Method of Simulating the Dynamics of Long-Range Interacting Spin Systems

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We introduce a novel method that eases the daunting task of simulating dynamics in spin systems with long-range interaction. Our Monte Carlo simulations of the long-range Ising model for the nonequilibrium phase ordering dynamics in two spatial dimensions perform $\sim 10^3$ times faster than the standard approach. Importantly, this enables us to establish agreement with the theoretical prediction for the time dependence of domain growth, in contrast to previous numerical studies. Our method can easily be generalized to applications in other systems.

Generic models of statistical physics exhibiting a transition from disordered to ordered states have been proved to be instrumental for understanding the dynamics in diverse fields, from species evolution \cite{1} to traffic flow \cite{2}, from economic dynamics \cite{3} to rainfall dynamics \cite{4}. An extensively used paradigm is the Ising model with nearest-neighbor (NNIM) interaction \cite{5,6}. Even the electrostatic forces, polarization forces, etc. Hence, a more complete picture calls for employing models that consider long-range interactions.

The simplest generic model system is the long-range Ising model (LRIM), which on a $d$-dimensional lattice is described by the Hamiltonian

$$\mathcal{H} = -\sum_i \sum_{j<i} J(r_{ij}) s_i s_j, \quad \text{with} \quad J(r_{ij}) = \frac{1}{r_{ij}^\gamma},$$

where spins $s_i = \pm 1$, $r_{ij}$ is the distance between the spins at site $i$ and $j$, and $J(r_{ij})$ is the interaction strength. The model exhibits a para- to ferro-magnetic phase transition. Naturally, simulations of such systems with long-range interaction are computationally far more expensive than its short-range counterpart. For equilibrium studies, the advent of various collective updates based on the Swendsen-Wang cluster algorithm \cite{7} allows one to perform efficient Monte Carlo (MC) simulations \cite{8,9}. Conversely, for understanding the nonequilibrium ordering kinetics following a quench from the high-temperature disordered phase into the ordered phase below the critical temperature $T_c$, one is restricted to use only local moves, viz., single spin flips. This makes MC simulations of ordering kinetics in LRIM severely expensive even with present-day computational facilities, and therefore, they have rarely been attempted \cite{10}.

The understanding of ferromagnetic ordering kinetics in NNIM is well developed \cite{4,5}. It is characterized by formation and growth of domains of like spins and is a scaling phenomenon, i.e., the characteristic length scale $\ell(t)$ at time $t$ follows the Lifshitz-Cahn-Allen (LCA) law \cite{11}: $\ell(t) \sim t^{1/2}$ which can be derived by considering that $\ell(t)$ grows via reduction of the curvature $1/\ell(t)$ of the domain walls. Similarly for the LRIM the growth is likely to be driven by interactions between domain walls. Assuming this growth as a scaling phenomenon and using an “energy scaling” argument it has been predicted that \cite{12,13}

$$\ell(t) \propto t^\alpha = \begin{cases} t^{1/2} & \sigma < 1 \\ \ln(t)^{1/2} & \sigma = 1 \\ t^{1/2} & \sigma > 1 \end{cases}$$

i.e., (i) in the “truly” long-range regime for $\sigma < 1$, the growth exponent $\alpha$ is $\sigma$ dependent, (ii) at the crossover point $\sigma = 1$, the growth follows the LCA law with a multiplicative logarithmic correction, and (iii) for $\sigma > 1$, LRIM behaves asymptotically as the NNIM with $\alpha = 1/2$. There exist few attempts to confirm these predictions via numerical solution of Ginzburg-Landau-type \cite{14} or Langevin-type \cite{15,16} dynamical equations. The only available results from MC simulations \cite{17} in this regard tackles the expensive calculation of the local energy involving all the spins by using a cut-off distance for $J(r_{ij})$ in \cite{18}. Importantly, in disagreement with \cite{17,18}, $\alpha$ is found there to be no different than in NNIM for all $\sigma$, thus suggesting a universal nonequilibrium behavior. In equilibrium it is well established both theoretically \cite{19,20} and in simulations \cite{21,22} that critical exponents are not universal. In the $d = 2$ LRIM, for $\sigma < 1$ the critical exponent $\eta$ takes its mean-field value, followed by an intermediate range $1 < \sigma < \sigma_\times$ where it is $\sigma$-dependent, and for $\sigma > \sigma_\times$ it behaves like in the NNIM. The value of the crossover point $\sigma_\times$ is still disputed \cite{23} and predicted to be $\sigma_\times = 2$ \cite{24} or $\sigma_\times = 7/4$ \cite{21}. In this Letter, thus along with the establishment of a novel and efficient approach of doing MC simulations for the kinetics of LRIM in $d = 2$ without involving any cut-off, we aim further to check the $\sigma$-dependence of the growth exponent $\alpha$.

