Cluster Simulations of Loop Models on Two-Dimensional Lattices

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We develop cluster algorithms for a broad class of loop models on two-dimensional lattices, including several standard O(n) loop models at $n \ge 1$. We show that our algorithm has little or no critical slowing-down when $1 \le n \le 2$. We use this algorithm to investigate the honeycomb-lattice O(n) loop model, for which we determine several new critical exponents, and a square-lattice O(n) loop model, for which we obtain new information on the phase diagram.

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From the beginning of the theory of critical phenomena, two models have played a central role: the *q*-state Potts model [1,2] and the O(n) spin model [3,4]. The parameter *q* or *n* is initially a positive integer, but the Fortuin-Kasteleyn (FK) representation [5] and the loop representation [6] show, respectively, how the models can be extended to arbitrary real or even complex values of *q* and *n* [7]. In particular, for *q*, n > 0 the extended model has a probabilistic interpretation as a model of random geometric objects: clusters [8] or loops [9], respectively. These geometric models play a major role in recent developments of conformal field theory [10] via their connection with stochastic Loewner evolution (SLE) [11,12].

Since nontrivial models of statistical mechanics are rarely exactly soluble, Monte Carlo simulations have been an important tool for obtaining information on phase diagrams and critical exponents [13]. Unfortunately, Monte Carlo simulations typically suffer from severe critical slowing-down, so that the computational efficiency tends rapidly to zero as the critical point is approached [14]. An important advance was made in 1987 with the invention of the Swendsen-Wang (SW) algorithm [15] for simulating the ferromagnetic Potts model at positive integer q, based on passing back and forth between the spin and FK representations. The SW algorithm does not eliminate critical slowing-down, but it radically reduces it [16]. Since then, many similar "cluster algorithms" have been devised, based on this principle [17] of augmenting the original spin model with auxiliary variables and then passing back and forth. But cluster algorithms have traditionally been limited to integer q, since they make essential use of the spin representation.

This limitation was first overcome in 1998 by Chayes and Machta [18], who devised a cluster algorithm for simulating the FK random-cluster model at any real $q \ge$ 1. For loop models, by contrast, efficient simulation at noninteger *n* has remained out of reach; to our knowledge only two Monte Carlo simulations at $n \ne 1$ have ever been published [19,20], and they used local algorithms. (Instead, PACS numbers: 05.10.Ln, 05.50.+q, 64.60.Cn, 64.60.Fr

numerical transfer-matrix techniques have been employed [21].) As a result, many open questions remain: for instance, the nature of the phase transition is unclear for the $n > \frac{3}{2}$ honeycomb-lattice loop model with vacancies [22]; and the phase diagrams and universality classes of loop models on lattices other than honeycomb are largely unexplored [23].

In this Letter we shall set forth a broad (but nonspecific) generalization of the Chayes-Machta idea, and then show how it can be adapted to provide a cluster algorithm for simulating loop models on two-dimensional lattices at any real $n \ge 1$. We shall furthermore present numerical evidence that this cluster algorithm has little or no critical slowing-down when $1 \le n \le 2$. Finally, we shall use this algorithm to obtain new results for the phase diagram of a loop model on the square lattice.

We begin by considering a generalized random-cluster (RC) model, defined as follows: Let G = (V, E) be a finite graph with vertex set V and edge set E. A configuration of the model is specified by a subset $A \subseteq E$ (the set of "occupied bonds"), and the partition function is

$$Z = \sum_{A \subseteq E} \left(\prod_{e \in A} \nu_e \right) \left(\prod_{i=1}^k W(H_i) \right), \tag{1}$$

where H_1, \ldots, H_k are the connected components of the graph (V, A); here $\{v_e\}$ are nonnegative edge weights, and $\{W(H)\}$ are nonnegative weights associated to the connected subgraphs H of G. The model (1) reduces to the FK random-cluster model if W(H) = q for all H; other special cases include an FK representation for the Potts model in a magnetic field, $W(H) = q - 1 + e^{h|V(H)|}$ [24], and the loop models to be discussed below.

Now let *m* be a positive integer, and let us decompose each weight W(H) into *m* nonnegative pieces, any way we like: $W(H) = \sum_{\alpha=1}^{m} W_{\alpha}(H)$. The first step of our generalized Chayes-Machta algorithm, given a bond configuration *A*, is to choose, independently for each connected component H_i , a "color" $\alpha \in \{1, ..., m\}$ with probabilities $W_{\alpha}(H_i)/W(H_i)$; this color is then assigned to all the vertices of H_i . The vertex set V is thus partitioned as $V = \bigcup_{\alpha=1}^{m} V_{\alpha}$. It is not hard to see that, conditioning on this decomposition, the bond configuration is nothing other than a generalized RC model with weights $\{W_{\alpha}(H)\}$ on the induced subgraph $G[V_{\alpha}]$, independently for each α .

