Reduced critical slowing down for statistical physics simulations

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Wang-Landau simulations offer the possibility to integrate explicitly over a collective coordinate and stochastically over the remainder of configuration space. We propose to choose the so-called “slow mode”, which is responsible for large autocorrelation times and thus critical slowing down, for collective integration. We study this proposal for the Ising model and the linear-log-relaxation (LLR) method as simulation algorithm. We firstly demonstrate super critical slowing down in a phase with spontaneously broken symmetry and for the heatbath algorithms, for which autocorrelation times grow exponentially with system size. By contrast, using the magnetisation as collective coordinate, we present evidence that super critical slowing down is absent. We still observe a polynomial increase of the autocorrelation time with volume (critical slowing down), which is however reduced by orders of magnitude when compared to local update techniques.

I. INTRODUCTION

Stochastic simulations of lattice theories combined with modern computer resources have rapidly evolved to an exceptional theoretical framework enlightening research areas such as Quantum Field Theory [1] and Statistical Physics [2]. Markov Chain Monte Carlo (MCMC) simulations in conjunction with a local update of the degrees of freedom are ubiquitous in the quiver of possibilities.

In MCMC simulations, a bunch of local updates - usually called MC sweep - result into a new configuration of degrees of freedom on the lattice. The simulations generates sequentially a string of lattice configurations. Under the Markov assumption, any configuration only depends on its predecessor. Objects of interests are expectation values. By virtue of the law of large numbers [3], those can be estimated using the $N$ configurations of the Markov set:

$$\langle A \rangle \approx \frac{1}{N} \sum_{i=1}^{N} A_i.$$ 

The price to pay for a finite reach $N$ is that the above estimator is afflicted by a statistical error $\epsilon_A$, which scales like $1/\sqrt{N}$ under the Markov assumption (and assuming that the variance of $A$ exists).

In practical Monte-Carlo simulations, configurations are correlated over a characteristic number of Monte-Carlo updates $t \approx \tau$, which is called autocorrelation time (we will give a proper definition below). An immediate impact is that the statistical error now scales like $\sqrt{\tau/N}$. Large autocorrelations times severely limit the usefulness of simulations at moderate computational costs, and a good deal of algorithmic research has been devoted to simulation methods with small autocorrelations.

The autocorrelation time depends on the simulation algorithm, the parameters of the simulated theory and the system size, say volume $V$, which could be the number of lattice sites. Of particular interest for many applications is a parameter regime that leaves the lattice degrees of freedom correlated over a typical spatial scale $\xi$ (correlation length). In Solid State Physics, $\xi$ diverges at a second order phase transition. In quantum physics simulations $1/\xi$ acts a regulator for the inherent divergencies of the underpinning quantum field theory, and the limit $\xi \to \infty$ is of crucial importance to extract physics relevant information from those computer simulations. Generating independent Markov ensembles in the case that degrees of freedom are correlated over many sites is a challenge for any algorithm and in particular for the important class of local update algorithms. This challenge is reflected by the monotonically increasing function $\tau(\xi)$ which describes the connection between correlation length $\xi$ and the autocorrelation time $\tau$. On a finite lattice, say with an extent $L$, spatial correlations are limited by $L$, leaving us with: $\tau = \tau(L)$. We will distinguish between a power-law and an exponential relation:

$$\tau(L) \propto L^z, \quad \text{(critical slowing down)}$$
$$\tau(L) \propto e^{mL}, \quad \text{(super critical slowing down)}.$$ 

Because of the connection between autocorrelation time $\tau$ and statistical error $\epsilon$, theories in the parameter regime afflicted by super critical slowing down can only be simulated for small or moderate lattice sizes $L$, and extrapolation to large $L$ might or might not be possible.

Over many decades, research has been analysing the combination of theories and algorithms studying autocorrelations times for particular observables. For Markov chain simulation that satisfy detailed balance, large autocorrelations times are traced back to low eigenvalues of the transition matrix [4]. The latter paper offers a detailed study for lattice QCD and the important Hybrid Monte Carlo approach [5]. In theories that admit a characterisation of configurations by topology, such as QCD and CP(N) models, critical slowing down is often related
to slowly-evolving topological modes [6, 7]. More generally, modes with slowest de-correlation typically correspond to long-wavelength modes of physical fields. For a free scalar field theory, a combination of particular order of updating the fields and tuning of stochastic overrelaxation can significantly reduce critical slowing down [8]. Albeit this is per se an interesting finding, we here do not consider algorithms that need significant fine tuning for reducing autocorrelations.

To alleviate the “slow mode relaxation” issue, multi-grid methods have been proposed already in the late eighties [9]. For specific models, targeted solutions can be found that either eliminate critical slowing down or strongly reduce it. Those attempts are a based on a reformulation, and simulations include non-local updates. For the CP(N-1) model, which is plagued by the slow mode issue due to topological sectors, a complete absence of critical slowing down was reported in [10] for two dimensions. Cluster update algorithms [11, 12] generically possess a small dynamical critical exponent $z$ and thus provide a practical solution to the critical slowing down issue. Whenever a model allows a cluster reformulation, the performance cluster algorithms are hardly outperformed by any other approach and hence are the preferred simulation method.

Lattice theories that show spontaneous symmetry breaking in the infinite volume limit are particularly prone to super critical slowing down when simulated in the broken phase. Let $φ_z$ be the fields of such a theory with partition function

$$Z(β) = \int Dφ \exp{β S(φ)} ,$$

and $M(φ)$ the order parameter. For any finite lattice size, the symmetry implies that the expectation value of the order parameter, i.e., $⟨M⟩$ vanishes. In the broken phase, stochastically “important” configurations cluster in domains with $M(φ) \neq 0$ [13], and $⟨M⟩$ vanishes upon averaging over these relevant domains. Local update algorithms usually fail to induce transitions between these domains leading to super critical slowing down. Yang-Mills theories with a gauge group $SU(N \geq 3)$ fall into this important class of models [14]. Gauge symmetry prevents the definition of meaningful (gauge invariant) clusters and corresponding non-local update algorithms. We are hence turning to other more conventional simulation techniques.

A promising class of such algorithms are multi-canonical algorithms [15] and Wang-Landau techniques [16, 17]. Although the algorithmic differences and similarities between both methods have been studied in the literature (see e.g. [18]), both employ reweighing techniques with respect to a marginal distribution, which is at the heart of solving the issue of super critical slowing down. This has been firstly demonstrated by Torrie and Valleau [19] in a thermodynamics setting and later by Berg, Hansmann and Neuhaus for the ising model in [20].

At the root of super critical slowing down is the double-peak marginal distribution $P(M)$ of the order parameter, say the magnetisation $M$. Rather than leave it to importance sampling to transition between the two equally important phases, we calculate the partition function by integrating explicitly over the order parameter $M$ and stochastically over the remainder of the configuration space. To this aim, we exploit the identity

$$Z(β) = \int dm \rho(m) ,$$

$$\rho(m) = \int Dφ δ(m - M(φ)) \exp{β S(φ)} ,$$

where $δ$ is the Dirac $δ$–function. Thereby, $ρ$ is called the density-of-states. Density-of-states techniques have seen remarkable successes over the last decade ranging from a study of the QCD phase diagram at significant baryon chemical potentials [21], a recent study of the topological density in pure Yang-Mills theories [22] and the first proof of concept of solving a strong sign–problem using the $Z_3$ theory [23].

