High-Order Stochastic Gradient Thermostats for Bayesian Learning of Deep Models

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Abstract

Learning in deep models using Bayesian methods has generated significant attention recently. This is largely because of the feasibility of modern Bayesian methods to yield scalable learning and inference, while maintaining a measure of uncertainty in the model parameters. Stochastic gradient MCMC algorithms (SG-MCMC) are a family of diffusion-based sampling methods for large-scale Bayesian learning. In SG-MCMC, multivariate stochastic gradient thermostats (mSGNHT) augment each parameter of interest, with a momentum and a thermostat variable to maintain stationary distributions as target posterior distributions. As the number of variables in a continuous-time diffusion increases, its numerical approximation error becomes a practical bottleneck, so better use of a numerical integrator is desirable. To this end, we propose use of an efficient symmetric splitting integrator in mSGNHT, instead of the traditional Euler integrator. We demonstrate that the proposed scheme is more accurate, robust, and converges faster. These properties are demonstrated to be desirable in Bayesian deep learning. Extensive experiments on two canonical models and their deep extensions demonstrate that the proposed scheme improves general Bayesian posterior sampling, particularly for deep models.

1 Introduction

The ability to learn abstract representations that support generalization to novel instances lies at the core of many problems in machine learning and computer vision. Human learners often can grasp concepts at multiple levels of abstraction from training examples, and make meaningful generalizations (Xu and Tenenbaum 2007; Kemp, Perfors, and Tenenbaum 2007). Intuitively, appropriately employed prior knowledge and hierarchical reasoning are necessary in this task. Bayesian learning and inference applied to deep models may naturally possess such characterization, and potentially could take a step towards this ability (Salakhutdinov, Tenenbaum, and Torralba 2013).

Deep models come in two broad categories. The first uses stochastic hidden layers, typically deep latent variable models. This includes the deep sigmoid belief network (Mnih and Gregor 2014; Gan et al. 2015b), the variational auto-encoder (Kingma and Welling 2014), and many others (Ranganath et al. 2015; Gan et al. 2015a; Pu, Yuan, and Carin 2015). The second category of deep models uses deterministic hidden layers. While the stochastic hidden units of the first category make this class of models naturally amenable to Bayesian learning (see the above references), for the second category appropriate priors on the weights of networks may be employed to consider weight uncertainty. Previous work has applied Bayesian methods to neural networks (MacKay 1992; Neal 1995), including feedforward neural networks (Blundell et al. 2015; Hernández-Lobato and Adams 2015; Korattikara et al. 2015) and convolutional neural networks (Gal and Ghahramani 2015). Deep learning may often be interpreted as a stacking of such neural networks.

Bayesian learning and inference methods have generated significant recent research activity. Stochastic gradient Markov Chain Monte Carlo (SG-MCMC) methods (Welling and Teh 2011; Chen, Fox, and Guestrin 2014; Ding et al. 2014; Li et al. 2016) are a family of Itô diffusion based algorithms that can efficiently sample target distributions, and can be applied to large datasets. In these algorithms, two approximations are made (Chen, Ding, and Carin 2015). (i) For practical scalability, stochastic gradients from mini-batches of data are used to estimate the true gradient; (ii) For numerical feasibility, a numerical integration with small step is used to solve the corresponding Itô diffusion (a continuous-time Markovian process).

The first attempt at SG-MCMC was the Stochastic Gradient Langevin Dynamics (SGLD) (Welling and Teh 2011). It is based on 1st-order Langevin dynamics. The Stochastic Gradient Hamiltonian Monte Carlo (SGHMC) (Chen, Fox, and Guestrin 2014) extends SGLD with 2nd-order Langevin dynamics, where momentum variables are introduced into the system. In an attempt to address the problem of estimating stochastic gradient noise, the Stochastic Gradient Noé-Hoover Thermostat (SGNHT) (Ding et al. 2014) was proposed, with one additional global thermostat variable. To further improve the efficiency of the SGNHT, a multivariate version of SGNHT (mSGNHT) was proposed by in-
Introducing multiple thermostat variables instead of a single one (Ding et al. 2014). It was shown that mSGNHT provides more adaptivity than SGHMT (Gan et al. 2015a).

