient and can model complex functions. Kinetic-\( L^2E \) guarantee the convergence without additional assumptions on \( g(\theta, r) \), but requires additional gradient computations and conditional distribution \( p(r|\theta) \) is restricted to specific distribution which can possibly harm the flexibility of the sampler. We recommend to consider these trade-offs for further applications.

**Table 5.** Results of kinetic energy parameterization of \( L^2E \).

<table>
<thead>
<tr>
<th>Dataset</th>
<th>( L^2E ) ACC ↑</th>
<th>NLL ↓</th>
<th>ECE ↓</th>
<th>KLD ↑</th>
</tr>
</thead>
<tbody>
<tr>
<td>fashion-MNIST</td>
<td>0.9175</td>
<td>0.2425</td>
<td>0.0104</td>
<td>0.1334</td>
</tr>
<tr>
<td></td>
<td>0.9166</td>
<td>0.2408</td>
<td>0.0078</td>
<td>0.0507</td>
</tr>
<tr>
<td>CIFAR-10</td>
<td>0.9131</td>
<td>0.2867</td>
<td>0.0527</td>
<td>0.4152</td>
</tr>
<tr>
<td></td>
<td>0.9123</td>
<td>0.2909</td>
<td>0.0563</td>
<td>0.4344</td>
</tr>
<tr>
<td>CIFAR-100</td>
<td>0.7013</td>
<td>1.114</td>
<td>0.1212</td>
<td>1.735</td>
</tr>
<tr>
<td></td>
<td>0.6999</td>
<td>1.131</td>
<td>0.1403</td>
<td>1.749</td>
</tr>
<tr>
<td>Tiny-ImageNet</td>
<td>0.5561</td>
<td>1.966</td>
<td>0.1731</td>
<td>1.907</td>
</tr>
<tr>
<td></td>
<td>0.5583</td>
<td>1.846</td>
<td>0.1423</td>
<td>0.9139</td>
</tr>
</tbody>
</table>

**Table 6.** Results on IMDB classification task, following the experimental setup of Izmailov et al. (2021b)

<table>
<thead>
<tr>
<th>Method</th>
<th>ACC ↑</th>
<th>NLL ↓</th>
</tr>
</thead>
<tbody>
<tr>
<td>DE</td>
<td>0.867 ± 0.000</td>
<td>0.386 ± 0.000</td>
</tr>
<tr>
<td>CSGMCMC</td>
<td>0.848 ± 0.007</td>
<td>0.401 ± 0.032</td>
</tr>
<tr>
<td>HMC</td>
<td>0.868 ± 0.000</td>
<td>0.308 ± 0.000</td>
</tr>
<tr>
<td>Meta-SGMCMC</td>
<td>0.820 ± 0.003</td>
<td>0.401 ± 0.001</td>
</tr>
<tr>
<td>( L^2E )</td>
<td>\textbf{0.873 ± 0.001}</td>
<td>\textbf{0.301 ± 0.001}</td>
</tr>
</tbody>
</table>

**E. Comparison with Gong et al. (2018)**

In Table 7, we demonstrate the difference between Meta-SGMCMC (Gong et al., 2018) and \( L^2E \). Meta-SGMCMC aims to build a sampler through meta-learning that rapidly converges to the target distribution and performs accurate simulation. This goal aligns with the objectives of all SGMCMC methods. However, our approach is specifically designed with the more concrete purpose of effectively simulating multi-modal DNN posterior distribution and also generalizing to unseen problems. We compare \( L^2E \) to Meta-SGMCMC in various aspects in the following subsections.

**E.1. Difference in parameterization of meta models**

The update rule for \( z = (\theta, r) \in \mathbb{R}^{2d} \) presented in Gong et al. (2018) is as following.

\[
\begin{align*}
    r_{t+1} &= (1 - \epsilon_t D_f(z_t))r_t - \epsilon_t Q_f(z_t) \nabla \theta U(\theta_t) + \epsilon_t \Gamma_r(z_t) + \xi_t \\
    \theta_{t+1} &= \theta_t + \epsilon_t Q_f(z_t) r_t + \epsilon_t \Gamma_\theta(z_t) \\
    \xi_t &\sim \mathcal{N}(0, 2\epsilon_t D_f(z_t)), \quad \Gamma_i(z) := \sum_{j=1}^{2d} \frac{\partial}{\partial z_j} (D_{ij}(z) + Q_{ij}(z))
\end{align*}
\]

In our main text, we point out that parameterizing \( D_f \) and \( Q_f \) makes additional computational burden. Additionally, since \( D_f \) and \( Q_f \) mainly function as multipliers for gradient and momentum, learning them may not be as effective as it should...
Table 7. Comparison between Gong et al. (2018) and L2E