Key to the success of the density-of-states techniques is a robust method to estimate the density-of-states $ρ$ including control over its stochastic errors. In this paper, we explore the Linear-Log-Relaxation (LLR) method [24–26], which belongs to the class of the Wang-Landau techniques. The LLR method is based upon a systematical expansion of the marginal distribution $ρ(m)$ in a given $m$-interval leading to a stochastic non-linear equations for the expansion parameters (see below for details). In its lowest order, the LLR approach has similarities with the “multi-magnetic ensemble” method by Berg, Hansmann and Neuhaus [20]. The LLR-approach is also markedly different: it confines the MC simulation part to a window of size $2δ$ around a given value of the magnetisation $m_0$, which is a non-local constraint. We will be interested in the limit $δ \rightarrow 0$.

In this paper, we offer a systematic and large scale study of the phenomenon of critical slowing down using the LLR method. Since we are interested in simulation methods, which can applied universally to a wide range of lattice models, we benchmark our findings against those from a heatbath approach rather than a cluster algorithm, which would be anyhow the method of choice if applicable. We find evidence that super critical slowing down is absent (in line with the findings from a multi-canonical simulation [20]). We still find a correlation length that increases polynomial with the volume. We observe, however, that those correlations are strongly suppressed even at criticality.
II. UNDERSTANDING CRITICAL SLOWING DOWN

A. Accessing autocorrelations

The well-studied Ising model in a finite volume also serves here to illustrate the breakdown of importance sampling due to a failure of sampling the configuration space within an acceptable amount of computational resources. The purpose of this section is to quantify this breakdown for the popular Markov-Chain Monte-Carlo (MCMC) approach. We are particularly interested in the parameter dependence of failure, foremost its dependence on the system size. All numerical illustrations of this section are carried out using shockingly small lattice sizes. This illustrates the severeness of the issue: These small sizes are mandatory because of the rapid breakdown of ergodicity at even moderate lattice sizes.

Protagonists are the Ising spins \( s_x = \pm 1 \) associated with each lattice site \( x \) of the lattice of size \( V = L \times L \). We use periodic boundary with periodic boundary conditions [27] throughout the paper. Partition function \( Z \) and action \( S \) are given by

\[
Z = \sum_{\{s_x\}} \exp\{\beta S\}, \quad S = \sum_{\langle xy \rangle} s_x s_y , \tag{1}
\]

where the sum in the action extends over all nearest neighbours \( x \) and \( y \). Results for autocorrelations will depend on the algorithm. We therefore present details of the simulation here. We are employing the standard heatbath algorithm as benchmark:

1. Choose a site \( x \) of the lattice at random, and calculate the sum over the neighbouring spins:

\[
b_x = \sum_{y \in \langle xy \rangle} s_y .
\]

2. Define

\[
p_x = \frac{1}{1 + \exp\{-2\beta b_x\}} ,
\]

and choose \( s_x = 1 \) with probability \( p_x \) and set \( s_x = -1 \) otherwise.

3. Repeat both steps 1-2 above \( V \) times to complete one lattice sweep.

4. The spin configuration \( \{s_x\}_k \) after \( k \) sweeps is considered as part of a chain of configurations labeled by the Monte-Carlo time \( k = 1 \ldots N \). Define a sequence of random numbers for an observable \( f(\{s_x\}) \) by

\[
f_1 \to f_2 \to \ldots f_N , \quad f_i = f(\{s_x\}_i) .
\]

5. Obtain estimators for observables by

\[
f := \frac{1}{N_0} \sum_{i=1}^{N} f_i .
\]

6. Repeating steps 1-5 many times defines a random process for \( f \) itself. We denote the corresponding average by \([f]\). Note that \([f]\) is hence independent of, e.g., the random numbers used for a particular run, but does depend on \( N \). Approximate

\[
\langle f \rangle \approx [f] .
\]

A variable of particular interest is the magnetisation per spin

\[
\langle m \rangle = \left\langle \frac{1}{V} \sum_{x} s_x \right\rangle = \langle s_x \rangle ,
\]

which does not depend on the site \( x \) due to translation invariance. The corresponding elements of the chain of random variables are given by

\[
m_i = \frac{1}{V} \sum_{x=1}^{V} s_x^{(i)} , \tag{2}
\]

where \( s_x^{(i)} \) is the spin at site \( x \) of the configuration \( \{s_x\}_i \).

By the law of large numbers, we find

\[
\langle m \rangle = \lim_{N \to \infty} [m](N) .
\]
Any stochastic simulation, however, resorts to a finite length $N$ of the chain, and the central question is to what extent is the approximation

$$\langle m \rangle \approx [m]$$

valid?

To avoid a cluttering of notation, we preemptively use a result of the next subsection. By virtue of a symmetry argument, we have

$$\langle s_x \rangle = 0 \ , \ [m](N) = 0 \ , \ \forall N \ .$$

As usual, the error for the approximation (3) is given by the standard deviation

$$\epsilon^2 = [m^2] - [m]^2 = [m^2] \ .$$

We find

$$\epsilon^2 = \left[ \left( \sum_{i=1}^{N} \left( \sum_{\ell=1}^{N} m_i \right) \right) \right] = \frac{1}{N^2} \sum_{i=1}^{N} \sum_{\ell=1}^{N} [m_i m_\ell] \ .$$

Apparenty, the latter equation depends how the random variable $m_i$ is correlated to the variable $m_\ell$, and the average $m_i m_\ell$ is called autocorrelation. A key assumption here is that this correlation decreases exponentially with the distance $|k|$ between the positions in the chain:

$$[m_i m_\ell] = m_0^2 \exp \left\{ - \frac{k}{\tau} \right\} \ , \ k = |i-\ell|$$

$$m_0^2 := [m_i^2] \ ,$$

where $\tau$ is called autocorrelation time. This is expected to be the case for large separations $k$. A rather stark assumption is that the exponential behaviour dominates the double sum in (5). This assumption only can be justified afterwords in the numerical experiment but it seems to be the case for the parameter range explored in this paper. Inserting (6) into (5), the double sum can be performed analytically:

$$\epsilon^2 = \frac{m_0^2}{N^2} \sum_{i=1}^{N} \sum_{\ell=1}^{N} a^{i-\ell}$$

$$= \frac{m_0^2}{N} \frac{1+a}{1-a} - \frac{2a m_0^2}{N^2 (1-a)^2} \left( 1 - a^N \right) ,$$

$$a = \exp \{-1/\tau\} .$$

For a moderately sized autocorrelation time, we might find ourselves in a situation where we have $1 \ll \tau \ll N$. Expanding (7) yields for this case:

$$\epsilon^2 = \frac{2 m_0^2 \tau}{N} + \mathcal{O} \left( \frac{\tau^2}{N^2} \right) .$$

This the famous $1/\sqrt{N}$ law of MCMC simulations taking into account an autocorrelation time $\tau \gg 1$.

