# Deep Learning Hamiltonian Monte Carlo [1]

Building topological samplers for lattice gauge theories

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## bit.ly/l2hmc-qcd

#### **Abstract**

• We generalize the Hamiltonian Monte Carlo (HMC) algorithm with a stack of trainable neural network (NN) layers and evaluate its ability to sample from different topologies in a two-dimensional lattice gauge theory. We demonstrate that our model is able to successfully mix between modes of different topologies, significantly reducing the computational cost required to generate independent gauge field configurations.

#### **Motivation**

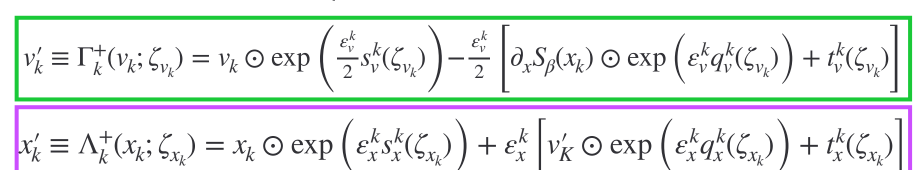
- Markov Chain Monte Carlo (MCMC) methods are pervasive throughout science and are used in applications ranging from epidemiological modeling to election forecasting.
- Recent developments in ML, together with ever-more-capable hardware has led to a resurgence in developing faster, more efficient simulation techniques.
- In particular, the development of invertible NN architectures has opened the flood-gates for new approaches that are capable of outperforming traditional techniques on particularly challenging distributions.
- Simulations in lattice gauge theory / lattice QCD are currently limited by our ability to generate independent configurations, making it a prime target for testing novel approaches.
- Here, we propose a generalized version of the L2HMC [2] and demonstrate its ability to significantly reduce the computational effort required to generate configurations for a two-dimensional U(1) lattice gauge theory.

#### Method

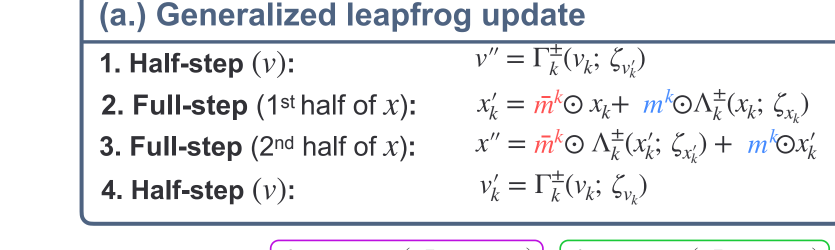
• We propose a generalized version of the L2HMC algorithm [2] that introduces distinct neural networks for each leapfrog step called leapfrog layers, shown in (a).

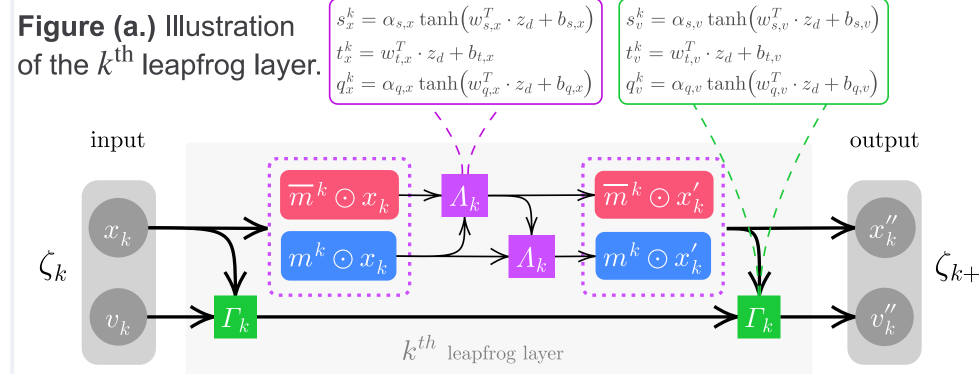
Denote the leapfrog step / layer by a discrete index

