Simulation of Potts models with real $q$ and no critical slowing down

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A Monte Carlo algorithm is proposed to simulate ferromagnetic $q$-state Potts model for any real $q > 0$. A single update is a random sequence of disordering and deterministic moves, one for each link of the lattice. A disordering move attributes a random value to the link, regardless of the state of the system, while in a deterministic move this value is a state function. The relative frequency of these moves depends on the two parameters $q$ and $\beta = \frac{1}{T}$. The algorithm is not affected by critical slowing down and the dynamical critical exponent $z$ is exactly vanishing. We simulate in this way a $3D$ Potts model in the range $2 < q < 3$ for estimating the critical value $q_c$ where the thermal transition changes from second-order to first-order, and find $q_c = 2.620 \pm 0.005$.

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I. INTRODUCTION

$Q$-state Potts model [1] is perhaps one of the simplest, non-trivial models in statistical mechanics. A broad set of techniques has been brought to bear on it in a variety of disciplines and it has been the subject of considerable theoretical attention over the last two decades (for a review, see [2]).

This model is theoretically well defined for any real or complex value of $q$ [3]. In particular, the limit $q \to 1^+$ corresponds to the random percolation problem and the limit $q \to 0^+$ has a fundamental role in enumerating the spanning trees of a graph [3]. Two-dimensional conformal field theory [4] suggests exact formulae for the critical indices and for other universal quantities as continuous functions of $q$ in the range $0 < q < 4$. Another interesting problem involving non-integer $q$ in three-dimensional Potts models is the determination of the universal value $q_c$ for which the thermal transition changes from second-order to first-order. A variety of techniques have been used [5–9], which locate $q_c$ in the range $2 < q_c < 3$. All these methods require extrapolations in $q$ because the standard simulations work only at integer values of $q$. Reweighting techniques [10,11] and transfer matrix methods [12] allow to estimate some thermodynamic functions [3] in a wider range of $q$, however there is no way to evaluate correlation functions there.

In this paper we remove this limitation by constructing a new Monte Carlo (MC) algorithm which works for any real $q > 0$. Although the time required for a sweep through the system grows faster than its size because at some step of the algorithm nonlocal information is required, the simulations are not affected by a critical slowing down and the dynamical critical exponent $z$ is exactly zero. We test the reliability of the method by comparison with some exact results for $2D$ Potts model at criticality. We probe its effectiveness by performing large scale MC simulations of a three-dimensional Potts model for estimating the universal value $q_c$.

II. THE ALGORITHM

Starting with the Hamiltonian $H = -\sum_{\langle ij \rangle} \delta_{\sigma_i \sigma_j}$ where the site variable $\sigma_i$ takes the values $\sigma_i = 1, 2, \ldots, q$, with $\langle ij \rangle$ ranging over the links of an arbitrary lattice or graph $\Lambda$, one can write the $q$-state Potts model partition function $Z = \sum_{\{\sigma\} \in \Lambda} e^{-\beta H}$ in the Fortuin Kasteleyn (FK) random cluster representation [3]

$$Z = \sum_{G \subseteq \Lambda} W(G) = \sum_{b,c} \Omega(b,c) v^b q^c,$$  (1)

where $v = e^\beta - 1 = \frac{q}{1-q}$, the summation is over all spanning subgraphs $G \subseteq \Lambda$, $W(G) = v^b q^c$ is their weight, expressed in terms of the number of $b$ edges of $G$, called bonds, and the number $c$ of connected components or FK clusters; $\Omega(b,c)$ is the number of subgraphs with $b$ bonds and $c$ clusters. This representation now defines a model for any real or complex $q$.

In principle, one could directly use Eq.(1) to define a Metropolis algorithm working for positive non integer $q$ [13], but this is a difficult problem to simulate because, for each proposed change of a link, the number $c$ of FK clusters, a nonlocal property, must be determined. Large lattices require a huge amount of CPU time. As a matter of fact, such a method has been applied only to two-dimensional systems, where special topological relations can be used [13].

Our strategy is different. We start by considering a useful identity which can be derived using the methods described in Ref. [14].