In case that the autocorrelation time is exceedingly large, we might face the ordering $1 \ll N \ll \tau$. Expanding (7) for this scenario yields an entirely different picture:

$$\epsilon^2 = m_0^2 \left[ 1 - \frac{N}{3 \tau} + \mathcal{O} \left( \frac{1}{N^2 \tau^2} \right) \right] .$$

In this case, the error is of order one, and we cannot expect that (3) yields a meaningful approximation. Note, however, that equation (10) still can provide information on the (large) autocorrelation time by virtue of the correction to the leading term even if $N \sim \tau$.

### B. Symmetry breaking and ergodicity

Partition function and action are invariant under a $Z_2$ transformation of the spins:

$$s_x \rightarrow (-1) s_x \quad \text{for} \quad \forall x .$$

This means that the configurations $\{s_x\}$ and $\{-s_x\}$ have the same probabilistic weight implying for any finite lattice size $V$:

$$\langle m \rangle = \langle s_x \rangle = -\langle s_x \rangle = -\langle m \rangle \ , \ \Rightarrow \ \langle m \rangle = 0 .$$

It also implies that, during the generation of the MCMC chain, the sequence

$$m_1 \rightarrow m_2 \rightarrow \ldots m_N \quad \text{and} \quad -m_1 \rightarrow -m_2 \rightarrow \ldots -m_N$$

occur with equal probability, meaning the average over chains vanishes as well, i.e.,

$$[m](N) = 0 .$$

The above symmetry enables us to cast each configuration of the MCMC chain into $Z_2$ classes. To this aim, we define

$$m_i = z_i \ [m_i] , \quad z_i = \pm 1 .$$

Thus, the mapping

$$\{s\}_i \rightarrow z_i$$

assigns a $Z_2$ sector (by virtue of the value of $z_i$) to each configuration. The symmetry transformation (11) maps each configuration onto a configuration with equal statistical weight of the other $Z_2$ sector.

The above conclusions are not necessarily true in the infinite volume limit $V \rightarrow \infty$. For infinite systems, the $Z_2$ symmetry ca be spontaneously broken. In fact, the Ising model is a prototype to explore this phenomenon. For $\beta > \beta_c$, the statistical system “freezes” in one of the $Z_2$ sectors with $\langle m \rangle \neq 0$. For $\beta < \beta_c$, we still find $\langle m \rangle = 0$ and the symmetry is realised. The critical value $\beta_c$ can be calculated analytically [28], and one finds:

$$\beta_c = \frac{1}{2} \ln \left( 1 + \sqrt{2} \right) \approx 0.440686 \ldots$$
This phenomenon is called spontaneous symmetry breaking and only applies to infinite volume systems.

Why should we be concerned with this phenomenon since we are only dealing with cases where $V$ is finite? The answer is that most importance sampling algorithms (if not all) for large enough $\beta \gg \beta_c$ and system size $L$, anticipate this phenomenon leading to the wrong result

$$[m](N < N_c) \neq 0$$

even at finite size $V$. The theorem of large numbers only guarantees $[m] = 0$ for $N \to \infty$, and on some practical applications $N_c$ can be unfeasibly large.

Let us study this statement in the context of an actual numerical simulation. We generate a chain for the magnetisation $m_i$ and for the $Z_2$ element $z_i$ as a function of the Monte-Carlo time $k$ for $\beta = 0.35$ and $L = 12$. We observe that system changes between $Z_2$ sectors during the run, which is expected since the $Z_2$ symmetry is unbroken at such small values of $\beta$. However, we realise that regions of positive (negative) $m_i$ cluster for some time. This indicate that we observe a significant autocorrelation time $\tau$ even at this small $\beta$. In order to quantify this, we present estimators for the autocorrelation functions for

$$m_k, z_k \text{ and } |m_k|.$$ 

Note that averages for $|m_k|$ and $|z_k|$ vanish but that for $[m_k]$ is non-zero due to the (semi-)positive nature of the observable. The simulation is carried out for a $12 \times 12$ lattice at $\beta = 0.35$, which is well placed within the symmetric phase with a moderate autocorrelation time. The simulation starts with a random spin configuration (hot-start) and initially discards 1000 configurations for thermalisation. The result for the autocorrelation functions is shown in figure 1 right panel. Our findings suggest that the autocorrelation functions of $m$ and $z$ are proportional (at least for sufficiently large a MC-time) difference, i.e.,

$$[m_i m_k] \approx m_z^2 [z_i z_k],$$

where $m_z^2$ is a parameter, which can be obtained comparing the fits in figure 1, right panel, and which is about 0.149. This finding signals that the autocorrelation of the centre sector drives the overall autocorrelation of the magnetisation.

We have systematically studied the error $\epsilon$ (as given by the equation (11)) for a $L = 12$ lattice size and the three $\beta$ values 0.3, 0.35 and 0.44. We fitted the theoretical expression for $\epsilon$ from (7) (the square root of (7) to be precise) to the numerical data. This yields an estimate for $m_z^2$ and the desirable autocorrelation time $\tau$. Our findings are summarised in figure 2 left panel. For beta 0.3 and 0.35 the observed autocorrelation time is small enough so that we can observe the characteristic $1/\sqrt{N}$ behaviour at large $N$. Note, however, that close to $\beta = \beta_c$, we observe a large autocorrelation time, which does not allow for the characteristic falloff for the range of $N$ explored. Note, however, that we still can get an estimate for $\tau$ by virtue of (7), which does not assume $N \gg \tau$.

The same Figure 2 right panel, shows the autocorrelation time as a function of $\beta$ for the three lattice size 12, 14 and 32. We observe that the autocorrelation time increases exponentially in all cases. Note, however, that
the slope of the increase changes around $\beta \approx \beta_c$ and is “steeper” for $\beta > \beta_c$, which corresponds to the symmetry broken phase in the infinite volume limit.

Equation (14) suggests that tunneling between $Z_2$ sectors is suppressed and that this suppression is at the heart of the practical ergodicity issue. For each step in of the MCMC chain, we can assign a probability $p$ that the configuration changes the $Z_2$ sector during this step. We then can calculate the average magnetisation $\langle z_i z_k \rangle$ analytically.

In a time series of $k + 1$ samples $z_i$, $i = 1 \ldots k + 1$ assume that $\ell$ transitions occur at $k$ possible locations (links between $i$ and $i + 1$). The probability for this event is given by

$$
\left( \frac{k}{\ell} \right) p^{\ell} (1 - p)^{k - \ell}.
$$

The contribution of this event to the autocorrelation function $\langle z_1 z_{k+1} \rangle$ is $(-1)^{\ell}$. Hence, we find

$$
\langle z_1 z_{k+1} \rangle = \sum_{\ell} \left( \frac{k}{\ell} \right) p^{\ell} (1 - p)^{k - \ell} (-1)^{\ell} = (1 - 2p)^k.
$$

(15)

Using the latter result in (14) and exploiting the connection to the autocorrelation time in (6), we find the connection between autocorrelation time $\tau$ and sector tunneling probability $p$:

$$
p = \frac{1}{2} \left( 1 - e^{-1/\tau} \right) \approx \frac{1}{2\tau}.
$$

(16)

The latter approximation holds for $\tau \gg 1$. For the example of the previous subsection, i.e., the heat-bath algorithm, a $12 \times 12$ lattice and $\beta = 0.35$, we found $\tau \approx 28$ leaving us with a tunneling probability of just $p \approx 1.8\%$.