<table>
<thead>
<tr>
<th>Purpose of meta-learning</th>
<th>Gong et al. (2018)</th>
<th>L2E</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fast convergence, low bias</td>
<td>Efficient exploration of multi-modal BNNS posterior</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Meta-training target</th>
<th>Gong et al. (2018)</th>
<th>L2E</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diffusion and curl matrix</td>
<td>Gradient of kinetic energy</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Meta-objective</th>
<th>Gong et al. (2018)</th>
<th>L2E</th>
</tr>
</thead>
<tbody>
<tr>
<td>KL(q_t</td>
<td>π)</td>
<td>− log 1/K ( \sum_{k=1}^{K} p(y_*</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Meta-gradient estimation</th>
<th>Gong et al. (2018)</th>
<th>L2E</th>
</tr>
</thead>
<tbody>
<tr>
<td>TBPTT (Werbos, 1990)</td>
<td>ES (Salimans et al., 2017)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Generalizes to</th>
<th>Gong et al. (2018)</th>
<th>L2E</th>
</tr>
</thead>
<tbody>
<tr>
<td>unseen classes in the same dataset</td>
<td>completely different datasets</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Scales to</th>
<th>Gong et al. (2018)</th>
<th>L2E</th>
</tr>
</thead>
<tbody>
<tr>
<td>small scale architecture</td>
<td>large scale architecture</td>
<td></td>
</tr>
</tbody>
</table>

Table 8. Experiments on CIFAR-10 following experiments in Gong et al. (2018). Without having seen CIFAR-10 dataset during meta training, L2E outperforms reported performance of Meta-SGMCMC while using significantly less computational cost than Meta-SGMCMC.

<table>
<thead>
<tr>
<th>Methods</th>
<th>NT ACC</th>
<th>NT+AF ACC</th>
<th>NT+Data ACC</th>
<th>NT NLL/100</th>
<th>NT+AF NLL/100</th>
<th>NT+Data NLL/100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meta-SGMCMC</td>
<td>78.12±0.035</td>
<td>74.41±0.11</td>
<td>89.97±0.04</td>
<td>68.88±0.15</td>
<td>79.55±0.057</td>
<td>15.66±0.28</td>
</tr>
<tr>
<td>L2E</td>
<td>79.21±0.203</td>
<td>75.91±0.200</td>
<td>92.49±0.234</td>
<td>63.23±0.46</td>
<td>72.11±0.22</td>
<td>11.82±0.81</td>
</tr>
</tbody>
</table>

be for effective exploration in low energy regions. In low energy regions where the norm of gradient and momentum are extremely small, it is difficult to make reasonable amount of update of \( \theta \) for exploration by multiplying \( Q_f \) to momentum and gradient. Also, \( Q_f \) should be clipped by some threshold for practical issue. By contrast, In the update rule of L2E (6), \( \alpha_\phi, \beta_\phi \) are added to the gradient, which is more suitable for controlling the magnitude and direction of update in low energy regions. Also, ablation study in Table 11 shows the inferior performance of learning \( D_f \) and \( Q_f \) in terms of classification accuracy and predictive diversity in our setting. Therefore, we choose to learning kinetic energy gradient is better than learning \( D_f \) and \( Q_f \), as it avoids additional computation and allows the sampler to mix better especially around the low energy region.

E.2. Difference in meta-training procedure

There are significant differences between two methods in meta-training pipeline. Firstly, the most notable distinction is that Meta-SGMCMC learns with only one task, requiring the training of a new learner for each problem. This poses an issue as a single learner may not generally apply well to various problems. In contrast, our approach involves sampling from a task set composed of diverse datasets and architectures for training. Another crucial difference is the choice of estimator for the meta-objective gradients. Gong et al. (2018) employs Truncated BackPropagation Through Time (TBPTT) which truncates computational graphs of a long inner-loop to estimate the meta gradient. While this approach can save computational cost by avoiding backpropagation through long computational graph, it results in a biased estimator of meta-gradient. To address these issues, we utilize ES to compute unbiased estimator of meta-gradient efficiently.

E.3. Experimental results

In order to compare L2E with Meta-SGMCMC, we exactly replicate the experimental setup of MNIST and CIFAR-10 experiments in Gong et al. (2018) and evaluate performance of L2E. These experiments evaluate how well each method generalizes to unseen Neural Network architecture (NT), activation function (AF) and dataset (Data) which were unseen during the meta-learning process. For the experimental details, please refer to the experiments section of Gong et al. (2018). We use same scale of metrics in Gong et al. (2018) for conveniently comparing two methods. We report Accuracy (ACC) and NLL on test dataset. Since L2E used MNIST dataset during meta-train, we do not evaluate Dataset generalization experiments on MNIST. Meta-SGMCMC used 20 parallel chains with 100 epochs for MNIST and 200 epochs for CIFAR-10.
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Table 9. Experiments on MNIST following experiments in Gong et al. (2018).

<table>
<thead>
<tr>
<th>Methods</th>
<th>NT ACC</th>
<th>NT+AF ACC</th>
<th>NT+Data ACC</th>
<th>NT NLL/100</th>
<th>NT+AF NLL/100</th>
<th>NT+Data NLL/100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meta-SGMCMC</td>
<td>98.36±0.02</td>
<td>97.72±0.02</td>
<td>98.62±0.02</td>
<td>640±6.25</td>
<td>875±3.19</td>
<td>230±3.23</td>
</tr>
<tr>
<td>L2E</td>
<td>98.39±0.05</td>
<td>98.07±0.08</td>
<td>-</td>
<td>558±3.19</td>
<td>679±6.65</td>
<td>-</td>
</tr>
</tbody>
</table>

Figure 8. 1-D synthetic regression of L2E, DE, HMC. See Appendix G for details and discussion.