By examining training with SG-MCMC algorithms, we note two issues. (i) as more variables are introduced, an accurate numerical method becomes more critical; and (ii) gradients in deep models often suffer from the vanishing/exploding problem (Bengio, Simard, and Frasconi 1994), which makes choosing a proper stepsize difficult in SG-MCMC. In this paper, we mitigate these concerns by utilizing a more accurate numerical integrator, the symmetric splitting integrator (SSI), to reduce discretization errors in mSGNHT. Furthermore, since SSI is more robust with respect to stepsizes than the default Euler integrator, it allows one to choose an appropriate stepsize much more easily. We justify that the Euler integrator used in mSGNHT is 1st-order, while the SSI is 2nd-order. Borrowing tools from (Chen, Ding, and Carin 2015), we show that mSGNHT with SSI (mSGNHT-S) converges faster and more accurately than mSGNHT with a Euler integrator (mSGNHT-E). Experiments across a wide range of model types demonstrate the utility of this method. Specifically, we consider latent Dirichlet allocation, logistic regression, deep neural networks and deep Poisson factor analysis.

2 Background

2.1 Itô Diffusion

Itô diffusion is a stochastic differential equation (SDE) defined as:

$$dX_t = F(X_t)dt + \sigma(X_t)dW_t,$$

where $X_t \in \mathbb{R}^n$, $W_t$ is Brownian motion, and $t$ is the time index. Functions $F: \mathbb{R}^n \rightarrow \mathbb{R}^n$ and $\sigma: \mathbb{R}^n \rightarrow \mathbb{R}^{n \times n}$ are assumed to satisfy the usual Lipschitz continuity condition (Knapp 2005). It has been shown that by designing appropriate functions $F$ and $\sigma$, the stationary distribution, $\rho(X)$, of the Itô diffusion (1) has a marginal distribution that is equal to the posterior distribution of interest (Chen, Ding, and Carin 2015; Ma, Chen, and Fox 2015).

To formulate mSGNHT (Gan et al. 2015a; Ding et al. 2014) into the Itô diffusion (1), let $X = (\theta, p, \xi)$, where $\theta \in \mathbb{R}^n$ are the model parameters, $p \in \mathbb{R}^n$ are momentums, and $\xi \in \mathbb{R}^n$ represent the thermostats (Gan et al. 2015a). Define $U \triangleq -\left(\sum_{i=1}^{N} \log p(d_i|\theta) + \log p(\theta)\right)$ as the unnormalized negative log-posterior, where $\{d_i\}$ represents the $i$th data sample, $p(d_i|\theta)$ the corresponding likelihood, and $p(\theta)$ the prior. For some constant $D > 0$, the mSGNHT in (Gan et al. 2015a) is shown to be a form of Itô diffusion, with

$$F = \begin{bmatrix} p \\ p \odot - \nabla_{\theta} U \\ p \odot - 1 \end{bmatrix}, \quad \sigma = \sqrt{2D} \begin{bmatrix} 0 & 0 & 0 \\ 0 & I_n & 0 \\ 0 & 0 & 0 \end{bmatrix},$$

where $\odot$ represents element-wise product, and $I_n$ is the $n \times n$ identity matrix. Based on the Fokker-Planck equation (Risken 1989), the marginal stationary distribution over $\theta$ can be shown to be $\rho(\theta) \propto \exp(-U(\theta))$, the posterior distribution we are interested in.

2.2 Euler Integrator

The continuous-time diffusion in (1) cannot be solved explicitly in general. As a result, numerical methods are required in SG-MCMCs to generate approximate samples. The standard numerical method used in SG-MCMC is the Euler integrator, which generates samples sequentially from a discrete-time approximation of (1). Specifically, conditioned on the current sample $X_t$ and step size $h$, the next sample at time $t+1$ is generated via the rule:

$$X_{t+1} = X_t + F(X_t)h + \sigma(X_t)\xi_{t+1}, \xi_{t+1} \sim N(0, hI_n).$$

In the case of mSGNHT, in each step, a stochastic gradient from a minibatch is used instead of the full gradient. We thus approximate $U$ with $\tilde{U}_t \triangleq -\left(\sum_{i \in S_t} \log p(d_i|\theta) + \log p(\theta)\right)$ for the $t$-th iteration, where $S_t \subseteq \{1, 2, \cdots, N\}$, and $|\cdot|$ is the cardinality of a set. This results in the following sampling rules:

$$\begin{cases}
\theta_{t+1} = \theta_t + p_t h \\
p_{t+1} = p_t - \nabla_{\theta} \tilde{U}_t(\theta_{t+1})h - \text{diag}(\xi_t) p_t h + \sqrt{2D} \xi_{t+1} \\
\xi_{t+1} = \xi_t + (p_{t+1} \odot p_{t+1} - 1) h 
\end{cases}$$

3 Symmetric Splitting Integrator for mSGNHT

The SSI has been studied in statistical physics (Leimkuhler and Matthews 2013; Leimkuhler and Shang 2015). It generalizes the idea of the leap-frog integrator used in the Hamiltonian Monte Carlo (Neal 2011) from the partial differential equation setting to the SDE setting. It was not until recently that SSI was introduced into machine learning to obtain a more accurate SGHMC algorithm (Chen, Ding, and Carin 2015). We adopt the idea and generalize it in this paper for mSGNHT.

The idea of SSI is to split the intractable SDE, i.e., (1), into several sub-SDEs such that each sub-SDE can be solved analytically. For the mSGNHT represented in (2), it is readily split into the following sub-SDEs:

$$A: \begin{cases}
d\theta = pdt \\
dp = -\xi \odot pdt, \\
d\xi = 0 
\end{cases}, \quad B: \begin{cases}
d\theta = 0 \\
dp = -\nabla_{\theta} \tilde{U}_t(\theta) dt + \sqrt{2D} dW. \\
d\xi = 0 
\end{cases}$$

All the sub-SDEs can be solved analytically, leading to the following rules to generate samples $\{\theta_{t+1,1}, p_{t+1,1}, \xi_{t+1,1}\}$ from mSGNHT for time $t+1$:

$$\begin{cases}
A: \theta_{t+1/2} = p_{t+1/2}h/2, \xi_{t+1/2} = \xi_t + (p_t \odot p_{t-1}) h/2 \rightarrow \\
B: p_{t+1/3} = \exp(-\xi_{t+1/2}h/2) \odot p_t \\
O: p_{t+2/3} = p_{t+1/3} - \nabla_{\theta} \tilde{U}_t(\theta_{t+1/2}) h + \sqrt{2D} \xi_{t+1/2} \rightarrow \\
B: p_{t+1} = \exp(-\xi_{t+1/2}h/2) \odot p_{t+2/3} \\
A: \theta_{t+1} = p_{t+1}h/2, \xi_{t+1} = \xi_{t+1/2} + (p_{t+1} \odot p_{t+1} - 1) h/2 
\end{cases}$$

\footnote{We write $\tilde{F}(X_t)$ from (2) as $\tilde{F}(X_t)$ if a stochastic gradient is used in the rest of the paper.}
From the update equations, SSI performs almost as efficiently as the Euler integrator. Furthermore, the splitting scheme for (2) is not unique. However, all of the schemes can be shown to have the same order of accuracy. In the following subsection, we show quantitatively that the SSI is more accurate than the Euler integrator in terms of approximation errors. To get an impression of how the SSI works, we illustrate it with a simple synthetic experiment.

**Illustrations with a Double-well Potential** To illustrate the proposed symmetric splitting scheme and its robustness to stepsize, fast convergence, and accurate parameter approximation, we follow (Ding et al. 2014), and consider the double-well potential with

\[ U(\theta) = (\theta + 4)(\theta + 1)(\theta - 1)(\theta - 3)/14 + 0.5, \]

and the target distribution \( \rho(\theta) \propto \exp(-U(\theta)) \). The unknown noise in the stochastic gradient is simulated as \( \nabla U(\theta)h = \nabla U(\theta)h + \mathcal{N}(0, 0.2h) \), where \( B = 1 \). No injecting noise is added. We examine a large range of stepsize \( h \) from \( 10^{-3} \) to 0.3.