C. Computational resources and precision

The strategy of comparing the performance of two different algorithms is as follows: we will agree at certain level of error $\epsilon^2$ and then ask the question how many “lattice sweeps” $N$ do we need to achieve this?

For the heatbath algorithm, we already worked out a connection between $\epsilon^2$ and $N$ (see (17)), and it depends on only two parameters, i.e., $m_0$ and $\tau$. It is to put this equation to the test. We have generated a time series of 6,000,000 magnetisations $m_k$, which we divide into subsequences of length $N$. For each subsequence, we calculate the average magnetisation

$$
m^{(\alpha)} = \frac{1}{N} \sum_{k=1}^{N} m_k^{(\alpha)},
$$

where $\alpha$ numbers the subsequences from 1 to $n_\alpha$, which fit into the series of 6,000,000 magnetisations. The error for the magnetisation estimator (4) is then estimated by

$$
\epsilon^2(N) \approx \frac{1}{n_\alpha} \sum_{\alpha=1}^{n_\alpha} \left[ m^{(\alpha)} \right]^2.
$$

(17)

Our numerical findings for $N = 10 \ldots 1500$ appear in figure 2, left panel, as solid lines. We show results for $\beta = 0.3$, $\beta = 0.35$, $\beta = 0.44$. Each curve is fitted by the theoretical prediction (7) with respect to only two fit parameters: $m_0$ and $\tau$. The agreement is excellent.

We can now ask the question: at least how many MCMC configurations do we need to achieve $\epsilon < 0.01$. For an answer, we use (17) with the readily obtained fit parameter $m_0$ and $\tau$. The agreement between theory and numerical data is that good that we can extrapolate to $N$ values bigger than 1500. We find that for our lattice size $L = 12$, $N$ has at least to be:

$$
\begin{align*}
\beta &= 0.30 : N = 10,800 \\
\beta &= 0.35 : N = 58,300 \\
\beta &= 0.44 : N = 10,460,000.
\end{align*}
$$

(18)

Note that the above $N$ values are vastly outside the fitting range of $N = 10 \ldots 1500$ and the application of (7) is an extrapolation. It is therefore in order to check the predictions (18). To this aim, we have created, for each $\beta$, an MCMC time series of length $N$ and have calculated the corresponding average magnetisation. We have repeated this 10 times. Since $\langle m \rangle = 0$, we expect these $m$ values to be scattered around zero with an error band $\epsilon = 0.01$ (one standard deviation). Our result is shown in figure 3. We observed the expected behaviour even for $\beta = 0.44$, for which $N = 10,460,000$. 

![Graph showing magnetisation vs count for different values of $\beta$.

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**FIG. 3.** Average magnetisation from a MCMC time series of length $N$ for three $\beta$ (see (18) for the $\beta$-$N$ pairs); $12 \times 12$ Ising model.
It appears that fitting ε-data with \( b_{0.46} \) is an economical way to calculate the autocorrelation time. We have done this for a range of \( \beta \) values and show the result in figure 2 right panel. We observe that the autocorrelation time \( \tau \) exponentially increases with \( \beta \). In the “symmetric phase” \( \beta \ll 0.44 \), the slope seems to be independent of the lattice size \( L \). In the “broken phase” \( \beta > 0.44 \), the picture changes: the slope of the exponential increase depends on the volume and is significantly bigger than in the symmetric phase. This signals a breakdown of validity of the heat-bath simulation for reasonable sized sample sizes \( N \).

D. Volume dependence and Critical Slowing Down

Of particular interest is to study the volume dependence of the autocorrelation time at give value of \( \beta \). For subcritical values, i.e., \( \beta < \beta_c \), we expect a power-law increase with the system size. This is simply because of that we operate with a local update algorithm, for which it is increasingly difficult to disorder a lattice configuration with increasing size. In the broken phase, i.e., \( \beta > \beta_c \), the picture is entirely different: the tunneling between centre-sectors is exponentially suppressed and a changing a \( Z_2 \) sector needs resources with exponentially increase with volume. In this subsection, we will verify this picture with unprecedented numerical evidence.

For extracting the autocorrelation time \( \tau \) for given size \( L \) and \( \beta \), we calculate the autocorrelation function as a function of the Monte-Carlo time \( t \). We fit the asymptotic tail to a the exponential form:

\[
C(t) = [m_0 m_t] \propto \exp\{-t/\tau\}.
\]

For small \( t \), we expect power-law corrections to the above functional form and, for large \( t \), the signal might be drowning in the statistical noise of the estimator. Let \( E(t) \) be the estimated error of the function \( C(t) \) at time \( t \). For the parameters \( L, \beta \) explored in this section, we only take data into with

\[
t > 200, \quad t < t_{\text{max}},
\]

where

\[
t_{\text{max}} : \text{largest } t \text{ with: } C(t) > 5E(t)
\]
or \( t_{\text{max}} = 2000 \) whatever is smaller. This is necessary to keep memory usage under control during the simulation. One of our many results is shown in figure 4 top panel. Parameters have been \( L = 16,32 \) and \( \beta = 0.43 \). Not all data are shown since the figure would become too crowded. The numerical data is well fitted by exponential form. Throughout, we monitor the \( \chi^2 \) of the fit. Errors for the fit parameter and hence the autocorrelation time is obtained by bootstrap. For the fits shown in figure 4 we obtained specifically

\[
\tau(L = 16) = 808.5(6), \quad \tau(L = 32) = 1794(1).
\]

We have repeated this analysis for \( L \in [8, 39] \) and \( \beta = 0.43, 0.44, 0.45, 0.46, 0.48 \). The results for the autocorrelation time \( \tau \) is shown in the same figure 4 bottom panel. We observe that \( \tau \) rapidly grows for \( \beta \) values instigating spontaneous symmetry breaking. We observe that the numerical data for \( \tau \) are well fitted by the formula

\[
\tau(L) = b_0 L^{b_1} \exp\{b_2 L\}.
\]

In the absence of the exponential \( (b_2 = 0) \), the formula describes a power-law growth of \( \tau \) with size \( L \) while, for \( b_2 > 0 \), the formula suggests an dominating exponential growth. The fits are also shown in the bottom panel of figure 4. They well describe the data. In particular, we find:
With the normalisation with the action $S$ because of the definition (14) and that of the partition in the usual way:

$$M_{b}$$

ability with which magnetisations $i.e.,$ symmetry transformation (11), the density is symmetric, $tional functions (19).

