NEURAL TRANSFORMATIONS FOR EFFICIENT TOPOLOGICAL MIXING

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ABSTRACT

We propose a generalized version of the L2HMC algorithm (Lévy et al., 2018), and evaluate its ability to sample from different topologies in a two-dimensional lattice gauge theory. In particular, we demonstrate that our model is able to successfully mix between modes of different topology, significantly reducing the computational cost required to generate independent gauge configurations.

1 TODO

- 1. Instead of using a time step to vary the network inputs, we use completely different networks for each leapfrog step in our trajectory.
- 2. Explain that "instead of using ||x'-x||, in \mathcal{L}_{θ} we use ..."
- 3. Remove "During training we maintain a buffer of M=2048..." from Section 2
- 4. Combine (5.), (6.), (7.) in Section 2
- 5. Remove Equation 12, combine with Equation 10 and move inline
- 6. Explain $Q_{\mathbb{Z}}$ is a physical quantity (winding number), explain we want to maximize difference but need continuous variable for training, hence $Q_{\mathbb{R}}$
- 7. Change notation for lattice gauge theory, $\varphi_{\mu}(x) \to x_{\mu}(n)$ to be consistent with Section 4.1.

2 Main Contributions

- 1. We propose a generalized version of the L2HMC algorithm that uses different networks (with different step sizes) for each distinct leapfrog step. We represent this generalization by carrying a discrete index $k=0,1,\ldots,N_{\rm LF}$ through the augmented leapfrog equations, (Equation 2, Equation 3) indicating that these functions are allowed to vary.
- 2. We propose a modified loss function, defined in terms of the topological charge metric $\delta_{\mathcal{Q}_{\mathbb{R}}}(\xi',\xi)$,

$$\mathcal{L}_{\theta}[\xi', \xi, A(\xi'|\xi)] = \frac{-A(\xi'|\xi) \cdot \delta_{\mathcal{Q}_{\mathbb{R}}}(\xi', \xi)}{a^2},\tag{1}$$

where $\delta_{\mathcal{Q}_{\mathbb{R}}}(\xi',\xi) \equiv (\mathcal{Q}'_{\mathbb{R}} - \mathcal{Q}_{\mathbb{R}})^2$, and $\mathcal{Q}_{\mathbb{R}} \in \mathbb{R}$ is the *real-valued topological charge*, defined in Section 6. We evaluate this loss at the end of each trajectory during training, and use the gradient information to update the weights θ parameterizing the auxiliary functions $s_i^k, t_i^k, q_i^k, i = x, v$ introduced in the augmented leapfrog updates in Section 4.1.

- 3. During training we maintain a buffer of M=2048 chains ("batch size") which are updated in parallel. Expectation values of physical quantities are calculated as averages over these M chains (after dropping the first $\sim 25\%$ of configurations for thermalization), and we use blocked jackknife resampling to compute the error estimates.
- 4. We introduce an annealing schedule (Section 5) $\{\gamma\}_{t=0}^N = \{\gamma_0, \gamma_1, \ldots \gamma_N\}$, with $\gamma_0 < \gamma_1 < \cdots < \gamma_N$, with $|\gamma_t| < 1$ and $\gamma_{t+1} \gamma_t \ll 1$ that is slowly varied during training (and removed entirely during inference) that scales the potential energy function in our target distribution, $p_t(x) \propto e^{\gamma_t S(x)}$, that allows our sampler to explore previously inaccessible regions of the phase space by shrinking the height of the various energy barriers separating isolated modes.

- 5. We apply the proposed method to a 1 + 1-dimensional U(1) lattice gauge theory defined on a $N_x \times N_t$ lattice with periodic boundary conditions, and report a significant reduction in the computational cost required to generate independent gauge configurations.
- 6. We use the *integrated autocorrelation time* of the topological charge $\tau_{\text{int}}^{\mathcal{Q}}$ as a metric for determining the efficiency of our trained sampler. The improvement for the trained model compared to generic HMC can be seen in Figure 3b.
- 7. We compare our results to traditional HMC across a variety of trajectory lengths and inverse coupling constants β , and show that our trained model consistently outperforms traditional HMC. These results can be seen in Section A.3.