- $k = \{0,1,...,N_{\mathrm{LF}}\} \in \mathbb{N}$  where  $N_{\mathrm{LF}}$  is the total number of leapfrogs. • Introduce  $d \sim \mathcal{U}(+,-)$  (direction—forward/backward) and denote the complete state  $\xi = (x, v, d)$ , then the *target distribution* is given by  $p(\xi) = p(x) \cdot p(v) \cdot p(d).$
- Each leapfrog step transforms  $\xi_k \equiv (x_k, v_k, d_k) \to (x_k'', v_k'', d_k) = \xi_k''$  by passing it through  $k^{\text{th}}$  leapfrog layer (note the direction,  $d_k$  is persistent).
- Consider the forward d = +1 direction and introduce the notation:



• We can write a complete leapfrog update as: (compare with **HMC** (i.)





- In order to keep our model reversible, we split the x update into two subupdates using a binary mask,  $m^k = 1 - \bar{m}^k$  that updates half of the components of x sequentially.
- We've introduced the shorthand notation for the networks' inputs:

$$\zeta_{v_k} \equiv (x_k, \partial_x S_\beta(x)), \zeta_{x_k} \equiv (m_k \odot x_k, v_k)$$

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 We must now explicitly keep track of the Jacobian factor of the update  $\xi o \xi'$ , these are easily computed to be

$$\left| \frac{\partial v_k''}{\partial v_k} \right| = \exp\left(\frac{1}{2} \varepsilon_v^k s_v^k (\zeta_{v_k}), \left| \frac{\partial x_k''}{\partial x_k} \right| = \exp\left(\varepsilon_x^k s_v^k (x_k)\right).$$

#### Method (contd.)

- For target distributions in Euclidean space,  $x \in \mathbb{R}^n$ , we define a loss function that encourages our sampler to move large distances in the phase space.
- To do this, we can maximize the expected squared jump distance (ESJD):

$$\mathcal{L}(\theta) \equiv \mathbb{E}_{p(\xi)} \left[ \delta(\xi, \xi') \cdot A(\xi' | \xi) \right] , \ \delta(\xi, \xi') \equiv \|x - x'\|^2.$$