We thus find evidence that $b_{2}$ starts growing to non-zero values around the critical values $\beta \approx \beta_{c}$ for the phase transition. In the symmetric phase at $\beta = 0.43$, we find that the autocorrelation time $\tau$ approximately grows with the volume $L^{2}$.

III. REDUCED CRITICAL SLOWING DOWN WITH THE LLR METHOD

A. Brief introduction to the LLR approach

We are aiming to estimate the magnetisation $M$ with reliable errors over a wide spectrum of $\beta$-values stretching from the symmetric phase deep into the symmetry broken phase for $\beta \gg 0.44$. We start by defining the density-of-states $\rho(M)$ for the magnetisation:

$$\rho(M) = \frac{1}{Z} \sum_{\{s_{x}\}} \delta \left( M, \sum_{x} s_{x} \right) \exp \{ \beta S \} \quad (20)$$

with the action $S$ in [1]. The Kronecker delta is defined in the usual way:

$$\delta(i,k) = 1 \quad \text{for} \quad i = k \, , \quad 0 \quad \text{else.}$$

The magnetisation is then given by

$$\langle m \rangle = \frac{\sum_{M} M \rho(M)}{\sum_{M} \rho(M)} ,$$

$$M = -V, -V + 2, \ldots, V - 2, V . \quad (21)$$

With the normalisation

$$\sum_{M} \rho(M) = 1 \quad (22)$$

because of the definition [14] and that of the partition function $Z$ in [1], $\rho(M)$ can be interpreted as the probability with which magnetisations $M$ contribute to expectation values such as the one in (21). By virtue of the $Z_{2}$ symmetry transformation [11], the density is symmetric, $i.e.,$

$$\rho(-M) = \rho(M) ,$$

leading to $\langle m \rangle = 0$ as expected. In our numerical study we will $not$ exploit the above symmetry relation but rather will study the stochastic errors for our estimate for $\langle m \rangle$.

At the heart of the LLR approach is the expectation value

$$\langle f \rangle(a) = \frac{1}{N} \sum_{\{s\}} f(s) e^{\beta S + a \delta_{m}} W_{\delta}(m_{0}, m(s)) \quad (23)$$

$$m(s) = \sum_{x} s_{x} ,$$

where we here use a Heaviside function for the window function:

$$W_{\delta}(m_{0}, m(s)) = \left\{ \begin{array}{ll}
1 & \text{for } m_{0} - \delta \leq m(s) \leq m_{0} + \delta . \\
0 & \text{else.} \end{array} \right. \quad (24)$$

Note that $\langle f \rangle(a)$ depends also on the parameters $\delta$ and $m_{0}$, and $a$ is also called the LLR coefficient. You can obtain the density-of-states $\rho(m_{0})$ by carrying out the following steps:

1. For a given $\delta$ and $m_{0}$, solve the stochastic equation

$$\langle m(s) - m_{0} \rangle(a^{*}) = 0 \quad (25)$$

for $a$ (solution $a^{*}$), which depends smoothly on $m_{0}$ and $\delta$ for $m_{0} \in [-V, V]$.

2. Use

$$\frac{d}{dt} \ln \rho(m_{0}) = \lim_{\delta \to 0} a(\delta, m_{0}) \quad (26)$$

and evaluate (or estimate) $\rho(m_{0})$ up to a multiplicative factor by integrating the above equation.

3. Determine the multiplicative factor by normalising $\rho$ (see [22]).

The last step might be optional since a normalisation constant drop out of expectation values such as the one in [20].

As for the heat-bath MCMC approach, we are interested in the question: what type of precision can we achieve as a function of the invested computational resources. We therefore will critically investigate the parameter dependence of the numerical error.

Let us first comment on solving the stochastic equation of the type [25]. This task has been extensively studied firstly by Robbins and Monro [29] and then taken up by number of authors (see [30] for a review). If $F(a)$ is a noisy estimator for

$$f(a) := \langle m(s) - m_{0} \rangle(a) \quad (27)$$

Robbins and Monro propose an under-relaxed iterative approach. Starting with some $a_{1}$, consider the recursion

$$a_{n+1} = a_{n} - \alpha_{n} F(a_{n}) \quad (28)$$
with a sequence of positive weights \( \alpha_n, n = 1, 2, 3 \ldots \) satisfying
\[
\sum_{n=1}^{\infty} \alpha_n \to \infty, \quad \sum_{n=1}^{\infty} \alpha_n^2 \to \text{finite}.
\]
The sequence converges with probability one to the solution \( a^* := a_\infty \). A particular sequence was suggested by Robbins and Monro:
\[
\alpha_n = \frac{\kappa}{n}.
\]
The algorithm reaches \textit{asymptotically} the optimal convergence rate of \( 1/\sqrt{n} \), but the initial (low \( n \)) performance crucially depends on the sequence. Chung \cite{32} and Fabian \cite{33} showed that optimal convergence is reached with the choice:
\[
\alpha_n = \frac{1}{f'(a^*)n}.
\]
This choice, however, hinges on the solution \( a^* \). For the specific problem at hand, i.e., \( \beta = 0.30 \), we can, however, find a good value \( \kappa \). For small enough \( \delta \), the marginal for the magnetisation \( m \) in the window \([m_0-\delta, m_0+\delta]\) is Poisson distributed, i.e., \( \propto \exp\{-a^*m\} \). Together with the 'reweighting' factor \( \exp\{am\} \) in \( \delta \), the \( m \) distribution becomes flat for values \( m \) inside the window. We then find with \( \alpha_n \), the definition \( \delta \) and the solution \( \beta \):
\[
f'(a^*) = \langle (m(s) - m_0)m(s) \rangle(a^*) = \frac{1}{\delta^2 + 1} \sum_{m=-\delta}^\delta m^2.
\]
The latter hold for \( \delta \gg 1 \), which would also be the result if the degrees of freedoms have a continuous domain of support. Note that by the nature of the task at hand \( \beta \), \( f'(a^*) \) does not depend on the solution \( a^* \). We arrive at the iteration that we will study in the remainder of the paper:
\[
a_{n+1} = a_n - \frac{3}{\delta^2 n} F(a_n).
\]  
We put the above iteration to the test for a \( V = 12 \times 12 \) lattice, \( \beta = 0.3 \), \( m_0 = \text{INT}(0.8V) \) and several \( \delta \) values. The estimator \( F(a) \) is obtained by 20 successive lattice sweeps. Our findings for the error \( \epsilon_n \) in the LLR coefficient \( a \) as a function of the Robbins Monro iteration time \( n \) is shown in figure 5. We performed 1,000 independent Robbins Monro runs to estimate the error for \( \epsilon_n \). We find optimal convergence behaviour already for \( n > 200 \). The error for small \( \delta \) are smaller than those for large \( \delta \). This is expected since for larger \( \delta \) the window function is wider and hence includes more spin in the averaging.

\[\text{FIG. 5. Left: The error in the LLR coefficient} \quad a \text{ as a function of the number of Robbins-Monro iterations} \quad n \] 

The following study is done for the 2D Ising model on a \( 32 \times 32 \) lattice. The objective is to find the amount of 'lattice sweeps' is needed to calculate the magnetisation \( m \) to a given accuracy. In the last section, we saw that the heat bath algorithm needs a rapidly increasing amount of resource if \( \beta \) approaches the regime of a spontaneously broken symmetry.

\[\text{Our simulations parameters are "ball park" figures and are not fine tuned.}\]
1. We use a step function as window function $m \in [m_0 - \delta, m_0 + \delta]$ with $\delta = 8, 16, 24, 32$.