We use single chain with 100 burn-in epochs for both experiments, and use 10 thinning epochs for collecting 20 samples.

In Table 8 and Table 9, we confirm that L2E outperforms Meta-SGMCMC for all generalization types in terms of ACC and NLL despite using significantly less computational cost. Notably, in CIFAR-10 experiment, despite L2E was not trained on the CIFAR-10 during meta-learning, L2E significantly outperforms Meta-SGMCMC with a wide margin indicating that our approach better generalizes to unseen datasets compared to Meta-SGMCMC.

F. Discussion on meta-training objectives

Previous studies have proposed various meta-objectives to achieve goals similar to ours. Gong et al. (2018) minimizes the KL(qt|π), where π is target distribution and qt is the marginal distribution of θ at time t for good mixing. On the other hand, Levy et al. (2017) employs meta-objective maximizing the jump distances between samples in weight space and simultaneously minimizing the energy in order to make sampler rapidly explore between modes. However, explicitly maximizing the jump distances in weight space can be easily cheated, as the distances between weights does not necessarily lead to the difference in the functions, resulting in trivial sampler with which achieving the balance between convergence and exploration is hard. Also, minimizing the divergence with target distribution seems sensible, but due to intractable qt, computing the gradient of qt should resort to gradient estimator. Gong et al. (2018) used Stein-gradient estimator, which requires multiple independent chains so it harms scalability. Also, this objective does not lead the learned sampler to explore multi-modal distribution. Gong et al. (2018, Figure 3) shows that the learned sampler quickly converges to low energy region, but learned friction term Df restricts the amount of update in low energy region, limiting the exploration behaviour. Among choices, we find out that BMA meta-loss is a simple yet effective meta-objective that naturally encodes exploration-exploitation balance without numerical instability and exhaustive hyperparameter tuning.

G. 1-D synthetic regression

We conduct 1-D synthetic regression task to visually check whether L2E can capture the epistemic uncertainty. For the training data, we generate 1000 data points from underlying true function y = sin(x), within the interval [-5, 1] and [1, 4]. We use DE and HMC as baselines. We collect 50 parameters for each methods and plot the mean prediction and 95% confidence interval of the prediction. For L2E, we do not fine-tune the learned sampler used in the main experiments. We use thinning interval of 50 training steps and 1000 burn-in steps for L2E, 300 training steps for each single solution of DE and 1000 burn-in steps and 100 leap-frog steps for HMC. We use 2 layers MLP with 100 hidden units and ReLU activation to estimate the function.
Effective method for capturing epistemic uncertainty should make confident predictions for the training data and should be uncertain on OOD data points. Figure 8 shows that L2E better captures ‘in-between’ uncertainty than DE. L2E generally produces diverse predictions for out-of-distribution data points, especially for the input space between two clusters of training data while DE shows relatively confident prediction in that region. HMC is known for the golden standard for posterior inference in Bayesian method. Our experimental result aligns with this common knowledge since HMC is the best in terms of capturing epistemic uncertainty among three methods especially in areas out of the range of training points. While our approach falls short of HMC, it demonstrates better uncertainty estimation than that of DE and shows similar predictive uncertainty with HMC between two training points cluster. These experimental results align with the OOD detection experiments and convergence diagnostics presented in our main text, indicating L2E performs effective posterior inference. It is important to note that even without meta-training on the regression datasets, L2E can adapt well to the regression problem.

**H. Image classification with Data Augmentation**

Since applying data augmentation violates Independently and Identically Distributed (IID) assumption of the dataset which is commonly assumed by Bayesian methods, this can lead to model misspecification (Wenzel et al., 2020; Kapoor et al., 2022) and under model misspecification, bayesian posterior may not be the optimal for the BMA performance (Masegosas, 2020). Therefore, prior work such as Izmailov et al. (2021b) argued the incompatibility between Bayesian methods and data augmentation. However, data augmentation is an indispensable technique in modern machine learning so it is also interesting to see how L2E performs with data augmentation. We run the image classification experiments on CIFAR-10 and CIFAR-100 with random crop and horizontal flip for the data augmentation. Since using data augmentation introduces Cold Posterior Effect (Wenzel et al., 2020) for Bayesian methods such as CSGMCMC and L2E, we additionally tune the temperature hyperparameter for these methods. We tune the temperature for each methods, using $T = 0.0001$ in both experiments. Please refer to Wenzel et al. (2020) for detailed analysis of Cold Posterior Effect. We collect 10 samples for all methods for these experiments.

