Invaded cluster algorithm for Potts models

J. Machta, Y. S. Choi, A. Lucke and T. Schweizer

Department of Physics and Astronomy, University of Massachusetts, Amherst, MA 01003-3720

L. M. Chayes

Department of Mathematics, University of California, Los Angeles, CA 90095-1555

Abstract

The invaded cluster algorithm, a new method for simulating phase transitions, is described in detail. Theoretical, albeit nonrigorous, justification of the method is presented and the algorithm is applied to Potts models in two and three dimensions. The algorithm is shown to be useful for both first-order and continuous transitions and evidently provides an efficient way to distinguish between these possibilities. The dynamic properties of the invaded cluster algorithm are studied. Numerical evidence suggests that the algorithm has no critical slowing for Ising models.
I. INTRODUCTION

This paper discusses a new cluster method for simulating both continuous and first-order transitions. The method, called the invaded cluster (IC) algorithm, was introduced in Ref. [1] in the context of the Ising model. In this paper we give a more extended discussion of the IC method, present new data for Ising and Potts critical points and show how to apply the method to first-order transitions in Potts models.

The Swendsen-Wang (SW) algorithms [2] and other cluster methods [3,4] have led to vast improvements in the efficiency of simulating the critical region of a variety of spin models. For the Potts models, these algorithms are based on the Fortuin-Kastelyn (FK) [5] representation of the system as a correlated bond percolation problem. Each Monte Carlo step in a cluster algorithm consists of generating an FK bond configuration from the spin configuration by occupying some of the satisfied bonds (i.e. bonds across which the spins are in agreement) of the lattice. Clusters of connected sites are randomly and independently assigned a new spin value that is the same throughout the cluster. This creates the updated spin configuration.

The IC algorithm shares these basic features with other cluster algorithms but differs in how the bond configurations are generated. For other cluster algorithms, satisfied bonds are independently occupied with a probability that depends on the temperature. For the invaded cluster algorithm, bonds are occupied in a random order until the bond configuration fulfills a stopping condition. For example, the stopping condition may be a requirement on the size of the largest cluster. For judicious choices of stopping condition the IC algorithm simulates the transition point of the model.

The relation between the SW algorithm and the IC algorithm is analogous to the relation between ordinary percolation and invasion percolation. In ordinary (bond) percolation [6], bonds are independently occupied with probability \( p \) forming connected clusters of sites. At the percolation threshold, \( p_c \), in a large finite box, the probability that any one of these cluster has an extent comparable to the system size “jumps” from nearly zero to nearly one.

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In invasion percolation \([7–12]\), at least the version most relevant to the current work, \(b\) bonds are randomly ordered and then successively occupied until a stopping condition is reached. If, in a system of scale \(L\), the stopping rule is that some cluster is comparable in extent to \(L\) then, as is easily shown (cf. footnote (11) in [1]) the fraction of occupied bonds will approach \(p_c\) as \(L \to \infty\). The IC algorithm for the \(q\)-state Potts models can be loosely described as the generalization of invasion percolation to the FK random cluster models.

The IC algorithm has several very attractive features: First, the algorithm may be used to study a phase transition without \(a\ priori\) knowledge of the transition temperature. The IC algorithm thus enjoys the property of “uniformity” or “self-organized criticality.”

The transition temperature is an \textit{output} of the IC algorithm just as \(p_c\) is an output of invasion percolation. In cases where the critical temperature is unknown or not known with sufficient accuracy, this can be a significant advantage. Histogram reweighting \([13,14]\) is now the method of choice for high precision measurements of the critical temperature. This method involves extrapolating from a guessed critical temperature and its systematic errors are difficult to judge.

The IC algorithm is also an extremely fast way to simulate Ising-Potts critical points. In Sec. [IVD] we show that autocorrelation times for the IC algorithm are significantly smaller than for related cluster algorithms. Indeed, for certain quantities such as the energy and

\(^1\)Conventionally, invasion percolation is formulated as a growth process that is initiated at a limited number of seed sites, e.g. a single site.

\(^2\)In the computer science literature, an algorithm is called “uniform” if it can be applied to problems of arbitrary size without the need for significant precomputation. Conventional Monte Carlo sampling of critical points is non-uniform because precomputation is required to obtain the critical temperature. As the system size increases, the critical temperature must be computed to increasing accuracy. For the IC algorithm the critical temperature is not used as an input. In this setting the concept of “uniformity” is akin to the idea of “self-organized criticality.”
the finite-size critical temperature, the integrated autocorrelation time appears to approach zero as the system size increases as a result of anti-correlations between successive Monte Carlo steps.

Although first devised for critical points, the IC algorithm is also effective for studying first-order transitions. For first-order transitions, it is convenient to have the stopping rule control the number of sites in the largest cluster. In this way the average magnetization is essentially fixed and a point in the coexistence region can be explored. By sampling a single point in the coexistence region the problem of exponentially long tunneling times between phases is avoided. The multicanonical Monte Carlo method \cite{13} also avoids exponential tunneling times and has some features in common with the IC algorithm. We also present a criterion for distinguishing first-order from continuous transitions. This criterion is effective for small system sizes and easily reveals the first-order nature of the 5-state two-dimensional Potts model and the 3-state three-dimensional Potts model, both of which have very weak first-order transitions.

In finite volume, the IC algorithm does not sample the canonical ensemble. We call the invariant measure sampled by the algorithm the “IC ensemble.” A fundamental supposition of this paper is that the IC ensemble is equivalent to the usual ensembles of statistical mechanics in the thermodynamic limit. Just as the microcanonical and canonical ensembles agree for all local observables in the infinite volume limit, we conjecture that the IC ensemble agrees with these two ensembles for all local observables. On the other hand, global fluctuations differ in different ensembles. For example, energy fluctuations are proportional to the heat capacity in the canonical ensemble but this is not the case for the IC or microcanonical ensembles. Although we have no proof yet of our assertions concerning the validity of the algorithm, in Sec. \textit{III} we carefully state our claims and supply (non-rigorous) arguments to back them up.