2. We perform 10,000 Robbins Monro iterations for each $m_0$ and for each $\delta$ leaving us with an estimate for the LLR parameter $a(\delta)$. We perform a quadratic fit for extrapolating to $\delta \to 0$ and set: $a = a(0)$.

3. Each double expectation value is estimated with 20 lattice sweeps.

4. We generate LLR parameters $a$ for 63 values of $m_0$, i.e., $(m_0)_k = -32^2 + k \times 32$, $k = 1 \ldots 63$.

5. For each $m_0$, we generate 80 potential LLR parameters $a_i$ for the subsequent statistical analysis.

We will measure resource in units of 'lattice sweeps' ($ls$), i.e., one resource unit corresponds to $V$ spin updates. This choice allows to measure resource independent of hardware employed for the calculations. All algorithms studied here - heat bath update, cluster algorithms, LLR method - uses 'lattice sweeps' at low level of the calculation. Although Ising spin updates are low cost, the 'lattice sweep' might be the most expensive computational element for other systems such as gauge theories with fermions (QCD) where a lattice sweep could be defined by a Hybrid Monte-Carlo trajectory.

To generate the above data set for the LLR coefficients (steps 1-4), the resources needed are

$$4 \times 20 \times 10,000 \times 63 \, ls = 5.04 \cdot 10^7 \, ls.$$  \hspace{1cm} (30)

From this data set, we can already estimate expectations values of functions of the magnetisation, and the objective here is to estimate the precision with which we can calculate $\langle m \rangle$ (which equals zero for a simulation with infinite resources). To this aim, we will repeat the calculation 80 times. This, the analysis uses the resources of $5.04 \cdot 10^7 \times 80 \, ls = 4.032 \cdot 10^9 \, ls$, which must not be confused with resource (30) needed to produce one sample result.

The density of states $\rho$ for the magnetisation $m$ is obtained by integration of the LLR coefficient:

$$\rho(m) = \exp\left\{ \int_0^m a(m') \, dm' \right\}. \hspace{1cm} (31)$$

The normalisation is arbitrarily chosen to be $\rho(0) = 1$. Expectations values are then obtained by a second integration, e.g.,

$$\langle m \rangle = \int m \rho(m) \, dm / \int \rho(m) \, dm. \hspace{1cm} (32)$$

Early studies \cite{23,24} used a trapezium rule and summation, which leads to an accumulation of error for increasing $m$. Representing the function $a(m)$ by high degree polynomial and performing the integrations (semi-) analytically has proven very successful \cite{26,34,35}. One can prove that the density of states for Ising model is an even function in $m$ by virtue of its $Z_2$ symmetry. Correspondingly, the LLR coefficient $a(m)$ is an odd function. A numerical approach exploiting this observation would approximate $a(m \geq 0)$ by polynomial of odd powers of $m$. This would lead to the exact result $\langle m \rangle = 0$.

The prime objectives here is to avoid any assumptions on symmetry and to observe to what extent the exact result $\langle m \rangle = 0$ is obtained. For this purpose, we approximate $a(m)$ over the full domain by polynomial containing...
even and odd powers of $m$. We find that a polynomial of degree 16 represents the numerical data for $a$ very well.

The result for $\rho(m)$ on a logarithmic scale is shown in figure 6. Error bars are obtained by the bootstrap method:

1. For each $m_0$, calculate a set of $n_B$ LLR coefficients from independent runs. We have chosen here $n_B = 60$.
2. For each of the (discrete) $m_0$ choose an LLR coefficient out of the $n_B$ possibilities.
3. Fit a polynomial of degree 16 to the data.
4. Perform the integration (31) analytically and obtain one sample for $\rho(m)$.
5a. Repeat this procedure many times and calculate the average for $\rho(m)$ and the standard deviation (error bar).

Step 5a gives rise to the graphs in figure 6 left panel. We find that for $\beta = 0.25, 0.30, 0.40$ the density-of states is maximal at $m = 0$ making $m = 0$ the most likely magnetisation. We also observe that, for a finite $L = 32$ lattice, the curve for $\beta = 0.44$ develops a double peak structure, which is characteristic for the spontaneous breakdown of symmetry. We expect that for increasing lattice size, the $\beta$ for which the double peak structure occurs will approach $\beta_c$ in (13).

We are here not primarily interested in the density of states $\rho$ but the expectation value of the magnetisation

$$m = M/V = \frac{1}{V} \sum_x z_x .$$

In this case, we replace step 5a by:

5b. For the sample $\rho(m)$, calculate the two integrals in (31) analytical and, thus, obtain a sample value for $\langle m \rangle$. Repeat this procedure many times and calculate the average for $\langle m \rangle$ and the standard deviation (error bar).

Figure 6 left panel, shows the (log of the) density of states as a function of the intrinsic magnetisation $m = M/L^2$. For the finite volume $L = 32$, we see that the most likely magnetisations are at $m \neq 0$ for $\beta = 0.44$. This is a precursor of spontaneous symmetry breaking. Increasing the volume, it is expected that this bifurcation moves up in $\beta$ to approach $\beta_c$. (13) in the infinite volume limit.

Having calculated the density of states, we estimated the magnetisation $m$ using (32). The precision with which the exact result $\langle m \rangle = 0$ is recovered depends on the quality of the symmetry $\rho(m) = \rho(-m)$. Our result for the error of $m$ is shown in figure 6 right panel, as a function of $\beta$, where we have kept fixed the number of Robbins Monro iterations and the bootstrap copies.

We find a moderate increase with increasing $\beta$, which can be explained by the larger variation of $\rho(m)$ with $m$ due its peak structure, which makes it harder to control the numerical precision of the integration over $m$ in the integrals of (23).

C. Autocorrelations and density-of-states

The so-called double expectation values such as in (17) are at the heart of the LLR approach since they ultimately give rise to $a$ and hence the density of states (see (25)). These expectation values can be viewed as ordinary Monte-Carlo expectation values, and, as such, they are susceptible to autocorrelations of the Markov chain.

We already established that there is a close link between spontaneous symmetry breaking and the exploding autocorrelation time for local update algorithms operating close to criticality. We expect that the double expectation values are much less affected by this phenomenon simply because they are not operating a close to criticality “most of the time”.