We now have the right to update these generalized RC models by any valid Monte Carlo algorithm. One valid update is "do nothing"; this corresponds to the "inactive" colors of Chayes and Machta [18]. Of course, we must also include at least one nontrivial update. The basic idea is to have at least one color for which the weights $W_{\alpha}(H)$ are "easy" to simulate. The original Chayes-Machta choice, when W(H) = q for all H, is to take $W_{\alpha}(H) = 1$ for one or more colors α (the so-called "active" colors); the corresponding model on $G[V_{\alpha}]$ is then independent bond percolation, which can be trivially updated. Since we must have $W_{\alpha}(H) \leq W(H)$, this works whenever $q \geq 1$.

We hope that this explication-extension of the Chayes-Machta framework will inspire other researchers to invent diverse algorithms of Chayes-Machta type. Here we would like to exhibit two such families of algorithms, for simulating a general class of "loop models" on two-dimensional lattices. More precisely, the models we have in mind should be called *Eulerian-subgraph models*, because the connected components are not necessarily loops; we recall that a bond configuration A is called Eulerian if every vertex has even degree (i.e., an even number of incident occupied bonds; zero is allowed). So a generalized RC model is an Eulerian-subgraph model if W(H) = 0 whenever H is not Eulerian. The simplest such model (the "standard Eulerian-subgraph model") has

$$W(H) = \begin{cases} n & \text{if } H \text{ is Eulerian,} \\ 0 & \text{otherwise.} \end{cases}$$
(2)

Another such model (the "disjoint-loop model") has

$$W(H) = \begin{cases} n & \text{if } H \text{ is a cycle or an isolated vertex,} \\ 0 & \text{otherwise.} \end{cases}$$
(3)

These two models are identical if the underlying graph G has maximum degree ≤ 3 (e.g., the honeycomb lattice) but are different otherwise. More generally, we can consider the "degree-weighted model"

$$W(H) = n \prod_{i \in V(H)} t_{d_H(i)},\tag{4}$$

where $d_H(i)$ is the degree of the vertex *i* in the graph *H*, and t_0, t_1, t_2, \ldots are nonnegative weights satisfying $t_d = 0$ for all odd degrees *d*.

It is well known [6] that on any graph G of maximum degree ≤ 3 , the model (2)/(3) arises for positive integer n as a loop representation of an *n*-component spin model,

$$\tilde{Z} = \operatorname{Tr} \prod_{ij \in E} (1 + n x_{ij} \boldsymbol{\sigma}_i \cdot \boldsymbol{\sigma}_j),$$
(5)

where $\boldsymbol{\sigma} = (\sigma^1, \dots, \sigma^n) \in \mathbb{R}^n$ and Tr denotes normalized integration with respect to any *a priori* measure $\langle \cdot \rangle_0$ on \mathbb{R}^n satisfying $\langle \sigma^{\alpha} \sigma^{\beta} \rangle_0 = n^{-1} \delta^{\alpha\beta}$ and $\langle \sigma^{\alpha} \rangle_0 = \langle \sigma^{\alpha} \sigma^{\beta} \sigma^{\gamma} \rangle_0 = 0$. (In particular, uniform measure on the unit sphere is allowed, as are various "face-cubic" and "corner-cubic" measures [6].) For $n \neq 1$, the Boltzmann weight (5) with spins on a sphere defines a nonstandard O(n) spin model, which has positive weights only for $|x_{ij}| < 1/n$, but it is, nevertheless, expected to belong to the usual O(n) universality class.

Likewise, on any graph *G*, the model (2) [but not (3)] arises from the spin model (5) with a "face-cubic" *a priori* measure; i.e., $\boldsymbol{\sigma}$ is a unit vector $\pm \mathbf{e}_{\alpha}$ $(1 \le \alpha \le n)$ with probability 1/(2n).

Finally, let us observe that, on any *planar* graph *G*, there is a one-to-two correspondence between Eulerian bond configurations *A* on *G* and Ising configurations $\{\sigma\}$ on the dual graph *G*^{*} (namely, *A* is the Peierls contour for $\{\sigma\}$). Under this mapping, the bond model (2) on *G* with n = 1 is isomorphic to the Ising model on *G*^{*} with couplings J_e satisfying $v_e = e^{-2J_e}$. More generally, the model (4) with n = 1 and $t_d = ab^d$, i.e.

$$W(H) = \begin{cases} a^{|V(H)|} b^{2|E(H)|} & \text{if } H \text{ is Eulerian} \\ 0 & \text{otherwise} \end{cases}$$
(6)

gives an Ising model with $b^2 v_e = e^{-2J_e}$.