3 RELATED WORK:

Recently, there has been a significant interest in applying probabilistic programming techniques with autodifferentiation capabilities to develop more efficient simulations. [not sure what this means: probabilistic programming techniques]

In particular, following the development of the RealNVP (Dinh et al., 2016) architecture, there has been an explosion of different proposals that aim to take advantage of the invertible network architecture. Because of the enormous computational burden faced by generative techniques in lattice gauge theory, there have been multiple works (Albergo et al., 2019; 2021; Favoni et al., 2020; Medvidovic et al., 2020; Neklyudov & Welling, 2020; Neklyudov et al., 2020; Li et al., 2020; Boyda et al., 2020; Kanwar et al., 2020; Toth et al., 2019; Hoffman et al., 2019; Wehenkel & Louppe, 2020; Pasarica & Gelman, 2010; Dinh et al., 2016; Tanaka & Tomiya, 2017; Schaefer et al., 2009; Cossu et al., 2018; Rezende et al., 2020), that look at developing more efficient sampling techniques specifically tailored for lattice gauge/field theories.

4 Background

We provide a review of the generic Hamiltonian Monte Carlo (HMC) algorithm and setup some of the relevant notation in Section A.1.

4.1 GENERALIZING THE LEAPFROG INTEGRATOR: L2HMC

In (Lévy et al., 2018), the authors propose the L2HMC ("Learning to Hamiltonian Monte Carlo") algorithm, and demonstrate its ability to outperform traditional Hamiltonian Monte Carlo (HMC) on a variety of two-dimensional target distributions. For example, the trained L2HMC sampler is shown to be capable of exploring regions of phase space which are typically inaccessible with traditional HMC. Additionally, they show that the trained sampler is efficient at mixing between modes of a multi-modal target distribution, a feature which is highly desirable for MCMC simulations of lattice gauge theory.

We denote a complete state by $\xi=(x,v,d)$ with target distribution $p(\xi)=p(x,v,d)=p(x)\cdot p(v)\cdot p(d)$. Here we've introduced a binary direction variable (distributed independently of both x and v) $d\sim \mathcal{U}(+,-)$ which is resampled following each Metropolis-Hastings accept/reject step, and can be interpreted as determining the "direction" (forward/backward) of our update. The key modification of the L2HMC algorithm is the introduction of six auxiliary functions s_i, t_i, q_i for i=x,v into the leapfrog updates, which are parameterized by weights θ in a neural network.

For simplicity, we consider the forward d = +1 direction, and introduce the notation:

$$v_k' \equiv \Gamma_k^+(v_k; \zeta_{v_k}) = v_k \odot \exp\left(\frac{\varepsilon_v^k}{2} s_v^k(\zeta_{v_k})\right) - \frac{\varepsilon_v^k}{2} \left[\partial_x S(x_k) \odot \exp\left(\varepsilon_v^k q_v^k(\zeta_{v_k})\right) + t_v^k(\zeta_{v_k})\right], \quad (2)$$

$$x_k' \equiv \Lambda^+(x_k; \zeta_{x_k}) = x_k \odot \exp(\varepsilon_x^k s_x^k(\zeta_{x_k})) + \varepsilon_x^k \left[v_k' \odot \exp(\varepsilon_x^k q_x^k(\zeta_{x_k})) + t_x^k(\zeta_{x_k}) \right]$$
(3)

where (1.) $\zeta_{v_k}=(x_k,\partial_x S(x_k),\tau(k)),\ \zeta_{x_k}=(x_k,v_k,\tau(k))$ are a shorthand notation for the input passed to the respective s,t,q network functions 1 (2.) $\tau(k)=\left[\cos\frac{2\pi k}{N_{\mathrm{LF}}},\sin\frac{2\pi k}{N_{\mathrm{LF}}}\right],$

¹Note that ζ_x , ζ_v are chosen to be subsets of the augmented state space that are independent of the variable being updated.

 $k=0,1,\ldots,N_{\rm LF}$, is a discrete time variable parameterizing our trajectory, and (3.) we indicate the forward d=+1 direction by the + superscript on Γ^+ , and Λ^+ . Using this notation, we can write the complete leapfrog update (in the forward d=+1 direction) as:

1. Half-step momentum update: $v'_k = \Gamma_k^+(v_k; \zeta_{v_k})$

2. Full-step half-position update²: $x'_k = \overline{m}^t \odot x_k + m^t \odot \Lambda_k^+(x_k; \zeta_{x_k})$

3. Full-step half-position update: $x_k'' = \overline{m}^t \odot \Lambda_k^+(x_k'; \zeta_{x_k'}) + m^t \odot x_k'$

4. Half-step momentum update: $v_k'' = \Gamma^+(v_k'; \zeta_{v_k'})$

Note that in order to keep our leapfrog update reversible, we've split the x update into two subupdates by introducing a binary mask $m^t = m^t \odot \mathbb{1} + \bar{m}^t \odot \mathbb{1}$ that updates half of the components of x sequentially, as shown (with the general transformation Λ^{\pm}) in Figure 1.