In Fig. 1, we plot the KL divergences between the true distributions and the estimated density, based on \( 10^6 \) samples, using two types of integrators. mSGNHT-S consistently provides a better approximation. The significant gap at larger stepsize reveals that mSGNHT-S allows large updates.

Furthermore, we visualize the results of the first \( 10^5 \) samples for \( h = 10^{-3} \) and \( h = 0.2 \) in Fig. 2. When the stepsize is too small (\( h = 10^{-3} \)), conventional mSGNHT-E has not explored the whole parameter space; this is because it converges slower, as shown later. When the stepsize is large (\( h = 0.2 \)), large numerical error is potentially brought in, and mSGNHT-E over-concentrates on the mode. In both cases, mSGNHT-S approaches the theoretical value of thermostat variable \( \xi = 1 \) more accurately.

### 3.1 Theoretical Justification

In (Chen, Ding, and Carin 2015), the authors formally studied the roles of numerical integrators in general SG-MCMCs. We adopt their framework, and justify the advantage of the proposed scheme for mSGNHT. We first define the local generator of the SDE (1) at the \( t \)-th iteration (i.e., replacing the full gradient with the stochastic gradient from the \( t \)-th minibatch) as:

\[ \tilde{L}_t f(X_t) \triangleq \left( \tilde{F}_t(X_t) \cdot \nabla X + \frac{1}{2} \left( \sigma(X_t)\sigma(X_t)^T \right) : \nabla X \nabla X^T \right) f(X_t) \]

where \( a \cdot b \triangleq a^T b, A : B \triangleq \text{tr}(A^T B), f : \mathbb{R}^n \rightarrow \mathbb{R} \) is any twice differentiable function. Based on the definition, according to the Kolmogorov backward equation, we have

\[ \mathbb{E}[f(X)] = e^{h\tilde{L}_t} f(X) \]

where the expectation is taken over the randomness in the diffusion. The operator \( e^{h\tilde{L}_t} \) is called the Kolmogorov operator. Because a numerical integrator is adopted to solve the original SDE, the resulting Kolmogorov operator, denoted as \( \tilde{P}^h_t \), approximates \( e^{h\tilde{L}_t} \). To characterise the accuracy of a numerical integrator, we use the following definition.

**Definition 1** A numerical integrator is said to be a \( K \)-th order local integrator if for any smooth and bounded function \( f \), the corresponding Kolmogorov operator \( \tilde{P}^h_t f(x) \) from the \( t \)-th minibatch with stepsize \( h \) satisfies the following relation:

\[ \tilde{P}^h_t f(x) = e^{h\tilde{L}_t} f(x) + O(h^{K+1}) \quad (4) \]

We follow (Chen, Ding, and Carin 2015), and state in Lemma 1 that a Euler integrator satisfies Definition 1 with \( K = 1 \) when used in mSGNHT. Detailed proofs are provided in the Appendix.

**Lemma 1** The Euler integrator used in mSGNHT-E is a 1st-order local integrator, i.e.,

\[ \tilde{P}^h_t = e^{h\tilde{L}_t} + O(h^2) \quad (5) \]

Using the Baker–Campbell–Hausdorff formula for commutators (Rossmann 2002), the SSI can be shown to be a 2nd-order integrator in mSGNHT, stated in Lemma 2.

**Lemma 2** The symmetric splitting integrator used in mSGNHT-S is a 2nd-order local integrator, i.e.,

\[ \tilde{P}^h_t = e^{h\tilde{L}_t} + O(h^3) \quad (6) \]
The authors of (Chen, Ding, and Carin 2015) formalize the role of numerical integrators in terms of posterior bias and mean square error (MSE). Specifically, for a testing function \( \phi(x) \), they study the difference between the posterior average \( \bar{\phi} = \frac{1}{T} \sum_{t=1}^{T} \phi(x_t) \), where \( \rho(x) \) denotes the true posterior of a Bayesian model, and \( \{x_t\} \) denotes samples from a SG-MCMC algorithm. To study the role of the SSI applied in mSGNHT, we simplify the notation and conclude their results in the following lemma.