In Table 10, when applying data augmentation, the performance gap between CSGMCMC and L2E becomes far less stark than without data augmentation. Moreover, DE outperforms other baselines with a large margin in CIFAR-10 and shows similar results with L2E in CIFAR-100 since L2E significantly outperforms other method without data augmentation. With data augmentation, it is not very surprising that DE outperforms other Bayesian methods like L2E and CSGMCMC in terms of predictive accuracy and calibration since they suffer from model misspecification and temperature tuning can partially handle this problem (Kapoor et al., 2022). Observing the high agreement between L2E and HMC in the function space suggests that L2E effectively approximates the predictive distribution of the target posterior distribution. However, when techniques that violate assumptions for model likelihood are applied, correctly simulating target distribution does not necessarily mean the superior performance than other methods. As a result, there may be a reduction or even a reversal in the performance gap compared to other methods. Nevertheless, L2E demonstrates comparable performance to DE with less compute on datasets like CIFAR-100. Also, L2E shows slightly better performance than Bayesian method, CSGMCMC in CIFAR-10 and outperforms with a wide margin in CIFAR-100 experiment. When it comes to predictive diversity, L2E significantly outperforms baselines on both experiments. Although applying data augmentation introduces significant variations to the posterior landscape, we confirm that L2E still maintains the exploration property. To sum up, we argue that L2E is still practical method even with data augmentation since it shows competitive predictive performance and efficiently explores the posterior landscape.

![Table 10. Results of image classification with data augmentation.](image)

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Method</th>
<th>ACC ↑</th>
<th>NLL ↓</th>
<th>ECE ↓</th>
<th>KLD ↑</th>
</tr>
</thead>
<tbody>
<tr>
<td>DE</td>
<td>0.931±0.002</td>
<td>0.209±0.003</td>
<td>0.017±0.001</td>
<td>0.293±0.002</td>
<td></td>
</tr>
<tr>
<td>CIFAR-10</td>
<td>CSGMCMC</td>
<td>0.923±0.003</td>
<td>0.234±0.007</td>
<td>0.031±0.005</td>
<td>0.142±0.019</td>
</tr>
<tr>
<td>L2E</td>
<td>0.926±0.001</td>
<td>0.235±0.002</td>
<td>0.037±0.001</td>
<td>0.391±0.005</td>
<td></td>
</tr>
<tr>
<td>DE</td>
<td>0.708±0.001</td>
<td>1.048±0.001</td>
<td>0.092±0.001</td>
<td>0.765±0.003</td>
<td></td>
</tr>
<tr>
<td>CIFAR-100</td>
<td>CSGMCMC</td>
<td>0.683±0.001</td>
<td>1.114±0.005</td>
<td>0.013±0.001</td>
<td>0.319±0.008</td>
</tr>
<tr>
<td>L2E</td>
<td>0.705±0.003</td>
<td>1.066±0.001</td>
<td>0.095±0.003</td>
<td>0.996±0.017</td>
<td></td>
</tr>
</tbody>
</table>
Learning to Explore for Stochastic Gradient MCMC

Figure 9. Meta loss at each outer iteration during the meta training process. For Meta-SGMCMC, it is evident that the meta loss is highly unstable depending on hyperparameters such as Q clip and step size. Conversely, with L2E, it can be observed that the meta loss converges stably even with varying step size values.

Algorithm 2 InnerLoop

1: Input: Meta parameter \( \phi \), inner iterations \( N_{\text{inner}} \), initial parameter \( \theta_0 \), step size \( \epsilon \), burn-in steps \( B \), thinning interval \( T \).
2: Output: Loss \( L(\phi) \)
3: Initialize \( \Theta = \emptyset \) and \( r_0 \sim \mathcal{N}(0, I_d) \).
4: for \( i = 1, \ldots, N_{\text{inner}} \) do
5: \( r_{t+1} = r_t - \epsilon_t (\nabla \tilde{U}(\theta_t) + \alpha_\phi + c\beta_\phi) + \xi_t \) where \( \xi_t \sim \mathcal{N}(0, 2c) \).
6: \( \theta_{t+1} = \theta_t + \epsilon_t \beta_\phi \)
7: if \( i > B \land \mod(i, T) = 0 \) then
8: \( \Theta \leftarrow \Theta \cup \{\theta_i\} \)
9: end if
10: end for
11: \( L(\phi) \leftarrow -\log \frac{1}{|\Theta|} \sum_{\theta \in \Theta} p(y^*|x^*, \theta) \)

I. Details for meta-training

For meta-training, we construct our experiment code based on JAX learned-optimization package (Metz et al., 2022a). Also, we build our own meta-loss and L2E with specific input features and design choice. Therefore, we construct our own meta-learning task distribution that L2E can efficiently learn knowledge for better generalization for large-scale image classification task.

I.1. Input features

We use the following input features for L2E:

- raw gradient values
- raw parameter values
- raw momentum values
- running average of gradient values

Running average feature is expanded for multiple time scale in that we use multiple momentum-decay values for averaging. We use 0.1, 0.5, 0.9, 0.99, 0.999 and 0.9999 for momentum decay so that running average feature is expanded into 6-dimensions. Therefore, we have total 9-dimensional input features for each dimension of parameter and momentum. Input features are normalized so that \( l_2 \) norm with respect to input features of different dimensions become 1. \( \alpha_\phi \) and \( \beta_\phi \) share weights of neural network except for the last layer of MLP, so that we can get two quantities with single forward pass. This weight sharing method is employed in Levy et al. (2017) or other recent literature in learned optimization like in Metz et al. (2022b) and Metz et al. (2019).
I.2. Task distribution

**Dataset** We use MNIST, Fashion-MNIST, EMNIST and MedMNIST for meta training. We do not use resized version of dataset for meta training. For MedMNIST, we use the BloodMNIST in the official website.