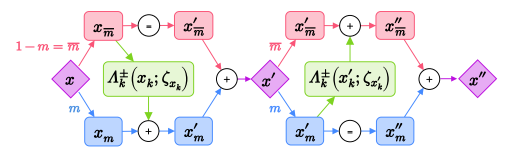


Figure 1: Illustration of the split x update. Here we include the general form of the transformation $\Lambda_k^{\pm}(x_k;\zeta_{x_k})$.

As in HMC, we form a complete trajectory by performing $N_{\rm LF}$ leapfrog steps in sequence, followed by a Metropolis-Hastings accept/reject step as described in Equation 16. However, unlike in the expression for HMC, we must take into account the overall Jacobian factor from the update $\xi \to \xi'$, which can be easily computed as

$$\left| \frac{\partial v_k'}{\partial v_k} \right| = \exp\left(\frac{\varepsilon_v^k}{s} s_v^k(\zeta_{v_k})\right), \quad \left| \frac{\partial x_k'}{\partial x_k} \right| = \exp\left(\varepsilon_x^k s_x^k(\zeta_{x_k})\right). \tag{4}$$

So far we've restricted our attention to the forward (d=+1) update, however we can obtain the expressions for the reverse direction simply by inverting the two functions $\Gamma^- \equiv (\Gamma^+)^{-1}$, $\Lambda^- \equiv (\Lambda^+)^{-1}$, and performing the updates in the reverse order.

In order to perform the updates in the generalized leapfrog integrator, we need to evaluate each of the functions s, t, q. Without loss of generality³, we temporarily ignore the discrete leapfrog index k, and restrict our attention to the s_x, t_x, q_x functions used in the x update, Equation 3.

Each of the s_x, t_x, q_x functions takes as input $\zeta_x = (x, v, \tau)$, with $x \in \mathbb{R}^n$, $v \in \mathbb{R}^n$, and $\tau \in \mathbb{R}^2$. The network splits these inputs and constructs the following intermediate variable (where σ denotes an arbitrary activation function)

$$z_1 = \sigma(w_x^T x + w_y^T v + w_\tau^T \tau + b). {5}$$

This intermediate variable z_1 is then passed through another series of fully-connected layers,

$$z_n = \sigma(w_n^T z_{n-1} + b_n), \ z_{n-1} = \sigma(w_{n-2}^T z_{n-2} + b_{n-2}), \ \dots, \ z_2 = \sigma(w_2^T z_1 + b_2).$$
 (6)

The network outputs s_x, t_x, q_x are then defined in terms of this final hidden variable z_n as

$$s_x(\zeta_x) = \alpha_s \tanh(w_s z_n + b_s), \quad t_x(\zeta_x) = w_t^T z_n + b_t, \quad q_x(\zeta_x) = \alpha_q \tanh(w_q z_n + b_q)$$
 (7)

²By this we mean we are performing a complete update step that only updates half of the components of x determined by the mask m^t and its complement \bar{m}^t .

 $^{^3}$ Because we maintain a separate network with identical architecture for evaluating the s,t,q functions in the momentum update Equation 2, the procedure is identical

where α_s , and α_q are trainable scaling factors. The only requirement on the details of the network is that the dimensionality of the outputs s_x, t_x, q_x match the dimensionality of our physical variables $x, v \in \mathbb{R}^n$.

Next, we introduce a loss function

$$\mathcal{L}_{\theta}(\xi, \xi', A(\xi'|\xi)) = -\frac{A(\xi'|\xi) \cdot \delta(\xi, \xi')}{a^2}$$
(8)

where $\delta(\xi, \xi')$ is a suitably chosen *metric function*, and a is a scaling factor.