where  $\theta$  are (collectively) the weights in the NN

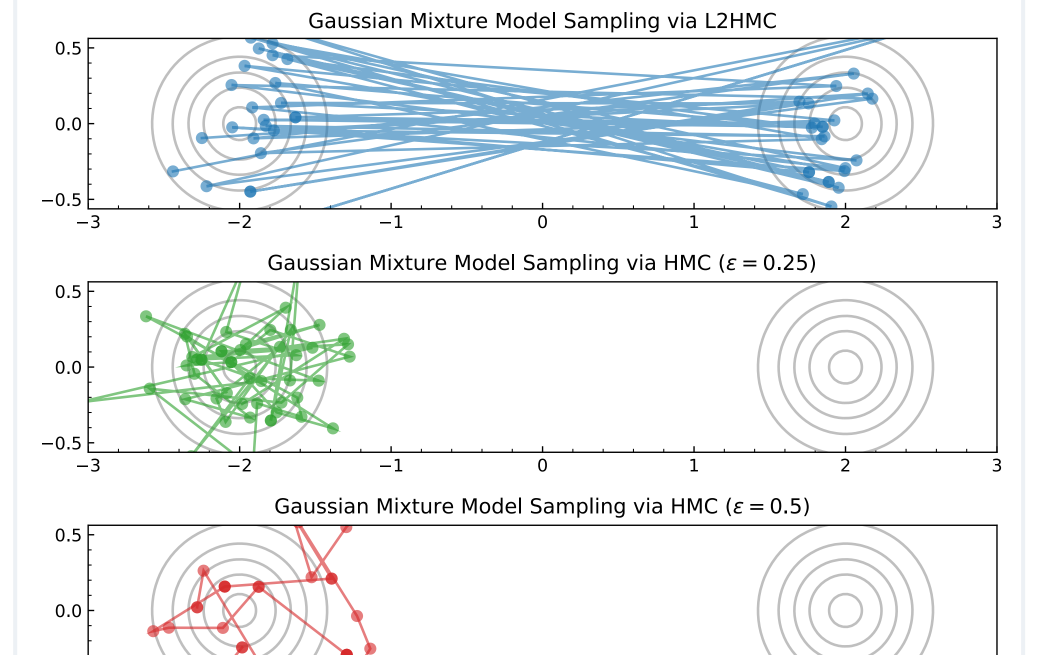


Figure (b.) Illustration of contours from the true target distribution vs samples obtained from both the trained model and generic HMC. We can see that HMC fails to mix between the two modes of the distribution whereas the trained model efficiently jumps between them.

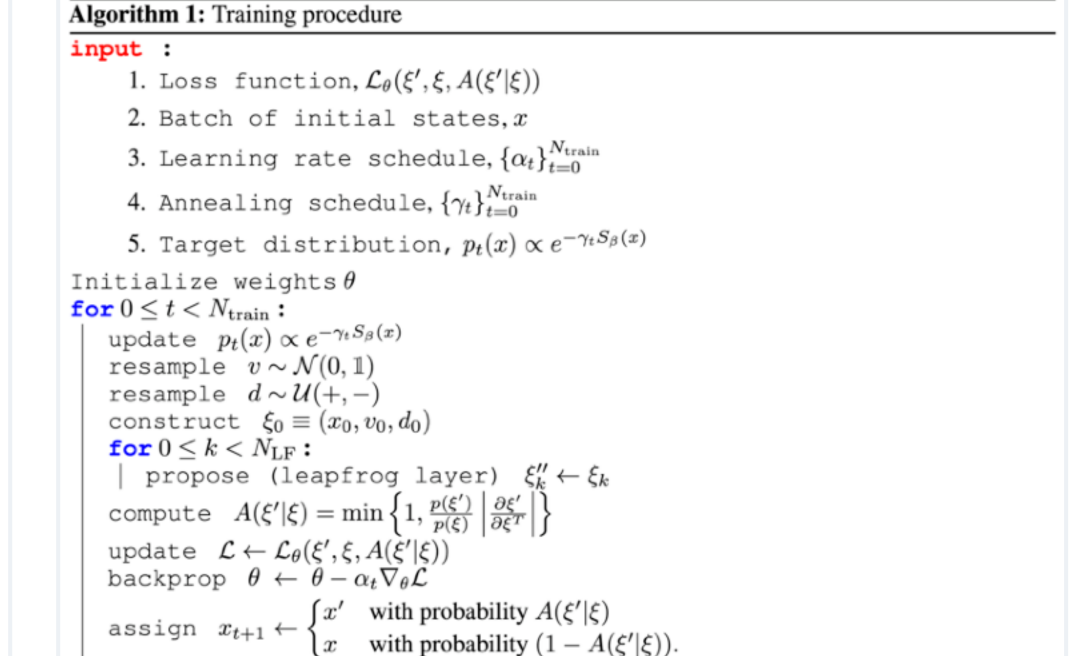
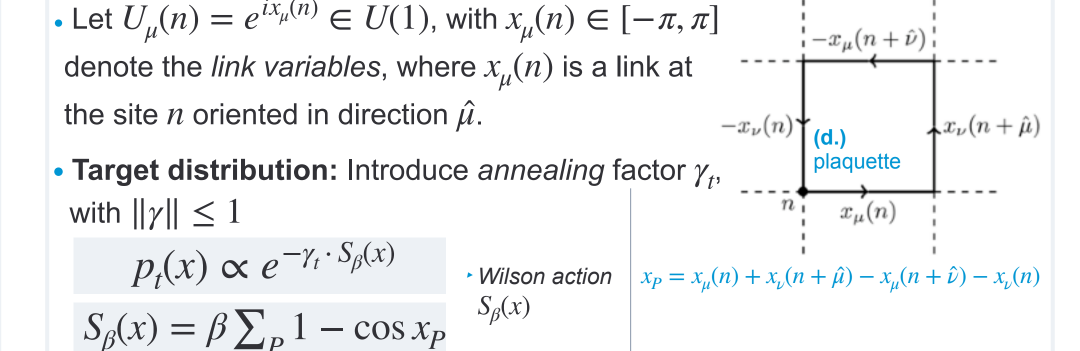
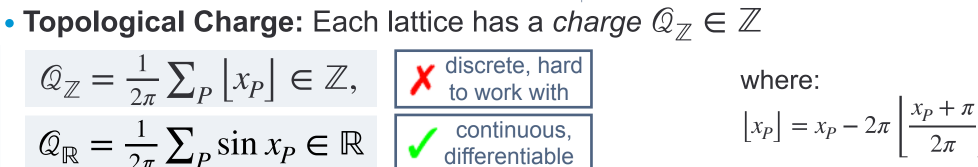