With this observation, we can present two algorithms of Chayes-Machta type for simulating Eulerian-subgraph models on a planar graph G. In these algorithms, we keep at all stages an Eulerian bond configuration A on G together with one of the corresponding Ising spin configurations $\{\sigma\}$ on G^* .

The first algorithm applies to those models in which there exist a, b > 0 such that $W(H) \ge a^{|V(H)|} b^{2|E(H)|}$ for all Eulerian *H*. This includes, in particular, the model (4) with $n \ge 1$ and $t_d \ge ab^d$.

Algorithm 1.—We use two colors: an active color ($\alpha = 1$) with $W_1(H)$ given by (6), and an inactive color ($\alpha = 2$) carrying the remaining weight. The first step of the algorithm leads to a partitioning $V = V_1 \cup V_2$. We now freeze all bonds having one or both vertices in V_2 in their current state, while leaving bonds within V_1 free to move. The latter bonds are updated by applying one step of the Swendsen-Wang algorithm to the correspondingly constrained Ising model on G^* . In detail, the algorithm proceeds as follows (see [25] for a proof of validity):

(1) Independently for each component H_i , color it $\alpha = 1$ with probability $W_1(H_i)/W(H_i)$, and $\alpha = 2$ otherwise.

(2) On each edge of G^* whose dual does not lie entirely within V_1 , place a bond. On each edge ij of G^* whose dual e lies entirely within V_1 and for which $J_e \sigma_i \sigma_j > 0$, place a bond with probability $1 - e^{-2|J_e|}$. (Here J_e is defined by $b^2 v_e = e^{-2J_e}$.)

(3) Form clusters on G^* from sites connected by bonds. Independently on each cluster, flip the Ising spins with probability $\frac{1}{2}$. (Note that a cluster typically contains spins of both signs.) (4) The new bond configuration is the Peierls contour for the new Ising configuration.

This algorithm is easily extended to allowing more than one active color, in case $W(H) \ge ka^{|V(H)|}b^{2|E(H)|}$ for some integer $k \ge 2$. We freeze all bonds having one or both vertices in an inactive color as well as all bonds connecting two distinct active colors.

Please note that Algorithm 1 is actually a *family* of algorithms corresponding to different choices of a, b. Among the maximal allowed pairs (a, b), some choices may be more efficient than others.

Alternatively, one can use other choices of $W_1(H)$ and then update the resulting bond model using either a local algorithm [19] or a worm-type algorithm [26].

Our second algorithm applies to the standard Euleriansubgraph model (2) with $n \ge 1$. Let us first observe, using the relation cycles = bonds - vertices + components, that we can replace n by $n^{c(H)} [c(H) =$ cyclomatic number of H] in (2) if we simultaneously replace v_{ij} by $x_{ij} = v_{ij}/n$. Note also that the total number of faces (including the exterior face) in a bond configuration A equals its cyclomatic number plus 1. Therefore, we can attribute a weight n to each face.

Algorithm 2.—We again use two colors, but this time we color the *faces* of A (which are clusters of vertices of the dual graph G^*). In detail, the algorithm proceeds as follows (see again [25] for a proof of validity):

(1) Independently for each face of *A*, color it $\alpha = 1$ with probability 1/n and $\alpha = 2$ with probability 1 - 1/n. This decomposes the vertex set of G^* as $V_1 \cup V_2$.

(2) On each edge of G^* that does not lie entirely within V_1 , place a bond. On each edge ij of G^* that lies entirely within V_1 and for which $J_e \sigma_i \sigma_j > 0$, place a bond with probability $1 - e^{-2|J_e|}$. (Here *e* is the bond of *G* dual to ij, and J_e is defined by $x_e = e^{-2J_e}$.)

(3),(4) As in Algorithm 1.

This algorithm is again easily extended to allowing more than one active color, in case $n \ge 2$, using the same rule as in Algorithm 1.

Numerical results.—We began by investigating the loop model (2)/(3) on the honeycomb lattice, for which Nienhuis [9] and Baxter [27] found the exact critical point when $-2 \le n \le 2$: $x_c(n) = (2 + \sqrt{2 - n})^{-1/2}$. Nienhuis further observed [9,28] that the critical O(n) model with $n \ge 0$ corresponds to a *tricritical* Potts model with $q = n^2$. From Coulomb-gas theory, the leading and subleading thermal exponents and leading magnetic exponent for the Potts model are known to be [9]

$$y_{t0} = \frac{3g-6}{g}, \quad y_{t1} = \frac{4g-16}{g}, \quad y_{h0} = \frac{(g+2)(g+6)}{8g},$$
(7)

where g is the Coulomb-gas coupling and $q = 4\cos^2(\pi g/4)$; here $g \in [2, 4]$ for critical and [4,6] for tricritical. We also have a hull exponent [29] $y_H = (g+2)/g$.