**Lemma 3 (Roles of numerical integrators)** Under certain assumptions, the Bias and MSE of a SG-MCMC algorithm with stepsize \( h \) and a \( K \)-th order integrator are:

\[
\text{Bias: } \mathbb{E}(\hat{\phi} - \bar{\phi}) = B_{\text{bias}} + O(h^K)
\]

\[
\text{MSE: } \mathbb{E}((\hat{\phi} - \bar{\phi})^2) = B_{\text{mse}} + O(h^{2K})
\]

where \( B_{\text{bias}} \) and \( B_{\text{mse}} \) are functions depending on \( (h, T) \) but independent of \( K \).

Based on Lemma 3 and (Chen, Ding, and Carin 2015), we summarize the properties of mSGNHT-S in the following remarks. The detailed are provided in Appendix.

**Remark 1 (Robustness)** When applying Lemma 3 to mSGNHT, the bias and MSE of mSGNHT-S is bounded as:

\[
B_{\text{bias}} = O(h^2) \quad \text{and} \quad B_{\text{mse}} = O(h^4),
\]

compared to \( B_{\text{bias}} = O(h) \) and \( B_{\text{mse}} = O(h^2) \) for the mSGNHT-E, respectively. This indicates that mSGNHT-S is more robust to the stepsizes than mSGNHT-E.

**Remark 2 (Convergence Rate)** The higher order a numerical integrator is, the faster its optimal convergence rate is. Convergence rates in term of bias for mSGNHT-S and mSGNHT-E are \( T^{-2/3} \) and \( T^{-1/2} \), respectively, indicating mSGNHT-S converges faster.

**Remark 3 (Measure Accuracy)** In the limit of infinite time \((T \to \infty)\), the terms \( B_{\text{bias}} \) and \( B_{\text{mse}} \) in Lemma 3 vanish, leaving only the \( O(h^K) \) terms. This indicates mSGNHT-S is an order of magnitude more accurate than mSGNHT-E.

### 3.2 Advantages of mSGNHT-S for Deep Learning

Compared to optimization-based methods (Martens 2010), the mSGNHT-S is able to more fully explore the parameter space. Therefore, it is less sensitive to initialization, a nontrivial issue in optimization (Sutskever et al. 2013). Second, the mSGNHT is related to stochastic gradient descent (SGD) with momentum in optimization (Chen, Fox, and Guestrin 2014; Ding et al. 2014), with the additional advantage that the momentum is updated element-wise to automatically adapt stepsizes. Additionally, (Sutskever et al. 2013) shows that momentum-accelerated SGD is capable of accelerating along directions of low-curvature in the parameter space, leading to faster convergence speed. As a result, mSGNHT is more favorable than other momentum-free SG-MCMC algorithms, such as the “vanilla” SGLD (Welling and Teh 2011).

Specifically for mSGNHT, we know from previous analysis that (i) mSGNHT-S is less sensitive to stepsize as shown in Remark 1, and thus can tolerate gradients of various magnitudes. This provides a potential solution to mitigate the vanishing/exploding gradients problem (Rumelhart, E., and Williams 1986). Our empirical results on deep neural networks verifies this in Section 5.2. (ii) Convergence speed is a critical criteria for learning, and mSGNHT-S clearly outperforms mSGNHT-E in this regard as discussed in Remark 2. (iii) mSGNHT-S converges to a solution one order magnitude more accurate than mSGNHT-E, as discussed in Remark 3, and it is more accurate in estimating model parameters. The number of parameters in large-scale models may be significant, and small numerical error in individual parameters can accumulate, causing noticeable inefficiency. For these reasons, we advocate mSGNHT-S for training large deep models.