Figure (c.) Summary of training algorithm.

## **Application to Lattice Gauge Theory**





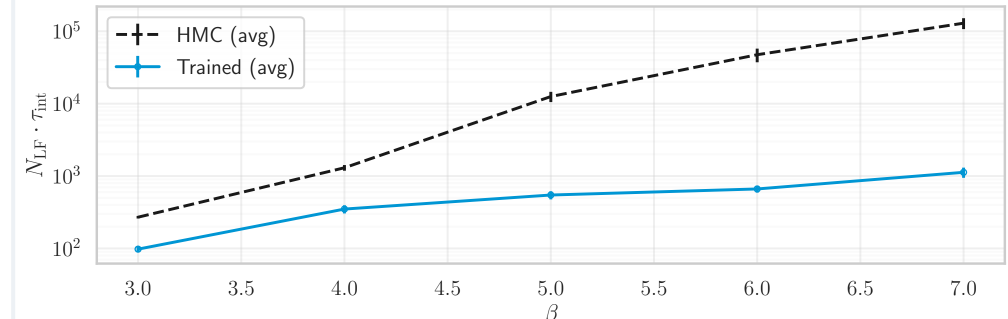
• Loss function: We maximize the expected squared charge difference

$$\mathcal{L}(\theta) = \mathbb{E}_{p(\xi)} \left[ -\delta(\xi', \xi) \cdot A(\xi' | \xi) \right]$$
$$\delta(\xi', \xi) = \left[ \mathcal{Q}_{\mathbb{R}}(x') - \mathcal{Q}_{\mathbb{R}}(x) \right]^{2}$$

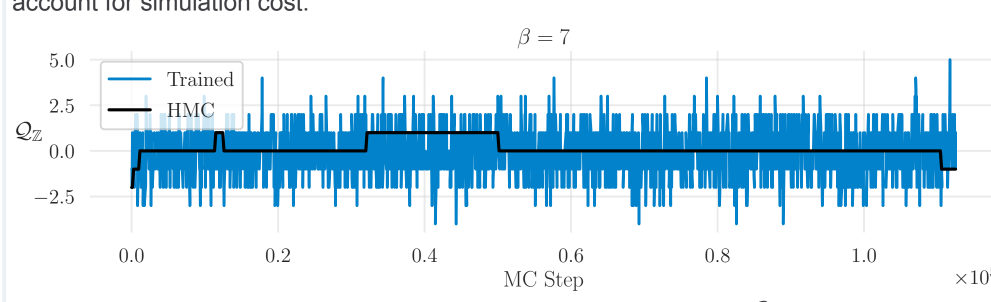
where  $A(\xi'|\xi)$  is given in **HMC** (j.)

#### Results

- In order to measure the computational cost of our approach we use the integrated autocorrelation time  $au_{\mathrm{int}}^{\mathbb{Q}_{\mathbb{Z}}}$ , which can be roughly interpreted as the number of trajectories (on average) before an independent sample is drawn.
- We can see in Figure (e.) that the trained model consistently outperforms generic HMC across  $\beta = 2,3,...,7$ .