**Neural network architecture** At each outer iteration, we randomly sample one configuration of neural network architecture which is constructed by the possible choice below. We use the following options for neural network architecture variation:

- Size of the convolution output channel : {4, 8, 16}
- Number of convolution layers: {1, 2, 3, 4, 5}
- Presence of the residual connection: boolean

![Diagram](image)

*Figure 10. Meta learning procedure of L2E.*

I.3. Meta-training procedure

**General hyperparameter** For meta-training, we fix the length of the inner loop for all tasks at 3000 iterations. We determine this by monitoring the meta-objective during training and set sufficient length of inner loop for L2E to enter to high-density region regardless of tasks. This setting can vary when task distribution is changed. To compute the meta-objective, we collect 10 inner-parameters with a thinning interval of 50 during the last 500 iterations of the inner-loop. We find out that this thinning interval and the number of collected inner parameters do not have a significant impact on the model performance. Also, we use 1000 iterations of outer-loop for training meta-parameter since meta-loss converges after 1000 outer iterations.

**Outer optimization** For training meta-parameter, we use Adam with learning rate 0.01 and $\beta_1 = 0.9, \beta_2 = 0.99$. We apply gradient clipping to the gradient of meta-objective to prevent unstable training due to the different gradient scale among tasks.

I.4. Meta-SGMC MC meta-training procedure

**General setup** For meta-training, we use the same task distribution as L2E do to fairly compare generalizability of two methods. We use 1-layer MLP with 10 hidden units for $Q_f$ and $D_f$. Since output of $D_f$ restricted to positive value, we take absolute value of output of $D_f$. We modify some original input preprocessing method different from Gong et al. (2018). Firstly, they heuristically find some good values to scale the input for $Q_f$ and $D_f$. This does not work in multi-task training as hyperparameter tuning is task specific, we scale the norm of input of neural networks as L2E do.

For meta-training, we use TBPTT as the gradient estimator for meta-parameter. We use total 5000 inner-steps as a single problem and use 50 steps for truncation length. After running 5000 steps for one task, we freshly sample the task for training meta-sgmc, which is completely different from Gong et al. (2018) and similar to L2E. Since Gong et al. (2018) used 15 steps for truncation length, we find out that lengthen the truncation length can improve the performance. Also, we tune the band width length and use 100 for stein gradient estimator.
Table 11. Results of ablation studies. We use 10 samples for ablation studies. \(L_2E\) shows the best performance among other choices. Notable result is that the performance gap between small \(L_2E\) gets larger as the size of task gets bigger. This implies that task distribution of meta-training plays important role for generalization capability. Precond \(L_2E\) shows worse performance in general and diverge in Tiny-ImageNet task. This parameterization shows worse performance and less generalizability than kinetic gradient parameterization.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>(L_2E)</th>
<th>ACC ↑</th>
<th>NLL ↓</th>
<th>ECE ↓</th>
<th>KLD ↑</th>
</tr>
</thead>
<tbody>
<tr>
<td>fashion-MNIST</td>
<td>Small (L_2E)</td>
<td>0.9158</td>
<td>0.2520</td>
<td>0.0120</td>
<td>0.0797</td>
</tr>
<tr>
<td></td>
<td>Precond (L_2E)</td>
<td>0.8801</td>
<td>0.3461</td>
<td>0.0382</td>
<td>0.2597</td>
</tr>
<tr>
<td></td>
<td>(L_2E)</td>
<td>0.9166</td>
<td>0.2408</td>
<td>0.0078</td>
<td>0.0507</td>
</tr>
<tr>
<td>CIFAR-10</td>
<td>Small (L_2E)</td>
<td>0.8952</td>
<td>0.3465</td>
<td>0.0610</td>
<td>0.5008</td>
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<tr>
<td></td>
<td>Precond (L_2E)</td>
<td>0.7503</td>
<td>0.7950</td>
<td>0.1349</td>
<td>0.4126</td>
</tr>
<tr>
<td></td>
<td>(L_2E)</td>
<td>0.9009</td>
<td>0.3231</td>
<td>0.0480</td>
<td>0.6263</td>
</tr>
<tr>
<td>CIFAR-100</td>
<td>Small (L_2E)</td>
<td>0.6536</td>
<td>1.3067</td>
<td>0.1237</td>
<td>1.4391</td>
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<td></td>
<td>Precond (L_2E)</td>
<td>0.3397</td>
<td>2.6761</td>
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<td>0.7449</td>
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<tr>
<td></td>
<td>(L_2E)</td>
<td>0.6702</td>
<td>1.2345</td>
<td>0.1224</td>
<td>1.2632</td>
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<tr>
<td>Tiny-ImageNet</td>
<td>Small (L_2E)</td>
<td>0.4759</td>
<td>2.3180</td>
<td>0.0672</td>
<td>2.8437</td>
</tr>
<tr>
<td></td>
<td>Precond (L_2E)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>(L_2E)</td>
<td>0.5287</td>
<td>1.9790</td>
<td>0.1478</td>
<td>0.8124</td>
</tr>
</tbody>
</table>

**Hyperparameters** For step size, we use 1e-06 since larger step size does not work in our task distribution for Meta-SGMCMC. Please refer to Figure 9 for sensitivity of Meta-SGMCMC compared to \(L_2E\). We use 5 for clipping values of \(Q_f\) and 100 for \(D_f\). For implementing meta-objective, we follow the official implementation of the meta-objective except for number of chains due to memory consumption. We use 2 chains for running Meta-SGMCMC on our task-distribution.