Let $l$ be any link of $\Lambda$. Denote by $\{G_l^+\}$ the set of spanning subgraphs where $l$ is a bond and by $\{G_l^-\}$ those in which this bond is missing. We have $Z = Z_l^+ + Z_l^-$, with $Z_l^\pm = \sum_{G_l^\pm} W(G_l^\pm)$. Introducing a bond variable $\alpha_l$ equal to 1 when $l$ is a bond and 0 otherwise yields

$$\langle \alpha_l \rangle = \frac{Z_l^+}{Z}.$$  (2)
The same quantity can be evaluated in a different way by addition of a bond to each graph of type $G^l_t$. There are two kinds of missing bonds. Those joining two different clusters, called potential bridges, are picked out by a variable $\beta_l$ which takes the value 1 only on them and is zero otherwise; their addition lowers the number $c$ of FK clusters. We have

$$\beta_l = 1 \Rightarrow W(G^l_t) \frac{v}{q} = W(G^l_t) .$$

The remaining missing bonds, described by a similar variable $\gamma_l$, join two sites of the same cluster; their addition keeps $c$ invariant, then

$$\gamma_l = 1 \Rightarrow W(G^l_t) v = W(G^l_t) .$$

Combining Eqs. (2), (3) and (4) yields $\langle \alpha_l \rangle = \frac{\varepsilon}{q}(\beta_l) + v(\gamma_l)$ which is the wanted identity. Since of course $\alpha_l + \beta_l + \gamma_l = 1$, we can rewrite it as

$$\langle \alpha_l \rangle = p \langle \alpha_l \rangle + \frac{p}{q} \langle \beta_l \rangle + p \langle \gamma_l \rangle ,$$

where the weighting factors can now be interpreted in terms of probabilities. The idea is now to regard this identity as the limit of a recursion relation of the type

$$\pi^{(n+1)}_l = p \pi^{(n)}_l + \frac{p}{q} \beta^{(n)}_l + q \gamma^{(n)}_l ,$$

where $\pi^{(n+1)}_l$ is the probability of having a bond on the link $l$ in the configuration $G^{(n+1)}$. It is expressed as a state function $\langle \alpha_l, \beta_l, \gamma_l \rangle$ of the same link in the $G^{(n)}$ configuration. This generates a Markov process $\cdots \rightarrow G^{(n)} \rightarrow G^{(n+1)} \cdots$ where the equilibrium distribution yields Eq. (5). This stochastic chain fulfills two important conditions: i) there is a non-zero probability of going from any configuration to any configuration in a single sweep through $\Lambda$, ii) the equilibrium distribution maps to itself as Eq. (5) is kept invariant by the process. One can then argue that detailed balance is satisfied.

To see it directly, assume for instance that in the $n^{th}$ configuration $l$ is a potential bridge ($\beta^{(n)}_l = 1, \ G^{(n)} = G^l_\Lambda$) which is promoted to a bond in the $(n+1)^{th}$ configuration ($\alpha^{(n+1)}_l = 1, \ G^{(n+1)} = G^l_\Lambda$). The transition rate is $P(G^l_\Lambda \rightarrow G^l_\Lambda) = \frac{\varepsilon}{q}$. Conversely Eq. (5) yields $P(G^l_\Lambda \rightarrow G^l_\Lambda) = 1 - p$. Then, according to Eqs. (5) and (6),

$$P \left( G^l_\Lambda \rightarrow G^l_\Lambda \right) = \frac{W(G^l_\Lambda)}{W(G^l_\Lambda)} \frac{W(G^l_\Lambda)}{W(G^l_\Lambda)} \frac{W(G^l_\Lambda)}{W(G^l_\Lambda)}$$

as detailed balance requires. The same conclusion can be reached in all the other cases.

A straightforward, preliminary, implementation of the recursion relation (5) is the following: i) go over each link $l \in \Lambda$ of the configuration $G^{(n)}$ and generate a pseudo-random number $X_l$ uniformly distributed from 0 to 1. ii) Create a bond on $l$ only in the following two cases: a) $X_l < \frac{\varepsilon}{q}$ and $l$ is a bond ($\alpha_l = 1$) or a missing bond joining two sites of the same FK cluster ($\gamma_l = 1$); b) $X_l \geq \frac{\varepsilon}{q}$ and $l$ is a potential bridge ($\beta_l = 1$). This generates uniquely the configuration $G^{(n+1)}$.