**Figure (e.)** Estimate of the integrated autocorrelation time  $N_{\rm LF} \cdot \tau_{\rm int}^{\mathcal{Q}_{\mathbb{Z}}}$  vs  $\beta$ , scaled by  $N_{\rm LF}$  to account for simulation cost.



**Figure (f.)** Illustration of the integer valued topological charge  $Q_{\mathbb{Z}}$  vs MC step for both HMC (black) and the trained model (blue).

- In order to understand the mechanism driving this improved behavior, we looked at how different physical quantities evolve during a single trajectory in the trained model as shown in Figures (g.), (h.).
- We see that our sampler artificially increased the energy of the physical system during the first half of the trajectory, before returning back to its original physical value.
- We believe that this ability to vary the energy during the trajectory helps the sampler to overcome energy barriers between topological sectors whereas HMC remains stuck.

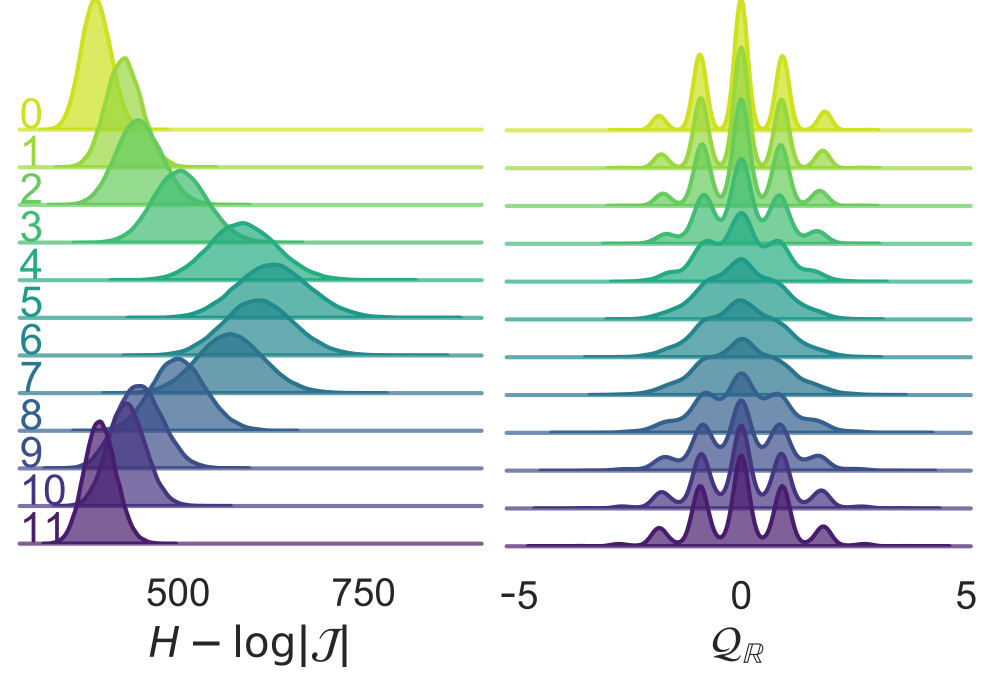
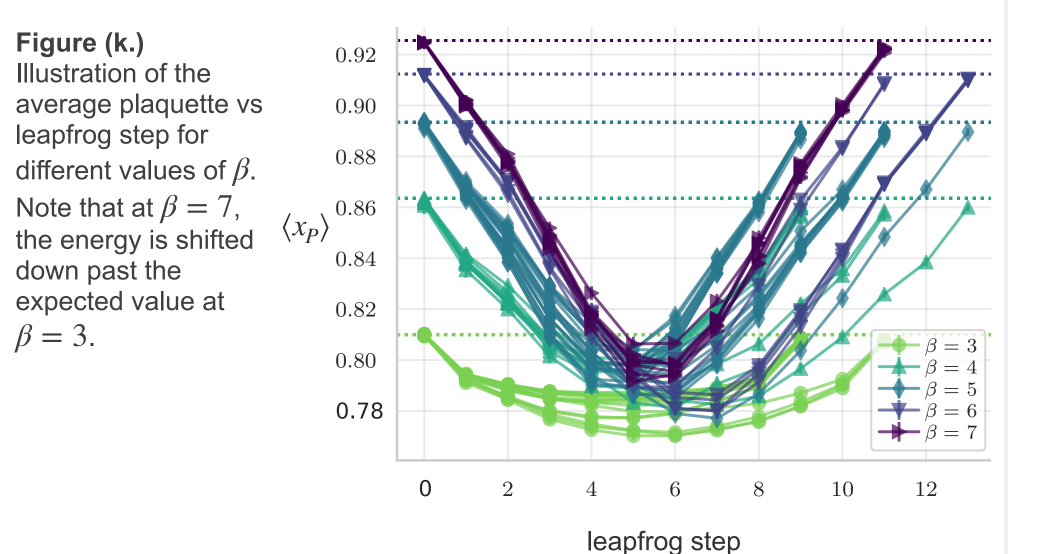