TABLE I. A comparison of the critical exponents determined by Monte Carlo simulations and those predicted by (7) *et seq.*

Ν		y_{t0}	y_{t1}	y_{h0}	y_H
1.25	Exact	1.83277	0.88740	1.93436	1.38908
	Num.	1.8327(1)	0.887(2)	1.9343(2)	1.3890(1)
1.50	Exact	1.78054	0.748 11	1.91989	1.406 49
	Num.	1.7805(1)	0.749(5)	1.9198(1)	1.406 5(1)
1.75	Exact	1.707 86	0.55428	1.903 47	1.43071
	Num.	1.7080(1)	0.55(1)	1.903 5(1)	1.4307(1)
2.00	Exact	1.5	0	1.875	1.5
	Num.	1.5000(1)	-0.01(2)	1.8750(1)	1.500 1(1)

We simulated the honeycomb-lattice loop model (2)/(3) at $x = x_c(n)$ for n = 1.25, 1.5, 1.75, and 2, using Algorithm 1, for 14 linear lattice sizes *L* in the range $4 \le L \le 1024$. Periodic boundary conditions were applied [30]. We measured the observables $\mathcal{N} =$ number of occupied bonds, $S'_2 =$ sum of squares of loop lengths, $\mathcal{D}_2 =$ sum of squares of face sizes, and $\mathcal{M} =$ total Ising magnetization. From these we obtained the specific-heat-like quantity $C = L^{-d} \langle \mathcal{M}^2 \rangle - \langle \mathcal{M} \rangle^2$ and the Ising susceptibility $\chi_{\text{Is}} = L^{-d} \langle \mathcal{M}^2 \rangle$ (here d = 2).

Our numerical results for the static critical exponents are shown in Table I. We find empirically that $\chi_{Is} \propto L^{2y_{t0}-d}$, $L^{-d}\langle \mathcal{N} \rangle \propto \text{const} + L^{y_{t1}-d}$, $C \propto \text{const} + L^{2y_{t1}-d}$, $L^{-d}\langle \mathcal{D}_2 \rangle \propto L^{2y_{h0}-d}$, and $L^{-d}\langle S'_2 \rangle \propto L^{2y_H-d}$. (We shall give elsewhere [20,25] theoretical arguments for several of these.) The leading Potts exponents y_{t0} and y_{h0} are absent in the spin and loop observables of the O(n) model, but they can be seen in observables associated to the faces. To our knowledge, this is the first observation of y_{t0} , y_{h0} , and y_H in O(n) models (see also [20]).

We have also determined the integrated autocorrelation time τ_{int} for the observables $\mathcal{N}, \mathcal{S}'_2, \mathcal{D}_2$, and \mathcal{M}^2 . For $1 < n \leq 2$, critical slowing-down is either entirely absent or

3.4 3 2.6 2.2 1.8 1.4 10 L 100 1000

FIG. 1 (color online). Integrated autocorrelation times τ_{int} for the O(2) loop model on the honeycomb lattice versus lattice size L.

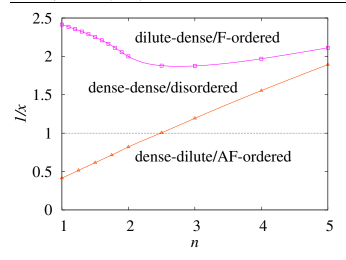


FIG. 2 (color online). Location of the ferro- and antiferromagnetic phase transitions of the standard Eulerian-subgraph model (2) on the square lattice, as a function of n.

very nearly absent. The τ_{int} data for n = 2 are shown in Fig. 1, and they strongly suggest that the dynamic critical exponent z is zero for n = 2.

We next simulated model (2) on the square lattice, using Algorithm 2, for lattices of linear size $4 \le L \le 512$ with periodic boundary conditions [30]. The phase diagram of this model is largely unexplored, especially when n > 2. Phase transitions were located by analyzing the data for several Binder-type ratios. Two distinct phase transitions were found, which correspond to the ferromagnetic and antiferromagnetic transitions in terms of the dual Ising spins (see Fig. 2). The three phases are, respectively, dilute-dense, dense-dense, and dense-dilute, where A-Bmeans that the bond configuration is of type A and its complement is of type B (here "dense" = "percolating"). The locations of the ferromagnetic critical point for $n \le 2$ agree accurately with those given in [31]. Our data show that the lines of phase transitions extend to n > 2. We shall discuss elsewhere [25] the nature of this new phase transition. The critical slowing-down is essentially absent for $1 \le n \le 2$, but is strong for n > 2.

We also applied Algorithm 1 to model (4) on the square lattice. The results, along with a conjectured renormalization-group flow in the (n, v, t_4) space, will appear elsewhere [25].

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