Let $q > 1$ for definiteness. It is worth noting that when $X_l < \frac{\varepsilon}{q}$ the algorithm adds a bond to $l$ regardless of which configuration $G^{(n)}$ we are dealing with. Similarly, when $X_l > p$ no bond is added. In the remaining cases ($\frac{\varepsilon}{q} \leq X_l \leq p$). The value attributed to $l$ (bond or no bond) is unambiguously determined by $G^{(n)}$.

Inspecting all the cases leads to the following better implementation of the algorithm:

Step 1: Pick a link $l \in \Lambda$ and generate a pseudo-random number $0 \leq X_l \leq 1$.

Step 2: Update the link according to the following scheme

| move | current state | new state |
|------|--------------|-----------|
| a) $X_l < \frac{\varepsilon}{q}$ | any bond | bond |
| b) $X_l > p$ | any no bond | no bond |
| c) $\frac{p}{q} \leq X_l \leq p$ | $\alpha_l = 1$ bond | $\beta_l = 1$ no bond |
| | $\gamma_l = 1$ bond | |

Step 3: Return to step 1.

The first two moves do not need any information on the state of the system: they just disorder it. The last one is a purely deterministic move; its only effect is to put a bond whenever a link joins two sites of the same cluster. It requires distinguishing between the two kinds of missing bonds ($\beta_l = 1$ or $\gamma_l = 1$). One can infer this nonlocal property by identifying the connected components of the configuration, like in the Swendsen-Wang (SW) algorithm [15]. This cluster reconstruction is time demanding, however it gives a complete information on the state of the missing bonds of the whole lattice. As the update proceeds through the lattice this amount of information is progressively lost because of disordering moves (the deterministic moves never change c). We may partly keep track of the cluster structure by relabelling the cluster indices whenever a disordering move creates a bond between two of them. Cluster reconstruction is truly necessary only when a deterministic move touches a missing bond of a putative single cluster where some bond has been erased by previous disordering moves.

Because of non locality, the number of operations involved every MC step is $\propto N^a$, where $N$ is the number of links and $1 < a \leq 2$. The efficiency of the algorithm depends crucially on the actual number of cluster reconstructions per sweep. In our 3D simulations reported below the fraction of links requiring cluster reconstruction was about 3% with a decreasing trend for larger lattices.
TABLE I. The decorrelation time $\tau$ of the new algorithm for the critical 2D Ising model for different linear lattice sizes $L$ is compared with the same quantity of the SW algorithm $\tau_{sw}$ and with the upper bound $\tau_o$. The definition of $\tau$ and the $\tau_{sw}$ data are taken from Ref. [18].

| $L$ | $\tau$ | $\tau_{sw}$ | $\tau_o$ |
|-----|---------|-------------|---------|
| 8   | 2.65(3) | 5.17696(32) | 3.3869  |
| 16  | 3.16(5) | 6.5165(12)  | 4.5158  |
| 32  | 3.69(6) | 8.0610(18)  | 5.6448  |
| 64  | 4.3(1)  | 9.794(4)    | 6.7737  |

III. CORRELATION TIMES

An unusual feature of the described algorithm is the presence of randomly distributed disordering moves. The mean number of links subjected to disordering moves in a single sweep is $N_p r$ with $p_r = 1 + \frac{\xi}{q} - p$. For instance, in the Ising model ($q = 2$) at criticality more than 70% of the links are disordered every sweep. It is now easy to find an upper bound for the mean number $\tau$ of MC steps needed to generate effectively independent configurations. After $n$ sweeps the mean number of links which do not have yet undergone a disordering move is $N(1-p_r)^n$. When this number is of the order of 1 all the links have been touched by a disordering move and the upper bound $\tau_o \geq \tau$ is given by the obvious relation $N(1-p_r)^{\tau_o} \sim 1$, i.e. $\tau_o = -\frac{\log N}{\log(1-p_r)}$. Thus the dynamical exponent $z$ is 0, as critical slowing down manifests itself by the power law $\tau \propto N^z$ at the critical temperature where a second order phase transition occurs [16]. A numerical estimate of the decorrelation time of the dynamics of the new algorithm for the critical Ising model on a square lattice is reported in Tab. 1. Note that the actual value of $\tau$ does not saturate the upper bound and is much smaller than the analogous quantity of the SW algorithm.