In a standard Metropolis simulation \cite{25} for kinetics of LRIM one attempts to flip a randomly chosen spin $s_i$ with probability $p_i = \min[1, \exp(-\Delta E_i/k_B T)]$, where $k_B(=
1) is the Boltzmann constant, \( T \) is the temperature and \( \Delta E_i = E_a - E_b \) is the change between the energy before \((E_a)\) and after \((E_b)\) the flip. The aim of our approach is to abstain us from doing the expensive calculation of \( \Delta E_i \) at every attempted move. Instead we introduce a local pseudo heatbath, assigned to each spin, and only update other spin flips to this pseudo heatbath.

When simulating a long-range interacting system using periodic boundary conditions (via minimum-image convention), one encounters strong finite-size effects. To circumvent this we use Ewald summation [11, 24, 26]. Qualitatively all the subsequent results can be reproduced by using the simpler minimum-image convention, albeit with stronger finite-size effects. As a starting point we consider a square lattice with all spins aligned (up or down). Then we do the Ewald summation to calculate the effective interaction \( J(r_{ij}) \) on the chosen lattice \( (\text{dependent on } L \text{ and } \sigma) \). Now, for each spin \( s_i \), we assign a local pseudo heatbath

\[
h_i = \sum_{j \neq i} J(r_{ij})s_j. \tag{3}
\]

Conveniently, for an aligned configuration all \( h_i \) are identical. To prepare an initial configuration that mimics a paramagnetic phase \( (T \gg T_c) \) we flip half of the spins choosing randomly. For each spin flip we update the pseudo heatbath of all other spins, i.e., if a spin \( s_i \) is flipped the pseudo heatbath \( h_j \) of any other spin \( s_j \) accounts for a change of \(-2J(r_{ij})s_i\). This generates the initial random configuration with updated \( h_i \).

In the next stage we do the Metropolis simulations at any given temperature using the Boltzmann criterion with the advantage of the stored \( h_i \). Using Eq. (3) one can write down the change in energy due to an attempted flip of a randomly chosen spin \( s_i \) as

\[
\Delta E_i = E_a - E_b = 2s_i \sum_{j \neq i} J(r_{ij})s_j = 2s_i h_i. \tag{4}
\]

This enables us to calculate the acceptance probability \( p_i \) just by a multiplication of stored variables. Now, only if the attempted flip is allowed (by comparing \( p_i \) with a random number) we update \( h_j \) of every other spin \( s_j \). Updating the \( h_j \) here is the bottleneck. However, one does this update less often than the spin-flip attempts.

| \( \sigma \) | 0.4 | 0.6 | 0.8 | 1.0 | 1.5 |
|----------|-----|-----|-----|-----|-----|
| Clocks \((10^{10})\) | 1.42(4) | 2.1(2) | 3.7(2) | 5.3(3) | 8.1(3) |

**TABLE I.** Average number of POSIX clocks needed by our method for different values of \( \sigma \). Estimations are made from simulations of LRIM with \( L = 1024 \) averaged over 20 initial realizations, running up to \( 10^4 \) MCS. Corresponding clocks for the standard method are \( 5.6(1) \times 10^{13} \). All simulations were run on an Intel Xeon CPU E5-2640 V4.