### 4 Related Work

One direction for scalable Bayesian learning of deep models is stochastic variational inference. For deep models with stochastic hidden units, learning has been implemented using variational methods; when the latent variables are continuous, Stochastic Gradient Variational Bayes (SGVB) (Kingma and Welling 2014) has been employed, while models with discrete latent variables have been trained via Neural Variational Inference and Learning (NVIL) (Mnih and Gregor 2014). For deep models with deterministic hidden units, recent studies have shown that imposing uncertainty on global parameters helps prevent overfitting, yielding significant improvement on model performances. Representative works of this type include Bayes by Backprop (BBB) (Blundell et al. 2015) and Probabilistic Backpropagation (PBP) (Hernández-Lobato and Adams 2015), which approximate posteriors of network weights as a product of univariate Gaussian distributions.

Another direction for Bayesian deep learning is SG-MCMC, the line of work followed by this paper. These methods do not have to assume a simplifying form for the posterior, as in variational methods. The difference with respect to optimization methods is the injection of Gaussian noise in the parameter update, allowing better exploration of parameter space when learning. Works of this type include the SGLD (Welling and Teh 2011), SGHMC (Chen, Fox, and Guestrin 2014), SGHT (Ding et al. 2014), and mSGNHT (Gan et al. 2015a). It has been shown in (Sutskever et al. 2013) that carefully tuned momentum methods suffice for dealing with curvature issue in deep network training. mSGNHT belongs to the class of momentum-accelerated SG-MCMC algorithms. In terms of numerical integrators, recently work for HMC include (Chao et al. 2015; Shahbaba et al. 2014). For our SDE setting, (Leimkuhler and Wollust 2014) proposes a symmetric splitting scheme for the SGNHT with a specific stochastic gradient noise, which is different from our setting of mSGNHT for deep models. Recently, (Chen, Ding, and Carin 2015) provides a theoretical foundation for rigorous study of numerical integrators for SG-MCMC. Our work is complementary, providing implementation guidance for the numerical integrator for other SG-MCMC, and investigating its roles in practical applications.
5 Experiments

5.1 Canonical Models

We consider two representative Bayesian models to demonstrate that mSGNHT-S improves general posterior sampling:

Latent Dirichlet Allocation (LDA) (Blei, Ng, and Jordan 2003) for latent variable models, and logistic regression, and testing data consist of a9a respectively, with parameter dimension.

Latent Dirichlet Allocation We first evaluate our method on the ICML dataset (Chen et al. 2015) using LDA. This dataset contains 765 documents from the abstracts of ICML proceedings from 2007 to 2011. After removing stopwords, we obtained a vocabulary size of 1918 and total words of 44140. We used 80% of the documents (selected at random) for training and the remaining 20% for testing. Similar to (Patterson and Teh 2013), we used the semi-collapsed LDA whose posterior is provided in the Appendix. Following (Ding et al. 2014), a Gaussian prior \( \mathcal{N}(0, 1) \) is used for the reparameterized parameter. The Dirichlet prior parameter for topic distribution for each document is set to 0.1. The number of topics is set to 30. We use perplexity (Blei, Ng, and Jordan 2003) to measure the quality of algorithms.

Table 1: LDA on ICML.

<table>
<thead>
<tr>
<th>Method</th>
<th>Test Perplexity ↓</th>
<th>Test Accuracy ↑</th>
</tr>
</thead>
<tbody>
<tr>
<td>SGNHT-S</td>
<td>939.67</td>
<td>84.95%</td>
</tr>
<tr>
<td>SGNHT-E</td>
<td>960.56</td>
<td>84.72%</td>
</tr>
<tr>
<td>SGHMC-S</td>
<td>1004.73</td>
<td>84.56%</td>
</tr>
<tr>
<td>SGHMC-E</td>
<td>1017.51</td>
<td>84.51%</td>
</tr>
<tr>
<td>SGRLD</td>
<td>1154.68</td>
<td>84.80%</td>
</tr>
<tr>
<td>Gibbs</td>
<td>907.84</td>
<td>84.94%</td>
</tr>
</tbody>
</table>

Table 2: LR on a9a.