**Outer optimization** For training meta-parameter, we use Adam with learning rate 0.0003 and \(\beta_1 = 0.9, \beta_2 = 0.99\). We apply gradient clipping to the gradient of meta-objective to prevent unstable training due to the different gradient scale among tasks. We train meta-models for 1000 outer-iteration while saving the meta-models for every 100 outer iterations. Since training Meta-SGMCMC is significantly unstable as shown in Figure 9, we choose the best sampler by evaluating every saved meta-models for fashion-MNIST and CIFAR-10 tasks.

**J. Ablation studies**

In Table 11, we demonstrate the results from our ablations studies. We evaluate how the size of task set can affect the generalization performance of \(L_2E\) and the parameterization choice can make impact on the performance. We use following two variants of \(L_2E\):

- **Small \(L_2E\)**: meta-trained on only one dataset, using small architectures. In detail, this model only use Fashion-MNIST dataset and channel sizes of 4 and depths of 1 and 2 are possible choices of random configuration.

- **Precond \(L_2E\)**: Parameterization of \(L_2E\) changes from designing kinetic energy gradient to preconditioner.

Firstly, we confirm that \(L_2E\) trained with larger task distribution shows better generalization performance. Although small \(L_2E\) shows decent performance on Fashion-MNIST and CIFAR-10, it shows significant drop of performance in Tiny-ImageNet and CIFAR-100. This implies that further scaling of task distribution can lead \(L_2E\) to show better performance in diverse unseen tasks. Also, we compare the method parameterizing \(D(z), Q(z)\) with neural networks as in Gong et al. (2018). This shows that our parameterization is significantly better than designing preconditioner. Since preconditioner matrix can only work as multiplier of learning rate or noise scale, it has limitation for its expressivity which shows limitation in BNNs.

**K. Experimental Details**

We use JAX library to conduct our experiments. We use NVIDIA RTX-3090 GPU with 24GB VRAM and NVIDIA RTX A6000 with 48GB for all experiments. We implement meta-training algorithm based on the learned-optimization
Learning to Explore for Stochastic Gradient MCMC

Table 12. Hyperparameters for DE.

<table>
<thead>
<tr>
<th>Method</th>
<th>fashion-MNIST</th>
<th>CIFAR-10</th>
<th>CIFAR-100</th>
<th>Tiny-ImageNet</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optimizer</td>
<td>SGDM</td>
<td>SGDM</td>
<td>SGDM</td>
<td>SGDM</td>
</tr>
<tr>
<td>Num models</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>10</td>
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<tr>
<td>Total epochs</td>
<td>10000</td>
<td>10000</td>
<td>10000</td>
<td>2000</td>
</tr>
<tr>
<td>Initial learning rate</td>
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<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
</tr>
<tr>
<td>Learning rate schedule</td>
<td>Cosine decay</td>
<td>Cosine decay</td>
<td>Cosine decay</td>
<td>Cosine decay</td>
</tr>
<tr>
<td>momentum decay</td>
<td>0.5</td>
<td>0.1</td>
<td>0.05</td>
<td>0.3</td>
</tr>
<tr>
<td>Weight decay</td>
<td>$5 \times 10^{-4}$</td>
<td>$1 \times 10^{-3}$</td>
<td>$5 \times 10^{-4}$</td>
<td>$5 \times 10^{-4}$</td>
</tr>
<tr>
<td>Batch size</td>
<td>128</td>
<td>128</td>
<td>128</td>
<td>128</td>
</tr>
</tbody>
</table>

Table 13. Hyperparameters for CSGMCMC.

<table>
<thead>
<tr>
<th>Method</th>
<th>fashion-MNIST</th>
<th>CIFAR-10</th>
<th>CIFAR-100</th>
<th>Tiny-ImageNet</th>
</tr>
</thead>
<tbody>
<tr>
<td>Num burn in epochs</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Num models</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>10</td>
</tr>
<tr>
<td>Total epochs</td>
<td>5100</td>
<td>5100</td>
<td>5100</td>
<td>1100</td>
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<tr>
<td>Thinning interval</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>exploration ratio</td>
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<td>0.94</td>
<td>0.94</td>
<td>0.8</td>
</tr>
<tr>
<td>Step size</td>
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<td>$1 \times 10^{-6}$</td>
<td>$1 \times 10^{-6}$</td>
<td>$1 \times 10^{-6}$</td>
</tr>
<tr>
<td>Step size schedule</td>
<td>Cosine</td>
<td>Cosine</td>
<td>Cosine</td>
<td>Cosine</td>
</tr>
<tr>
<td>Momentum decay</td>
<td>0.5</td>
<td>0.01</td>
<td>0.01</td>
<td>0.05</td>
</tr>
<tr>
<td>Weight decay</td>
<td>$5 \times 10^{-4}$</td>
<td>$1 \times 10^{-3}$</td>
<td>$1 \times 10^{-3}$</td>
<td>$5 \times 10^{-4}$</td>
</tr>
<tr>
<td>Batch size</td>
<td>128</td>
<td>128</td>
<td>128</td>
<td>128</td>
</tr>
</tbody>
</table>

package (Metz et al., 2022a) with some modifications.