5 Annealing Schedule

To help our sampler overcome the large energy barriers between isolated modes, we introduce an *annealing schedule*, during the training phase

$$\{\gamma_t\}_{t=0}^N = \{\gamma_0, \gamma_1, \dots, \gamma_{N-1}, \gamma_N\},$$
 (9)

where $\gamma_0 < \gamma_1 < \cdots < \gamma_N \equiv 1, \gamma_{t+1} - \gamma_t \ll 1$, and N denotes the total number of training steps to be performed. Note that we are free to vary γ during the initial training phase as long as we recover the true distribution with $\gamma \equiv 1$ at the end of training and evaluate our trained model without this factor. Explicitly, for $\gamma_t < 1$ this rescaling factor helps to reduce the height of the energy barriers, making it easier for our sampler to explore previously inaccessible regions of the phase space. In terms of this additional annealing schedule, our target distribution picks up an additional index t to represent our progress through the training phase, which can be written explicitly as

$$p_t(x) \propto e^{-\gamma_t S(x)}, \quad \text{for} \quad t = 0, 1, \dots, N.$$
 (10)

6 LATTICE GAUGE THEORY

Note: We define the lattice gauge theory in terms of the link variables $-\pi \le \varphi_{\mu}(x) < \pi$, which we explicitly choose to identify as the physical (position) variable x in Section 4.1, i.e.

$$\varphi_{\mu}(x) \Longleftrightarrow x, \quad p(x) \Longleftrightarrow p(\varphi), \quad \text{etc.}$$
 (11)

Our theory is defined on an $N_x \times N_t$ two-dimensional lattice with periodic boundary conditions. Let $U_\mu(x) = e^{i\varphi_\mu(x)} \in U(1)$, with $\varphi_\mu(x) \in [-\pi,\pi]$ denote the *link variables*, where $\varphi_\mu(x)$ denotes the link oriented in the $\hat{\mu}$ -direction located at the site x. We can write our target distribution $p_t(\varphi)$ in terms of the Wilson action, $S(\varphi)$ as

$$p_t(\varphi) = e^{-\gamma_t \beta \cdot S(\varphi)}, \quad \text{with} \quad \beta \cdot S(\varphi) = \sum_P 1 - \cos(\varphi_P)$$
 (12)

and $\varphi_P \equiv \varphi_\mu(x) + \varphi_\nu(x+\hat{\mu}) - \varphi_\mu(x+\hat{\nu}) - \varphi_\nu(x)$ is the sum of the link variables around the 1×1 elementary plaquette, as shown in Figure 2. Here, $\beta=2/g_0^2$ is the inverse coupling constant and $\beta\to\infty$ recovers the continuum limit of the theory. We consider two physical quantities of interest, namely the *real* and *integer* valued *topological charge*, $(\mathcal{Q}_\mathbb{R}\in\mathbb{R}, \mathcal{Q}_\mathbb{R}\in\mathbb{R})$, and $\mathcal{Q}_\mathbb{Z}\in\mathbb{Z}$, respectively) defined as

$$Q_{\mathbb{R}}(\varphi) \equiv \frac{1}{2\pi} \sum_{P} \sin(\varphi_{P}), \quad Q_{\mathbb{Z}}(\varphi) \equiv \frac{1}{2\pi} \sum_{P} \lfloor \varphi_{P} \rfloor, \quad \text{where} \quad \lfloor \varphi_{P} \rfloor = \varphi_{P} - 2\pi \left\lfloor \frac{\varphi_{P} + \pi}{2\pi} \right\rfloor$$

$$U_{\mu}(x) = e^{i\varphi_{\mu}(x)}, \quad \varphi_{\mu}(x) \in [-\pi, \pi]$$
(13)

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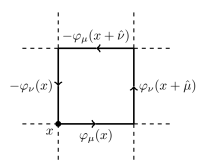
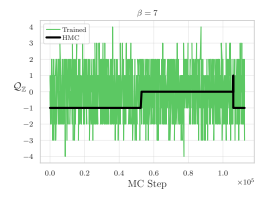
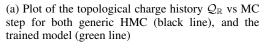
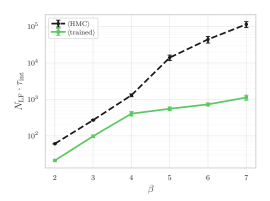


Figure 2: Elementary plaquette, P on the lattice.







(b) Plot of the estimated integrated autocorrelation time $au_{\mathrm{int}}^{\mathcal{Q}_{\mathbb{R}}}$ vs β for both generic HMC (black dashed line), and the trained model (solid green line).