Since we aim to study the ordering kinetics, after preparing the initial configurations, we quench the system to a temperature \( T_q = 0.1T_c(\sigma) \). Like in the standard method, here also the unit of time is one MCS, demonstrating the ferromagnetic ordering in LRIM with \( L = 1024 \) for different \( \sigma \). Only the up spins (+) are marked.

![Evolution snapshots at different times](image)

**FIG. 1.** Evolution snapshots at different times (in units of MCS), demonstrating the ferromagnetic ordering in LRIM with \( L = 1024 \) for different \( \sigma \). Only the up spins (+) are marked.

We now check the scaling of the morphology-
characterizing two-point equal-time correlation function $C(r, t) = \langle s_i s_j \rangle - \langle s_i \rangle \langle s_j \rangle$ and its Fourier transform, the structure factor $S(k, t) = \int dr C(r, t) e^{ikr}$. Figure 2(a) presents $C(r, t)$ at different times for $\sigma = 0.6$, showing the signature of a growing length scale with time. The multiplicative scaling during the growth is confirmed by the data collapse as shown in Fig. 2(b), on plotting the $C(r, t)$ against $r/\ell(t)$ where the length scale $\ell(t)$ is extracted from the criterion $C[r = \ell(t), t] = 0.5$. The data at large $r/\ell(t)$ for the latest time seems to show some discrepancy attributed to finite-size effects. However, the scaling of the structure factor $S(k, t)$ that forms a basic assumption when deriving the theoretical growth laws for LRIM [14, 15], is confirmed convincingly as shown in Fig. 2(c). Similar respective behavior is observed when scaled $C(r, t)$ and $S(k, t)$ at the same time is plotted for different $\sigma$ in Fig. 3. The slower decay of $C(r, t)$ for smaller values of $\sigma$ could be an indication of the inverse relation of the growth exponent $\alpha$ with $\sigma$, as found in Eq. 2. Contrasting, the scaled $S(k, t)$ for different $\sigma$ in Fig. 3(b) show reasonably good overlap. The solid lines in Fig. 2(c) and Fig. 3(b) depict the consistency of the data with the Porod tail [28]; $S(k, t) \sim k^{-(d+1)}$ at large wave number $k$.

The multiplicative scaling of the morphology-characterizing functions indeed suggests the presence of scaling of the growing length scale. Hence, shifting our focus on the growth exponent $\alpha$ in Fig. 4(a) we present the time dependence of the length scale $\ell(t)$ for $\sigma = 0.6$. The behavior is certainly not $\sim t^{1/2}$ (shown by the dashed line), but in fact the data for all $L$ follow the predicted behavior of $t^{1/(1+\sigma)}$ until they show deviations due to finite-size effects. This already indicates that the underlying scaling behavior is indeed consistent with [2]. Nevertheless, to further strengthen the claim and to gauge the effect of a finite system size we call for a finite-size scaling (FSS) analysis [29, 30] which recently has been successfully employed in kinetics of other systems [31–33]. Quantifying the growth including an initial crossover time $t_0$ and length $\ell_0 = \ell(t_0)$ one can write down the ansatz $\ell(t) = \ell_0 + A(t - t_0)^{\alpha}$ and construct a FSS function $Y(y) = (\ell(t) - \ell_0)/(L - \ell_0)$ with the scaling variable $y = (L - \ell_0)^{1/\alpha}/(t - t_0)$. In the scaling regime one expects $Y \sim y^{-\alpha}$. Thus on plotting $Y$ as a function of $y$ for different $L$ one must observe a data collapse with $Y \sim y^{-\alpha}$ behavior for large $y$ provided $\alpha$ is chosen appropriately. We did this exercise for different $\sigma$ choosing $\alpha$ from [2]. However, not all of them are presented here, but rather a representative plot for $\sigma = 0.6$ is shown in the inset of Fig. 4(a). The collapsed data is consistent with the underlying master curve $Y \sim y^{-\alpha}$. Considering the col-

FIG. 2. (a) Correlation functions $C(r, t)$ at different times for $\sigma = 0.6$. (b) Demonstration of the scaling of $C(r, t)$ as a function of $r/\ell(t)$ for the same times as in (a). (c) Scaling plots for the structure factor $S(k, t)$. The solid line there corresponds to the Porod tail behavior of $S(k, t) \sim k^{-3}$.