<table>
<thead>
<tr>
<th>Method</th>
<th>Test Perplexity ↓</th>
<th>Test Accuracy ↑</th>
</tr>
</thead>
</table>

To show the robustness of mSGNHT-S to stochastic gradient noise, we chose minibatch of size 5, and \( D \) in mSGNHT-S is fixed as 0.75. We test a wide range of values for step size \( h \). Generally, larger \( h \) imposes larger gradient-estimation error and numerical error. Learning curves of the test perplexity for \( h = 10^{-2}, 2.5 \times 10^{-3}, 5 \times 10^{-4} \) are shown in Fig. 3. We observe that the proposed SSI is consistently better than the Euler integrator. Furthermore, mSGNHT is shown to significantly outperform the SGHMC when \( h \) is large. We note that the gap between SSI and the Euler integrator is larger for mSGNHT’s than SGHMC’s, indicating the importance of numerical integrators in higher dimensional systems.

The best performances for each method are shown in Table 1. Note Gibbs sampling typically obtains the best perplexity because it uses the full dataset for each update. However, it is not scalable for large datasets. In our noisy gradient setup, we see that SGNHT-S provides the lowest perplexity among the SG-MCMC methods, including a stochastic sampling method for simplex-structured distributions, Stochastic Gradient Riemannian Langevin Dynamics (SGRLD) (Patterson and Teh 2013).

Logistic Regression We examine logistic regression (LR) on the a9a dataset (Lin, Weng, and Keerthi 2008). The training and testing data consist of 32561 and 16281 data points, respectively, with parameter dimension 123. The minibatch size is set to 10, and the Gaussian prior on the parameters is \( \mathcal{N}(0, 1) \). A thinning interval of 50 is used, with burn-in 300, and \( 3 \times 10^4 \) total iterations. Similar to the experiments for LDA, we test a large range of \( h \). We find that mSGNHT-S gives stable performances across varying \( h \) on the regression model. Test accuracies are compared in Table 2, from which mSGNHT-S also outperforms the recent doubly stochastic variational Bayes (SDVI) (Titsias and Lázaro-Gredilla 2014), and a higher-order variational autoencoder method (HFSVGI) (Fan et al. 2015). More details are provided in the Appendix.

5.2 Deep Models

To illustrate the advantages of the proposed algorithm for deep learning, two deep models are considered. Specifically, for the case of deterministic hidden layers, we consider deep Feedforward (Convolutional) Neural Networks (FNN), and for stochastic latent layers, we consider Deep Poisson Factor Analysis (DPFA) (Gan et al. 2015b).

Deep Neural Networks We evaluate FNN on the MNIST dataset for classification. The data contains 60000 training examples and 10000 testing examples, each being a 28×28 image of handwritten digit. A L-layer FNN is equivalent to compositing \( L \) times of a nonlinear function \( g_{\theta,} \) e.g., the sigmoid function used in the logistic regression model. At the top layer, a softmax function is used for multi-class classification, specifically

\[
P(y|x) \propto \text{softmax}(g_{\theta_L} \circ \cdots \circ g_{\theta_0}(x)),
\]

where \( \circ \) denotes function composition. For each data \( \{x, y\}, \ x \in \mathbb{R}^{784} \) is the raw image, \( y \) is the label. A Gaussian prior is placed on model parameters \( \theta = \{\theta_0, \ldots, \theta_L\} \propto \mathcal{N}(0, \sigma^2 I) \) with \( \sigma^2 = 1 \) in our experiment.

We use the Rectified Linear Unit (ReLU) (Glorot, Bordes, and Bengio 2011) as \( g_{\theta,} \) in each layer. The number of hidden units for each layer is 100, \( D \) is set to 5, stepsize \( h \) is set
to $10^{-4}$, 40 epochs are used. To reduce bias (Chen, Ding, and Carin 2015), $h$ is decreased by half at epoch 20. We test the FNNs with depth $\{2, 3, 4\}$, respectively. Fig. 4 displays learning curves on testing accuracy and training negative log-likelihood. It can be seen that mSGNHT-S consistently converges faster, better than mSGNHT-E for both training and testing. The gaps between mSGNHT-S and mSGNHT-E becomes larger in deeper models. Notably, in the 4-layer FNN, mSGNHT-E failed when $h = 10^{-4}$, while mSGNHT-S worked well. We therefore plot the results for $h = 5 \times 10^{-5}$. From the training plot, mSGNHT-E is failing as learning progresses. It starts to work, only because the stepsize is decreased by half. This confirms that mSGNHT-S is robust to stepsizes, thus is able to mitigate the vanishing/exploding gradient problem in deep models. In addition, we also consider $\theta_0$ as the sigmoid activation function, and the case of convolutional neural networks, empirical results are consistent with the ReLU case. More results are in the Appendix.