The new algorithm proves also useful in fighting against another dynamical problem which one deals with in the case of first-order transitions, namely the exponentially fast suppression of the tunnelling between metastable states with increasing lattice size. To reduce this type of slowing down the multi-canonical MC algorithm has been proposed [17] also the method of simulated tempering [18] proves useful [19]. In a few numerical tests for two-dimensional models with $q = 7$ and $20 \geq L \geq 100$ we found that the tunnelling time of the canonical algorithm described in the present paper grows with the system size $V$ as $\tau \propto V^\alpha$ with $\alpha = 1.03 \pm 0.03$, like in an optimal variant [20] of the multi-canonical method, but with a smaller proportionality factor.

The reason of this performance is that the random moves accelerate the tunnelling between order and disorder. The drawback is that the new algorithm is non-local, so the CPU time grows like $V^b$ with $b > 1$; for instance in the present case we found $b \sim 1.85$. Thus the new algorithm cannot certainly be recommended for integer $q$, although at a first-order transition it performs much better than any local canonical algorithm.

IV. SIMULATIONS

As a first, simple, application of the new algorithm we tested the reliability of our code by checking a percolation property of Potts model on a square lattice which is supposed to be exact in the range $0 < q < 4$, namely that the mean frequency of active bonds $\langle \alpha_l \rangle$ at criticality (corresponding to $\nu = \sqrt{q}$ in Eq. (1)) should coincide, in the thermodynamic limit, with the random percolation value, i.e. $\langle \alpha_l \rangle = \frac{1}{2}$, irrespective of the value of $q$ [21]. We simulated critical Potts models in a $128 \times 128$ square lattice with $q$ ranging from 1.5 to 3.5. In all the cases the mean number of bonds was compatible with the exact result. Finite size effects of this observable, which are visible on smaller lattices, allow to evaluate the critical thermal exponent $\nu$ as a function of $q$. This could be used to check a conjectural formula suggested by the 2D conformal field theory [18]. We plan to study this problem in a future publication.

The new algorithm allows us to deal with an important issue of three-dimensional Potts model, namely the estimate of the tricritical point $q_c$ in the range $2 < q_c < 3$, where the thermal transition changes from second-order for $q \leq q_c$ to first-order for $q > q_c$. Many different techniques have been used to locate this point [22]. We applied a method very similar to that described by Lee and Kosterlitz [23] by computing the double histogram $N(b, c)$ of bond and cluster number distribution in a cubic lattice of volume $L^3$ at a given $\beta$ and $q$ and then extrapolating the data to nearby values. Using Eq. (1) we can write
TABLE II. The simulations were performed on cubic lattices of side $L$ at the values of $q$ and $\beta$ listed below. MCS is the number of Monte Carlo steps considered.

| $L$  | $q$  | $\beta$  | MCS  |
|------|------|-----------|------|
| 12   | 2.70 | 0.52270   | 3.0 $10^7$ |
| 13   | 2.70 | 0.52270   | 3.0 $10^7$ |
| 14   | 2.75 | 0.52721   | 3.3 $10^7$ |

\[
N(b, c; \beta, q, L) = N \Omega(b, c) \frac{v^b q^c}{Z},
\]

where $N$ is the number of MC sweeps. We can trade the number of bonds $b$ for the energy per site $E$ using the relation $E = -\frac{\ln N(b, c)}{v^b}$ . Near a first-order transition the histogram $P(E) = \sum_q N(b, c)/N$ has a characteristic double peak structure corresponding to the ordered and the disordered phase. A suitable reweighting through Eq. (8) of the energy distribution yields the value $\beta_c(L, q)$ where the two peaks at $E_1(\beta)$ and $E_2(\beta, L)$ are of equal height. A typical plot of the quantity $A(E, q; \beta_c, L) = \sum c \ln(N(b, c)/N)$ is shown in Fig.1. A useful estimator of the interface free energy between the ordered and the disordered phase [21] is given by

\[
\Delta F(q, L) = A(E_m, q; \beta_c, L) - A(E_1, q; \beta_c, L),
\]

where $E_m$ is the local maximum which separates the two dips at $E_1$ and $E_2$ (see Fig.1). At a first-order transition $\Delta F(L)$ increases monotonically with $L$ and is expected to vanish at the tricritical point. Extrapolating the numerical data both in $\beta$ and $q$ one may locate this point. The region of reliable extrapolation [21] is $O(1/L^3)$ for both $\beta$ and $q$. This does not cause a problem for $\beta$, since it can be adjusted continuously, but $q$ cannot in standard simulations, being by necessity an integer value. Actually Lee and Kosterlitz performed their simulations at $q = 3$ and found that the extrapolated data become too noisy for $|\delta q| > 0.3$ [22]. In our case we can directly evaluate the range of reliable extrapolations. Indeed the main advantage of the algorithm described in this paper is that now also $q$ can be adjusted continuously.