FIG. 3. (a) Scaled correlation function $C(r, t)$ at $t = 100$ MCS for different $\sigma$ as mentioned. (b) Same as (a) but for the scaled structure factor $S(k, t)$. The solid line again corresponds to the Porod tail.

FIG. 4. (a) Demonstration of the scaling behavior $\ell(t)$ for $\sigma = 0.6$ as a function of $t$ for different system sizes. The data at large $t$ seems to follow the predicted behavior of $t^{1/(1+\sigma)}$ until they show deviations due to finite-size effects. The solid line again corresponds to the Porod tail behavior $\ell(t) \sim t^{1/2}$. The inset (b) depicts the consistency of the data with the underlying master curve $Y \sim y^{-6}$.
with multiplicative logarithmic correction: \( (t \ln t)^{1/2} \), albeit a power-law growth with \( \alpha > 1/2 \) cannot unprejudiced be ruled out. However, in accordance with \( \sigma \) for \( \sigma = 1.5 \) in the post-crossover regime \( (\sigma > 1) \) the growth appears to be \( t^{1/2} \), as expected for the NNIM. To consolidate the visual validation we also performed for each case least-square fits of prediction \( \alpha \) and verified the predicted exponent values \( \alpha \). In the inset of Fig. \( \alpha \) we show a plot of the length scale obtained from simulations using different cut-off radii \( r_c \) in Eq. \( \alpha \) for \( \sigma = 0.6 \). For the largest \( r_c \), the data follows \( t^{1/(1+\sigma)} \) behavior as is observed without any cut-off, whereas the cases with smaller \( r_c \) obey the LCA law. Thus, in conjunction with the previously reported simulation \( \alpha \) one can infer that the use of a relatively small \( r_c \) make the spins interact only on short range leading to \( \sigma \)-independent growth exponents.

To conclude, we have studied the kinetics of ferromagnetic ordering using the long-range Ising model in \( d = 2 \) spatial dimensions via Monte Carlo simulations using a novel method. We have introduced a pseudo heat bath for the calculation of the local energy that helps to reduce the expensive calculation of local energy involving all the spins at every step. Our approach speeds up the simulation by a factor of \( \sim 10^3 \) compared to the standard way. This enables us to simulate systems as big as \( 2048^2 \) spins without using any cut-off radius in the distance-dependent power-law interaction. Results obtained from our simulations are the first confirmation of the theoretical prediction \( \alpha \) for the growth laws in the long-range Ising model \( \alpha \). We have also demonstrated that the inappropriate use of a cut-off radius in the local-energy calculation may lead to a different growth exponent, explaining the mismatch between previous simulation results \( \alpha \) and theory.

In equilibrium, the long-range Ising model has a dimension-dependent crossover behavior of the critical exponents \( \alpha \), while in nonequilibrium the prediction \( \alpha \) is expected to be independent of the dimension. In this light, we take the ordering kinetics of the \( d = 3 \) case as our next endeavor to check this dimension independence \( \alpha \). Our method shall trigger interests to explore other aspects associated with ordering phenomena in the long-range Ising model \( \alpha \), viz., aging and related dynamical scaling \( \alpha \). The generic simple feature of the method shall ensure its facile adoptions to nonequilibrium simulations of other models, viz., \( g \)-state Potts and clock models. In view of the delicate cut-off dependence \( \alpha \), it would also be interesting to revisit the ordering phenomenon in long-range liquid crystals \( \alpha \). Although originally designed for simulating dynamics, our method should be proven to be handy for equilibrium simulations of systems with long-range interactions, for which there (currently) exist no cluster algorithms, e.g., (lattice-) polymers \( \alpha \).

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