Deep Poisson Factor Analysis  
DPFA (Gan et al. 2015a) is a recently proposed framework for deep topic modeling, where interactions between topics are inferred through a deep latent binary hierarchy. We adopt the deep sigmoid belief networks (DSBN) (Gan et al. 2015b) as the deep architecture in the experiment. We use mSGNHT to infer the parameters in DSBN, and the Expanded-Natural reparametrization method to sample from the probabilistic simplex (Patterson and Teh 2013). More details for model specification are in the Appendix.

We test the DPFA on a large dataset, Wikipedia, from which 10M randomly downloaded documents are used, using scripts provided in (Hoffman, Bach, and Blei 2010). We follow the setup in (Gan et al. 2015a), where 1K documents are randomly selected for testing and validation, respectively. The vocabulary size is 7702, and the minibatch size is set to 100, with one pass of the whole data in the experiments. We collect 300 posterior samples to calculate test perplexities, with a standard holdout technique. A three-layer DSBN is employed, with dimensions $128-64-32$ (128 topics right above the data layer). Step sizes are chosen as $10^{-4}$ and $10^{-5}$, and parameter $D = 40$.

The results are shown in Fig. 5, displaying the predictive perplexities on a held-out test set as a function of training documents seen. Clearly, mSGNHT-S converges faster than mSGNHT-E at both chosen stepizes. A magnified plot is shown at the top-right corner of the figure as well, displaying perplexities for the last 10K documents. mSGNHT-S outperforms other recent state-of-the-art methods (shown in semi-transparent plots). Specifically, we compare to, DPFA-SBN trained with Bayesian conditional density filtering (BCDF) (Guhaniyogi, Qamar, and Dunson 2014), DPFA with restricted Boltzmann machines (RBM) (Hinton 2002) trained with mSGNHT-E, and Negative Binomial Focused Topic Model (NB-FTM) (Zhou and Carin 2015) trained with BCDF. The shallow model LDA trained with BCDF is reported as the baseline.

6 Conclusion

A 2nd-order symmetric splitting integrator is proposed to solve the SDE within mSGNHT. This method is shown to be more accurate than the conventional Euler integrator, leading to higher robustness, faster convergence, and more accurate posterior samples. We apply the integrator on mSGNHT for four representative models, including latent Dirichlet allocation, logistic regression, deep neural networks, and deep Poisson factor analysis. Extensive experiments demonstrate that the proposed scheme improves large-scale sampling in terms of convergence speed and accuracy, particularly for deep models.
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References

Bengio, Y.; Simard, P.; and Frasconi, P. 1994. Learning long-term dependencies with gradient descent is difficult. TNN.


Chen, C.; Ding, N.; and Carin, L. 2015. On the convergence of stochastic gradient MCMC algorithms with high-order integrators. In NIPS.


Ding, N.; Fang, Y.; Babbush, R.; Chen, C.; Skeel, R. D.; and Neven, H. 2014. Bayesian sampling using stochastic gradient thermostats. In NIPS.


Kingma, D. P.; and Welling, M. 2014. Auto-encoding variational Bayes. In ICLR.


Korattikara, A.; Rathod, V.; Murphy, K.; and Welling, M. 2015. Bayesian dark knowledge. NIPS.


Ma, Y. A.; Chen, T.; and Fox, E. B. 2015. A complete recipe for stochastic gradient MCMC. NIPS.


Patterson, S., and Teh, Y. W. 2013. Stochastic gradient Riemannian Langevin dynamics on the probability simplex. In NIPS.


Salakhutdinov, R.; Tenenbaum, J. B.; and Torralba, A. 2013. Learning with hierarchical-deep models. TPAMI.


Sutskever, I.; Martens, J.; Dahl, G.; and Hinton, G. 2013. On the importance of initialization and momentum in deep learning. In ICML.