Our simulations were performed on three different lattices as listed in Tab. II. The statistics is good since in all the cases the mean flipping time between coexisting states was no larger than thirty MC steps. The errors were calculated by gathering the histogram $N(b, c)$ every $10^6$ MC steps and then performing a standard analysis.

In all the cases the energy histogram showed a double peak structure, providing a direct evidence of the first-order nature of the transition for these values of $q$ (see Fig.2). This yields the upper bound $q_c < 2.7$. In shorter simulations at $q = 2.6$ we found no trace of a double peak structure. This suggests $q_c > 2.6$. Using the reweighting method we estimated the values $\beta_c(L, q)$ where the two peaks are of equal height for each $L$ and for few values of $q$ near $q = 2.7$ and the corresponding values of $\Delta F$. The results are reported in Fig.3.

A further reweighting up to $q = 3$ allowed to compare the extrapolated data to those coming from a similar extrapolation of standard SW simulations at $q = 3$. This comparison showed that the range of reliable extrapolations is $|\delta q| < 0.25$. It has to be noted that we could not use for this comparison the high precision data of Ref. [22], because the energy distribution in terms of spin variables used there does not coincide with that expressed in terms of bond variables used by necessity in the present approach. In particular the $\beta_c(L, q)$’s are shifted and our $\Delta F(q, L)$ is always smaller.

Simple finite size scaling considerations suggest [8] that near $q_c$ the interface free energy has the simple form $\Delta F(q, L) \sim (q - q_c)^2 L^a$ which fits very well to our data (see Fig.3). To within our numerical accuracy $a = 4.8 \pm 0.1$ and $q_c = 2.620 \pm 0.005$. This agrees with the value $q_c = 2.55 \pm 0.12$ obtained in the large $q$ expansion of the latent heat [8]. Lee and Kosterlitz [8] extrapolating $q = 3$ data found a smaller value, $q_c = 2.45 \pm .010$. The difference could be due to the fact that extrapolations with $|\delta q| > 0.25$ gives an overestimate of $\Delta F$ (this is already visible in Fig.1). Other approximate methods give even smaller values: real space renormalization group methods [8] yield $q_c \sim 2.2$ while an Ornstein-Zernike approximation [8] gives $q_c \sim 2.15$.

V. CONCLUSIONS

This work provides a new MC algorithm to simulate ferromagnetic $q-$ state Potts model which has two very unusual features: it works for any real $q > 0$ and does not suffer of any critical slowing down. The former property is an obvious consequence of the fact that it is based on the Fortuin Kasteleyn random cluster representation, where $q$ acts as a continuous parameter. The latter is more tricky and is due to the implementation of the algorithm with a random sequence of disordering moves,
randomly distributed over the lattice. There is no reason to believe that this disordered mechanism is specific to Potts model and it would be very interesting to try to implement it in other, more general MC methods. A drawback of the new algorithm is that it is non-local, so the CPU time of a single sweep grows with the volume $V$ as $V^b$ with $1 < b < 2$, thus it is not recommended for integer $q$, where the SW algorithm works with $b = 1$. Actually at a first-order transition the new algorithm performs better than the SW method, but there the multi canonical MC algorithms are more suitable.

It is straightforward to extend the new algorithm in order to take into account quenched bond randomness, provided that all the couplings are ferromagnetic. On the contrary, generalising to systems with frustrations seems a rather difficult task, because it is not obvious how to define in this case the FK clusters for non-integer $q$.

We used such an algorithm to study the region $2 < q < 3$ of a three-dimensional Potts model in order to estimate the critical value $q_c$ for which the thermal transition changes from second to first-order. We obtain a rather precise estimate compared to other methods, the reason being that all the other methods are based on extrapolations from integer values of $q_c$, while the new algorithm simulates the system at nearby values of $q_c$.

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