K.1. Real-world image classification

Dataset We use tensorflow dataset for fashion-MNIST, CIFAR-10 and CIFAR-100. We utilize Tiny-ImageNet with image size of 64x64.

Architecture We use 1 layer convolution neural network on fashion-MNIST and ResNet56-FRN with Swish activation on Tiny-ImageNet.

Hyperparameter We report hyperparameters for each methods in Table 12, Table 13 and Table 14. We tune the hyperparameters of methods using BMA NLL as criterion with number of 10 samples. For all methods including $L^2_E$, we tune learning rate(step size), weight decay(prior variance) and momentum decay term. We use zero-mean gaussian as prior distribution for SGMCMC methods, so that prior variance is equal to half of the inverse of weight decay. For momentum decay, we grid search over $\alpha \in \{0.01, 0.05, 0.1, 0.3, 0.5\}$. For weight decay, we also search over $\lambda \in \{1e-04, 5e-04, 1e-03\}$ to find best configuration. For step size, except for $L^2_E$, we search over $\epsilon \in \{2e-7, 4e-7, 1e-6, 2e-6, 4e-6, 1e-5\}$. For $L^2_E$, due to scale of output of meta-learner, we additionally search over bigger step size including $2e-5, 1e-4, 2e-4$.

Baselines We use the following as baseline methods

- Deep ensembles (Lakshminarayanan et al., 2017): This method collects parameters trained from multiple different initialization for ensembling. DE is often compared with Bayesian methods in recent BNNs literature like in Izmailov et al. (2021b) in that DE induce similar function to HMC which is golden standard in BNNs with BMA.

- Cyclical Stochastic Gradient MCMC (Zhang et al., 2020): Zhang et al. (2020) introduced cyclic learning rate schedule to SGMCMC for improving exploration of sampler. CSGMCMC usually shows descent predictive performance comparing to other existing SGMCMC methods in large-scale experiments.
Learning to Explore for Stochastic Gradient MCMC

Table 14. Hyperparameters for $l_2$E.

<table>
<thead>
<tr>
<th>Method</th>
<th>fashion-MNIST</th>
<th>CIFAR-10</th>
<th>CIFAR-100</th>
<th>Tiny-ImageNet</th>
</tr>
</thead>
<tbody>
<tr>
<td>Num burn in epochs</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Num models</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>10</td>
</tr>
<tr>
<td>Total epochs</td>
<td>5100</td>
<td>5100</td>
<td>5100</td>
<td>1100</td>
</tr>
<tr>
<td>Thinning interval</td>
<td>50</td>
<td>50</td>
<td>50</td>
<td>20</td>
</tr>
<tr>
<td>Step size schedule</td>
<td>Constant</td>
<td>Constant</td>
<td>Constant</td>
<td>Constant</td>
</tr>
<tr>
<td>Momentum decay</td>
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<td>0.05</td>
<td>0.05</td>
<td>0.05</td>
</tr>
<tr>
<td>Weight decay</td>
<td>$5 \times 10^{-4}$</td>
<td>$5 \times 10^{-4}$</td>
<td>$5 \times 10^{-4}$</td>
<td>$5 \times 10^{-4}$</td>
</tr>
<tr>
<td>Batch size</td>
<td>128</td>
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<tr>
<td>Step size</td>
<td>$2 \times 10^{-5}$</td>
<td>$2 \times 10^{-5}$</td>
<td>$1 \times 10^{-4}$</td>
<td>$2 \times 10^{-4}$</td>
</tr>
</tbody>
</table>

- **Meta-SGMCMC** (Gong et al., 2018): Gong et al. (2018) proposed to meta-learn curl and diffusion matrix of SGMCMC to build sampler that can fastly converge to target distribution with small bias.

**Metrics**

Let $p(y|x, \theta) \in [0, 1]^K$ be a predicted probabilities for a given input $x$ with label $y$ and $\theta$ is model parameter. $p^{(k)}$ denotes the $k$th element of the probability vector. We have the following common metrics on the dataset $D$ consists of inputs $x$ and labels $y$:

- **Accuracy (ACC):**
  \[
  \text{ACC}(D) = \mathbb{E}_{(x,y) \in D} \left[ I \left[ y = \arg \max_k p^{(k)}(x) \right] \right].
  \]  (15)

- **Negative log-likelihood (NLL):**
  \[
  \text{NLL}(D) = \mathbb{E}_{(x,y) \in D} \left[ -\log p^{(y)}(x) \right].
  \]  (16)

- **Expected calibration error (ECE):** Actual implementation of Expected Calibration Error (ECE) includes dividing predicted probabilities with their confidence. We use following implementation
  \[
  \text{ECE}(D, N_{\text{bin}}) = \sum_{b=1}^{N_{\text{bin}}} \frac{n_b |\delta_b|}{n_1 + \cdots + n_{N_{\text{bin}}}},
  \]  (17)
  where $N_{\text{bin}}$ is the number of bins, $n_b$ is the number of examples in the $b$th bin, and $\delta_b$ is the calibration error of the $b$th bin. We use $N_{\text{bin}} = 15$ for computing ECE.