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A APPENDIX

A.1 HAMILTONIAN MONTE CARLO

The Hamiltonian Monte Carlo (HMC) algorithm is a widely used technique that allows us to sample from an analytically known target distribution p(x) by constructing a chain of states $\{x^{(0)}, x^{(1)}, \ldots, x^{(n)}\}$, such that $x^{(n)} \sim p(x)$ in the limit $n \to \infty$. For our purposes, we assume that our target distribution can be expressed as a Boltzmann distribution, $p(x) = \frac{1}{Z}e^{-S(x)} \propto e^{-S(x)}$, where S(x) is the *action* of our theory, and Z is often referred to as the *partition function*, which ensures our target distribution is correctly normalized to unity. In this case, HMC begins by augmenting the state space with a fictitious momentum variable v, normally distributed independently of x, i.e. $v \sim \mathcal{N}(0,1)$. Our joint distribution can then be written as

$$p(x,v) = p(x) \cdot p(v) \propto e^{-S(x)} \cdot e^{-\frac{1}{2}v^T v} = e^{-\mathcal{H}(x,v)}$$
 (14)

where $\mathcal{H}(x,v)$ is the Hamiltonian of the joint (x,v) system. Notably, this system obeys Hamilton's equations

$$\dot{x} = \frac{\partial \mathcal{H}}{\partial v}, \quad \dot{v} = -\frac{\partial \mathcal{H}}{\partial x}$$
 (15)

which can be integrated using the *leapfrog integrator* along iso-probability contours defined by $\mathcal{H}=\mathrm{const.}$ Explicitly, for a step size ε and initial state $\xi=(x,v)$, the leapfrog integrator generates a proposal configuration $\xi'\equiv(x',v')$ by performing the following series of updates:

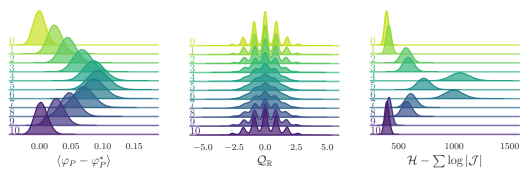
- 1. Half-step momentum update: $v^{1/2} \equiv v \left(t + \frac{\varepsilon}{2}\right) = v \frac{\varepsilon}{2} \partial_x S(x)$
- 2. Full-step position update: $x' \equiv x(t+\varepsilon) = x + \varepsilon v^{1/2}$
- 3. Half-step momentum update: $v' \equiv v(t+\varepsilon) = v^{1/2} \frac{\varepsilon}{2} \partial_x S(x')$

We can then construct a complete *trajectory* of length $\lambda = \varepsilon \cdot N_{\rm LF}$ by performing $N_{\rm LF}$ leapfrog steps in sequence. At the end of our trajectory, we either accept or reject the proposal configuration according to the Metropolis-Hastings acceptance criteria,

$$x_{i+1} = \begin{cases} x' & \text{with probability } A(\xi'|\xi) \\ x & \text{with probability } (1 - A(\xi'|\xi)), & \text{where} \quad A(\xi'|\xi) = \min\left\{1, \frac{p(\xi')}{p(\xi)} \left| \frac{\partial \xi'}{\partial \xi^T} \right| \right\}. \end{cases}$$
(16)

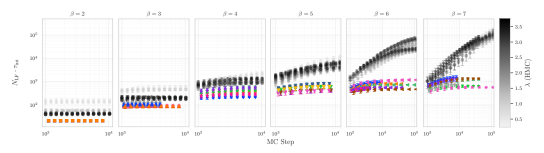
The generic leapfrog integrator is known to be symplectic (conserves energy), so the Jacobian factor reduces to $\left|\frac{\partial \xi'}{\partial \xi^T}\right| = 1$.

A.2 EXTRA FIGURES

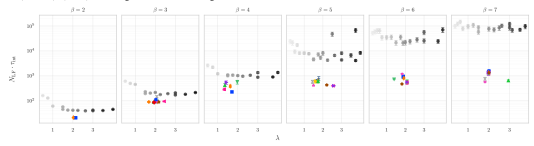


- (a) Error in the average plaquette, $\langle \varphi_P \varphi_P^* \rangle$ vs leapfrog step.
- (b) The real-valued topological charge, $\mathcal{Q}_{\mathbb{R}}$ vs leapfrog step.
- (c) The rescaled energy, $\mathcal{H} \sum \log \|\mathcal{J}\|$ vs leapfrog step.

Figure 4: Illustration of how various observables vary over a single trajectory for the trained sampler.