Figure (h.) The real-valued topological Figure (g.) The adjusted energy  $H - \log |\mathcal{J}|$  at intermediate leapfrog charge  $\mathcal{Q}_{\square}$  at intermediate leapfrog layers.



#### **Next Steps**

 Going forward we plan to continue development of this approach towards more complex theories in higher space-time dimensions (e.g. 2D, 4D SU(3)).

#### **Training Costs**

 Our models were trained using Horovod on the ThetaGPU supercomputer at the Argonne Leadership Computing Facility (ALCF). A typical training run on 1 node ( $8 \times NVIDIAA100 GPUs$ ) using a batch size M=2048, hidden layer shapes [256,256,256] for each of the  $N_{\rm LF} = 10$  leapfrog layers, on a  $16 \times 16$  lattice for  $5 \times 10^5$  training steps takes roughly 24 hours to complete.

#### Conclusion

- Presented a generalized version of the L2HMC algorithm—consisting of a stack of leapfrog layers—that improves the existing approaches' flexibility while remaining statistically exact.
- Shown that our trained model successfully mixes between modes of a two-dimensional Gaussian Mixture Model while HMC remains stuck in a local mode.
- Looked at applying the described approach to a two-dimensional U(1) lattice gauge theory
- Saw that for this lattice gauge model, our trained sampler is capable of significantly outperforming traditional HMC across a range of coupling constants.

#### References

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#### Hamiltonian Monte Carlo (HMC)

- **Goal**: Sample from (difficult) target distribution:  $p(x) \propto e^{-S(x)}$
- Method: For  $x \in U(1)^n$ , build chain  $x_0 \to x_1 \to \dots, \to x_N$  such that  $x_N \sim p(x) \text{ as } N \to \infty$
- Introduce  $v \sim \mathcal{N}\left(0, I_n\right) \in \mathbb{R}^n$ , write joint distribution:

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

• Evolve the system of equations  $\dot{x} = \frac{\partial H}{\partial y}$ ,  $\dot{y} = -\frac{\partial H}{\partial x}$  using the **leapfrog** 

#### integrator (a)

along  $H = \text{const: } \xi \equiv (x, v) \rightarrow (x', v') = \xi'$ 

• Accept or reject proposal configuration  $\xi'$  using **Metropolis-Hastings** 

### (i.) Leapfrog update 1. Half-step (v): $\tilde{v} = v - \frac{\varepsilon}{2} \partial_x S(x)$

2. Full-step (x):  $x' = x + \varepsilon \tilde{v}$ 3. Half-step (v):  $v' = \tilde{v} - \frac{\varepsilon}{2} \partial_x S(x')$ 

(j.) Metropolis-Hastings