- **Pairwise Kullback-Leibler Divergence (KLD):** Given set of ensemble members $\Theta = \{\theta_1, \ldots, \theta_M\}$, we construct matrix $A$ using pair of ensemble members which have $A_{ij} = p(y|x, \theta_i) \log \frac{p(y|x, \theta_i)}{p(y|x, \theta_j)}$ for a given input $x$ and $y$. We calculate the statistic as follows
  \[
  D_{\text{KL}}(D, M) = \mathbb{E}_{(x,y) \in D} \left[ \frac{1}{M(M-1)} \sum_{i \neq j} A_{ij} \right].
  \]  (18)

- **Agreement:** The Agreement is defined as the alignment between the top-1 predictions of the HMC and our own predictions. This metric is computed through the following formula:
  \[
  \text{Agreement}(D) = \mathbb{E}_{(x,y) \in D} \left[ I \left[ \arg \max_k \hat{p}^{(k)}(x) = \arg \max_k p^{(k)}(x) \right] \right],
  \]  (19)
  where $I[\cdot]$ is indicator function and $\hat{p}$ is predictive distribution of HMC. It indicates how well a method is able to capture the top-1 predictions of HMC.
Learning to Explore for Stochastic Gradient MCMC

Table 15. Wall clock time(sec) of SGMCMC methods per single epoch. Measured using NVIDIA RTX A6000.

<table>
<thead>
<tr>
<th>Method</th>
<th>fashion-MNIST</th>
<th>CIFAR-10</th>
<th>CIFAR-100</th>
<th>Tiny-ImageNet</th>
</tr>
</thead>
<tbody>
<tr>
<td>CSGMCMC</td>
<td>0.20</td>
<td>1.79</td>
<td>1.81</td>
<td>41.12</td>
</tr>
<tr>
<td>Meta-SGMCMC</td>
<td>0.32</td>
<td>2.44</td>
<td>2.45</td>
<td>47.22</td>
</tr>
<tr>
<td>L2E</td>
<td>0.32</td>
<td>2.64</td>
<td>2.66</td>
<td>48.43</td>
</tr>
</tbody>
</table>

• Total Variation(Total Var): Total Variation quantifies the total variation distance between the predictive distribution of the HMC and our own predictions averaged over the data points. Specifically, it compares the predictive probabilities for each of the classes as follows

\[
\text{TotalVar}(\mathcal{D}) = \mathbb{E}_{(x,y) \in \mathcal{D}} \left[ \frac{1}{2} \sum_k \left| \tilde{p}^{(k)}(x) - p^{(k)}(x) \right| \right].
\]

(20)

K.2. Computational cost

Table 15 shows the actual computational time for each methods. L2E does not incur significant computational overhead.

L. Implementation of Convergence Diagnostics

For Effective Sample Size (ESS), we use Tensorflow Probability (Lao et al., 2020) library for implementation. We use default parameter of the implementation. ESS is then normalized to the time consumed for running one thinning interval, which is equivalent to 50 epochs. Since scale of ESS is too large to report since dimension of neural network parameters are huge for our experiments, we divide it by \(10^{-5}\) and report for convenience.

\(\hat{R}\) (Gelman & Rubin, 1992) has been widely employed to measure the convergence of MCMC chain. However, in modern BNNs, due to the multimodality in BNNs posterior distribution, traditional \(\hat{R}\) without further modification is meaningless in the parameter space(Sommer et al., 2024). Sommer et al. (2024) demonstrated that parameter space convergence should be measured both chain- and layer-wise to fix these issues. Basically, \(c\hat{R}_\psi^{(\kappa)}\) diagnostic splits a single chain’s path into \(\kappa\) sub chains for \(\hat{R}\) calculation. Therefore, we report the proportion of parameters with \(c\hat{R}_\psi^{(\kappa)} < 1.1\) using code implementation of Sommer et al. (2024) for our main results. We set \(\kappa = 2\) considering the number of collected parameters in our experiments. Please refer to Sommer et al. (2024) for detailed description of this metric. In Figure 11, we also demonstrate layer-wise visualization of \(c\hat{R}_\psi^{(\kappa)}\). For weights, L2E shows good convergence performance in terms of layer-wise \(c\hat{R}_\psi^{(\kappa)}\).
while convergence of biases is mixed across the layers. Considering that weights account for the majority of the parameters, we confirm that \( L^2E \) shows strong convergence in terms of layer-wise \( \hat{cR}^{(\kappa)}_\psi \).

**M. Limitation of \( L^2E \)**

While effective, \( L^2E \) has some limitations that should be considered for applications. Firstly, \( L^2E \) requires additional meta-training costs. In our experiment, meta-training takes approximately 6 hours on a single NVIDIA RTX A6000 GPU, but it possibly requires more computational cost for meta-training using larger tasks. While \( L^2E \) demonstrates good performance across various data domains, in Table 11, we observe that the performance of \( L^2E \) is influenced by the size of the meta-training distribution as the scale of the target problem increases. In other words, there is a possibility that we may need to meta-train \( L^2E \) with larger datasets and architectures to apply \( L^2E \) to very large models and datasets.