(a) Plot of the integrated autocorrelation time $au_{\mathrm{int}}^{\mathcal{Q}_{\mathbb{R}}}$ vs MC step for both generic HMC and the trained model, for $\beta=2,3,\ldots,7$ arranged from left to right.



(b) Plot of the integrated autocorrelation time $au_{\mathrm{int}}^{\mathcal{Q}_{\mathbb{R}}}$ vs λ for both generic HMC and the trained model.

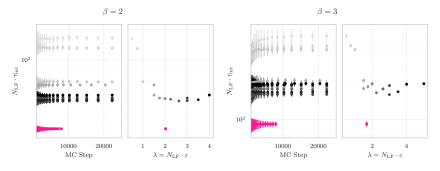


Figure 6: Plot of the estimated integrated autocorrelation time of the topological charge, $\tau_{\rm int}^{\mathcal{Q}}$ vs MC Step (left), and trajectory length λ (right) for both HMC (grey circles) and the trained model (pink squares)

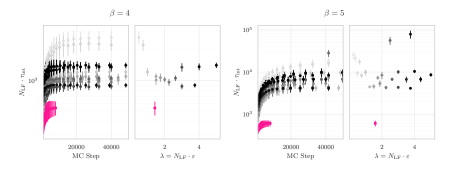


Figure 7: Plot of the estimated integrated autocorrelation time of the topological charge, $\tau_{\rm int}^{\mathcal{Q}}$ vs MC Step (left), and trajectory length λ (right) for both HMC (grey circles) and the trained model (pink squares)

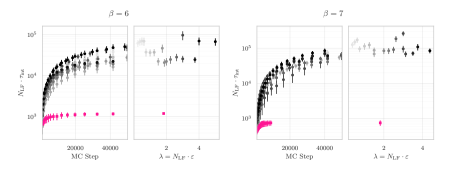


Figure 8: Plot of the estimated integrated autocorrelation time of the topological charge, $\tau_{\rm int}^{\mathcal{Q}}$ vs MC Step (left), and trajectory length λ (right) for both HMC (grey circles) and the trained model (pink squares)

A.3 AUTOCORRELATION PLOTS

A.4 TRAINING ALGORITHM

```
Algorithm 1: Training procedure
input :
                                                                                                                                                                            [1]
          1. Target distribution, p_t(x) \propto e^{-\beta_t U(x)}
          2. Loss function, \mathcal{L}_{\theta}(\xi', \xi, A(\xi'|\xi))
          3. Learning rate schedule, \{\alpha_t\}_{t=0}^{N_{\mathrm{train}}}
          4. Annealing schedule, \{\beta_t\}_{t=0}^{N_{\text{train}}}
          5. Batch of initial states, x
Initialize weights \theta
                                                                                                                                                                            [2]
for 0 \le t < N_{\text{train}}:
                                                                                                                                                                            [3]
      update: p_t(x) \propto e^{-\beta_t U(x)}
                                                                                                                                                                            [4]
       resample: v \sim \mathcal{N}(0, 1)
                                                                                                                                                                            [5]
       resample: d \sim \mathcal{U}(+,-)
                                                                                                                                                                            [6]
      construct: \xi \equiv (x, v, d) for 0 \le \ell < N_{\rm LF} :
                                                                                                                                                                            [7]
                                                                                                                                                                           [8]
        propose: \xi'_{\ell} \leftarrow \mathbf{FL}^{\pm}_{\ell} \xi_{\ell}
                                                                                                                                                                           [9]
      \begin{aligned} & \text{compute: } A(\xi'|\xi) = \min \left\{ 1, \frac{p(\xi')}{p(\xi)} \left| \frac{\partial \xi'}{\partial \xi^T} \right| \right\} \\ & \text{update: } \mathcal{L} \leftarrow \mathcal{L}_{\theta}(\xi', \xi, A(\xi'|\xi)) \\ & \text{backprop: } \theta \leftarrow \theta - \alpha_t \nabla_{\theta} \mathcal{L} \end{aligned}
                                                                                                                                                                          [10]
                                                                                                                                                                          [11]
                                                                                                                                                                          [12]
      assign: x_{t+1} \leftarrow \begin{cases} x' & \text{with probability } A(\xi'|\xi) \\ x & \text{with probability } (1 - A(\xi'|\xi)). \end{cases}
                                                                                                                                                                          [13]
                                                                                                                                                                          [14]
```