We first note that the double expectation values depend on a number of parameters, which are not present in a standard heat bath simulation. There is the LLR parameter $a$ which adds a term $a \sum_x s_x$ to the action. For $a \neq 0$ this parameter acts like a magnetic field, which breaks the $Z_2$ symmetry $s_x \rightarrow -s_x$. Secondly, the window function $W(m_0, m(s))$ (24) is part of the probabilistic measure. It restricts spin configurations to values of the magnetisation $m(s)$ close $m_0$. This means
that this factor also breaks the $Z_2$ symmetry as long as $m_0 \neq 0$. Note, however, that for $m_0 = 0$, the solution of the stochastic equation is $a = 0$ precisely because of the $Z_2$ symmetry. We thus expect that the calculation of $\rho(m \approx 0)$ might be affected by long autocorrelations. Note that for most of the observables in the broken phase, $\rho(m \approx 0)$ might be an entirely suppressed domain of integration for the integrals in e.g. (32). In this case, these autocorrelations have little impact on the precision of the calculation.

In a first step, we studied the autocorrelation time for the action and the spin-spin correlation function for different values of $m_0$, the centre of the window function:

\[
\begin{align*}
\text{action: } & \sum_{(x,y)} s_x s_y , \\
\text{spin-spin: } & s_x s_{x+L/2} .
\end{align*}
\]

Our findings are summarised in figure 7 left panel. Indeed, we observe that those autocorrelations are highest close to $m_0 = 0$ where the system can have critical behaviour.

Since the magnetisation is constrained to a region around $m_0$ in the LLR simulation, autocorrelations of the magnetisation are indeed very small. In search of an observable susceptible to longest autocorrelations, we introduce the Fourier transform of the magnetisation:

\[
\hat{M}(p_x, p_y) = \sum_{x} s_{x, y} \cos \left( \frac{2\pi}{L} (x p_x + y p_y) \right) ,
\]

For $p_x = 0, p_y = 0$, this quantity becomes the magnetisation, i.e., $M = \hat{M}(0)$. Another "infrared" observable, similarly prone to autocorrelations but unconstrained by the LLR approach, is $\hat{M}$ for the lowest momenta with either $p_x = 1, p_y = 0$ or $p_x = 0, p_y = 1$. The choice of these observables is motivated by the common observation that low-momentum modes typically have the slowest relaxation/ decorrelation rate in local, translationally invariant quantum field theories. We thus study the autocorrelation time for the observable

\[
M_1 \equiv \hat{M}(1,0) = \sum_{x, y} s_{x, y} \cos \left( \frac{2\pi}{L} x \right) .
\]

To this end, we firstly estimate the autocorrelation function $C(t)$ of $M_1$ and extract the autocorrelation time by analysing the exponential decrease at large values of $t$. If $t$ is too large, statistical noise drowns the signal. If $\sigma(t)$ is the standard deviation of the estimator for $C(t)$, we only use data with

\[
C(t) > 5 \sigma(t) .
\]

At small values of $t$, $C(t)$ is not well represented by an exponential function, which only hold asymptotically. We proceed as follows: starting at $t = t_0 = 0$, we fit an exponential function to the data and obtain the $\chi^2$/dof. We then systematically increase $t_0$ until $\chi^2$/dof falls below 0.8 for the first time. We thus extract the autocorrelation time $\tau$ from the fit:

\[
a_0 \exp\{-t/\tau\} .
\]

Figure 8 shows the correlations function $C(t)$ for a $32^2$ lattice and for four values of $\beta$ within the dynamically generated domain of support. Repeating this procedure for lattice sizes between $L = 8$ and 48, we find the result shown in figure 9. We indeed observe that the autocorrelation times for $M_1$ increase with increasing lattice size $L$, but not nearly to the extent as we have seen those for the heatbath simulation and the magnetisation $M$.

The central question is whether or not these autocorrelations times increase exponentially with $L$. In search of an answer, we have employed the same fit (19) of the data as in the case of the heatbath result. Of particular interest is the coefficient $b_2$, which indicates super critical slowing down for $b_2 > 0$. Our findings are summarised in the table below:

| $\beta$ | $\ln(b_0)$ | $b_1$ | $b_2$ |
|---------|-------------|-------|-------|
| 0.44    | $-1.28(1)$  | 1.965(4) | 0.0038(2) |
| 0.46    | $-1.26(3)$  | 1.942(3) | 0.025(1) |
| 0.48    | $-1.495(6)$ | 2.080(3) | 0.036(1) |
| 0.50    | $-2.194(7)$ | 2.484(4) | 0.029(2) |

Table II. Results of the fitting of the lattice size dependence of the autocorrelation time in LLR simulations with a product of power law and exponential functions (19).
FIG. 9. Autocorrelation time for the observable $M_1$ as a function of the system size $L$ for four values of $\beta$ and for the worst case scenario $m_0 = 0$.

FIG. 10. A comparison of the dependence of autocorrelation time for the observable $M_1$ on lattice size $L$ for the LLR approach for $m_0 = 0.8$ for several values of $\beta$. The quality are less convincing especially for $\beta = 0.48$. Here, figure 9 shows two fits: the exp-powerlaw fit for $\beta = 0.44$ and a power-law fit $b_2 = 0$. Both fits reasonable well present the data. We are carefully optimistic that any exponential growth is a quite small rate implying that autocorrelation times are manageable for realistic lattice sizes. Higher precision data and perhaps larger lattice sizes are needed to evidence this at a quantitative level.

As detailed above, only the double-expectation values for $m_0 = 0$ are afflicted by criticality since, for $m_0 \neq 0$, the $Z_2$ symmetry is explicitly broken by the window function and an LLR-coefficient $a \neq 0$. Nevertheless, it is important how the autocorrelation times scale with the lattice size $L$. In the broken phase, say for $\beta > 0.45$, the marginal distribution for the magnetisations peak at rather large values $M/V \approx \pm 0.9$. For generic observables with a broad domain of support from large portions of the domain of magnetisation, the dominant contributions from the LLR integration over the magnetisation raises from the region around $M/V \approx \pm 0.9$. Hence, we studied the volume dependence of the observable (34) as a function of the lattice size $L$ at $m_0 \neq 0$. The results for $m_0 = 0.9$ are shown in figure 10 in the double-log scale in comparison with the $m_0 = 0$ data. We observe that autocorrelation times are orders of magnitudes smaller than in the $m_0 = 0$ case. Most importantly however, we find that the increase of the autocorrelation time with size is at most polynomial in $L$ and for $\beta$ values away from its critical value even sub-polynomial. Log-log scale plot illustrates this in a particularly clear way, mapping any power-law dependence to a straight line. Therefore plots of functions that grow faster than a power of $L$ appear as bending upwards from a straight line, whereas plots of functions with sub-polynomial growth are bending down from a straight line.

This is an important finding since observables that receive their dominant contribution from the regions of large magnetisation are not affected by super critical slowing down.

IV. DISCUSSION AND CONCLUSIONS

Local update algorithms for Markov chains of a given sample size tend to fail exploring the full configuration space, and hence ergodicity, for theories in the regime of a spontaneously broken symmetry. In this regime, the marginal distribution of the order parameter exhibits several regions of equal stochastic importance but importance sampling generically selects only one of these regions and fails to transition between. Consequently, the autocorrelation function rises exponentially with the system size (super critical slowing down). A second question arising is whether the autocorrelation length still rises polynomial, say at criticality (critical slowing down). We addressed both issues in this study.

Our approach is to decompose the configuration space into the order parameter as a collective coordinate and the hyperspace orthogonal to this mode. Wang-Landau techniques (and the LLR method, in particular) are ideally placed to integrate the slow mode explicitly while the integration over the hyperspace is done stochastically using MCMC techniques.
In this paper, we used a simple two-dimensional Ising model to demonstrate to explore the performance of the LLR method. For the Ising model, there are efficient model-specific cluster algorithms that not only eliminate super critical slowing down but also largely alleviate critical slowing down as witnessed by a small dynamical critical exponent. Note, however, that cluster algorithms are only available for very specific models. The present research targets algorithms that work for a large class of models ‘out of the box’ without major fine-tuning.

For the Ising model, the mode that exhibits the longest autocorrelation time is the global magnetisation, that is, the sum of all spins. We expect that for all models that are well described by the Landau theory of phase transitions the global order parameter will always have the longest autocorrelation time. To confirm this, we also studied the autocorrelation time for the mode with lowest nonzero momentum $p = \frac{2\pi}{L}$, where $L$ is the linear system size. Our approach also resembles, to some extent, lattice QCD simulations in fixed topological sectors [7]. Indeed, global topological charge is known to be the observable with longest autocorrelation time in lattice QCD.

We found that the LLR algorithm has a potential for solving the issue of super critical slowing down for most observables. Only observables that are sensitive to the marginal distribution around $M \approx 0$, no matter how small it is, might be affected by critical slowing down. We only know one such observable: the order-disorder interface tension. We still see a polynomial rise of the autocorrelation time with the volume at criticality (and, hence critical slowing down), but we find that at a quantitative level the autocorrelation time is reduced by orders of magnitude when compared with that of a heatbath simulation with the same system size (see figure 11).

As a next step, it would be interesting to check whether explicit integration over more than one observable using higher-dimensional generalisation of the LLR algorithm could result in further reduction of computational time. It is also worth exploring whether the application of LLR method to fermionic systems could reduce ergodicity issues related to zeroes of the fermionic determinant. Finally, in a recent paper [37] it was suggested that normalising flows can eliminate the need to integrate the density of states over $m$ altogether, thus yielding an even larger speed-up for Monte-Carlo simulations. It would be interesting to see to what extent normalising flows can further reduce the critical slowing down in our situation.

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[1] H. J. Rothe, Lattice Gauge Theories, 4th ed. (WORLD SCIENTIFIC, 2012) https://www.worldscientific.com/doi/pdf/10.1142/8229
[2] K. Binder and D. Heermann, *Monte Carlo simulation in statistical physics*, 4th ed., 80 (Springer, Berlin ; Heidelberg ; New York ; Barcelona ; Hong Kong ; London ; Milan ; Paris ; Tokyo, 2002) pp. XII, 180 S.
[3] F.M.Dekking, *A Modern Introduction to Probability and Statistics Understanding Why and How* (Springer, London, 2005).
[4] S. Schaefer, R. Sommer, and F. Virotta (ALPHA), Nucl. Phys. B 845, 93 (2011) arXiv:1009.5228 [hep-lat]
[5] S. Duane, A. D. Kennedy, B. J. Pendleton, and D. Roweth, Phys. Lett. B 195, 216 (1987).
[6] C. Bonati and M. D’Elia, Phys. Rev. E 98, 013308 (2018) arXiv:1709.10034 [hep-lat]
[7] R. Brower, S. Chandrasekharan, J. W. Negele, and U. J. Wiese, Phys. Lett. B 560, 64 (2003) arXiv:hep-lat/0302005
[8] L. Horváth and A. Kennedy, Nuclear Physics B 510, 367 (1998)
[9] E. P. Stoll, J. Phys.: Cond. Matt. 1, 6959 (1989)
[10] U. Wolff, Nucl. Phys. B 832, 520 (2010) arXiv:1001.2231 [hep-lat]
[11] R. H. Swendsen and J.-S. Wang, Phys. Rev. Lett. 58, 86 (1987)
[12] U. Wolff, Phys. Rev. Lett. 62, 361 (1989)
[13] K. Binder, *The Frontier of Particle Physics* pp. 1009–1944.
[14] A. Billoire, T. Neuhaus, and B. Berg, Nuclear Physics B 396, 779 (1993)
[15] F. Wang and D. P. Landau, Phys. Rev. Lett. 86, 2050 (2001)
[16] F. Wang and D. P. Landau, Phys. Rev. E 64, 056101 (2001) cond-mat/0107006.
[17] C. Junghans, D. Perez, and T. Vogel, *Journal of Chemical Theory and Computation* 10, 1843 (2014) PMID: 26580515, https://doi.org/10.1021/ct500077d
[18] G. Torrie and J. Vallecio, *Journal of Computational Physics* 23, 187 (1977)
[19] B. A. Berg, U. Hansmann, and T. Neubea, *Physical Review B: Condensed Matter; (United States)* 47:1 (1993), 10.1103/PhysRevB.47.497
[20] Z. Fodor, S. D. Katz, and C. Schmidt, JHEP 03, 121 (2007) arXiv:hep-lat/0701022
[21] S. Borsanyi and D. Sexty, Phys. Lett. B 815, 136148 (2021) arXiv:2101.03383 [hep-lat]
[22] K. Langfeld and B. Lucini, Phys. Rev. D 90, 094502 (2014) arXiv:1404.7187 [hep-lat]
[23] K. Langfeld, B. Lucini, and A. Rago, Phys. Rev. Lett. 109, 111601 (2012) arXiv:1204.3243 [hep-lat]
[24] K. Langfeld, B. Lucini, R. Pellegrini, and A. Rago, Eur. Phys. J. C 76, 306 (2016) arXiv:1509.08391 [hep-lat]
[25] K. Langfeld, PoS LATTICE2016, 010 (2017) arXiv:1610.09856 [hep-lat]
[26] E. Ising, Zeitschrift fur Physik 31, 253 (1925)
[27] L. Onsager, Phys. Rev. 65, 117 (1944)
[28] H. Robbins and S. Monro, Ann. Math. Stat. 22, 400 (1951)
[29] H. J. Kushner and G. G. Yin, *Stochastic Approximation Algorithms and Applications* (Springer, 1997).
[30] J. R. Blum, Ann. Math. Stat. 25, 382 (1954)
[31] K. L. Chung, Ann. Math. Stat. 25, 463 (1954)
[32] V. Fabian, Ann. Math. Stat. 39, 1327 (1968)
[33] N. Garron and K. Langfeld, Eur. Phys. J. C 76, 569 (2016) arXiv:1605.02709 [hep-lat]
[34] N. Garron and K. Langfeld, Eur. Phys. J. C 77, 470 (2017) arXiv:1703.04649 [hep-lat]
[35] O. Francesconi, M. Holzmann, B. Lucini, and A. Rago, Phys. Rev. D 101, 014504 (2020) arXiv:2203.01243 [hep-lat]