Finally, we remark that in this study we have restricted our attention to Potts models. However, as shown by Wolff \cite{3}, it is possible to generalize the cluster methods to a much wider class of models using an embedding procedure. Presumably, the same ideas should
work for the IC algorithm but these matters will not be pursued here. Some additional spin
models and graphical representations appropriate for the IC algorithm are discussed in [16].

The rest of this paper is organized as follows, in Sec. [II] the invaded cluster algorithm
is described in more detail and in Sec. [III] the algorithm is justified and compared to other
simulation methods. In Sec. [IV] numerical results are presented.

II. INV ADED CLUSTER METHOD FOR POTTS MODELS

A. Potts models

The $q$-state Potts models are defined by a collection of spins, $\{\sigma_i\}$ with $i$ belonging to
some lattice and $\sigma_i = 1, 2 \ldots q$. The Hamiltonian is given by

$$H = -\sum_{\langle i, j \rangle} (\delta_{\sigma_i, \sigma_j} - 1)$$

where the summation is over the bonds of the lattice. If $q = 2$, this corresponds to an Ising
system. Here we consider hypercubic lattices of size $N = L^d$, usually with periodic boundary
conditions and $\langle i, j \rangle$ denotes a nearest neighbor pair.

In the seminal work of Fortuin and Kastelyn [5] it was shown that the above defined spin
system gives rise to a set of percolation-type problems known as random cluster models.
These models are defined by weights $W$ on bond configurations $\omega$ (collections of occupied
bonds) that are given by

$$W(\omega) = p^{\mid\omega\mid}(1-p)^{|E|-|\omega|}q^{C(\omega)}$$

where $|\omega|$ is the number of occupied bonds, $|E|$ is the number of bonds in the lattice (here
d$N$), $C(\omega)$ is the number of connected components of $\omega$ (counting isolated sites) and

$$p = p(\beta) = 1 - e^{-\beta}$$

is the relationship between the bond density parameter and the temperature in the spin
system. In particular,
\[
\sum_{\omega} W(\omega) = Z_\beta \equiv \text{Tr}[e^{-\beta H}]
\]

is the graphical expression for the partition function in finite volume. The weights make sense for any positive real \(q\) and the case \(q = 1\) is manifestly the usual bond percolation problem. In this paper we only consider positive integer values of \(q\). It has been gradually realized, with increasing degrees of sophistication, \([5,17–19,2,20]\) that the graphical model and the spin model are “equivalent” e.g. expectations of observables in the spin model are easily calculable in terms of appropriate probabilities in the random cluster system. From the modern perspective, the two descriptions are regarded as different facets of a single larger problem that is none other than the annealed bond–diluted version of the original Potts model.

In \(d \geq 2\), for all \(q\), there is a phase transition and for \(q = 1\) and 2, this transition is continuous. (See \([21, 22]\) for details.) For \(q \gg 1\), it can be proved that the transition is first-order \([23]\). In two dimensions, it is widely accepted that the transition is continuous for \(q = 3\) and 4 and discontinuous for \(q \geq 5\). In \(d \geq 3\), the transition is believed to be discontinuous for all \(q \geq 3\).

**B. The algorithm**

The invaded cluster algorithm for the \(q\)-state Potts models is defined as follows: Consider a finite lattice on which there is some spin configuration \(\{\sigma_i\}\). From the spin configuration, a bond configuration is constructed via a modified form of invasion percolation. First, the bonds of the lattice are assigned a random order. Bonds, \(\langle i, j \rangle\), are tested in this order to see if \(\sigma_i = \sigma_j\). If the latter occurs, the bond is said to be satisfied and it is added to the bond configuration. These bonds are called the occupied bonds. The unsatisfied bonds are not considered for the remainder of the current Monte Carlo step.

The set of occupied bonds partitions the lattice into clusters of connected sites. The cluster structure evolves until a stopping condition is achieved. The stopping condition is typically based on the size or topology of a single cluster as detailed below. When the
stopping condition has been satisfied, a new spin configuration is obtained by randomly assigning one of the \( q \) spin types to each cluster (including isolated sites) and setting each spin in the cluster to that value. Statistics are collected on the new spin/bond configuration thereby completing a single Monte Carlo step.

Among the quantities typically measured on each Monte Carlo step is the ratio \( f \) of occupied to satisfied bonds. As we shall see, \( f \) serves as an estimator of the temperature of the system through the relation, \( \langle f \rangle \approx 1 - e^{-\beta} \).

**C. Stopping rules**

The IC method does not use the temperature as an input parameter. Instead, an appropriate choice of stopping rules allows us to simulate the critical point, the coexistence region of a discontinuous transition or a point away from the phase transition. The numerical results in Sec. [V] feature the following three stopping rules:

- **Extension rule**
  
  Some cluster has maximum extent \( L \) (the system size) in at least one of the \( d \) directions.

- **Topological rule**
  
  Some cluster winds around the system in at least one of the \( d \) directions. (This is the usual rule for the identification of spanning clusters in percolation.)

- **Mass rule**
  
  Some cluster has at least \( mN \) sites.

It is noted that for any of the above rules, the invasion stops when exactly one cluster satisfies the condition. Thus, during the evolution of the bond configuration, when a given bond is occupied, it is only necessary to check its cluster to see if the stopping condition is fulfilled. The extension and topological rules involve no parameters and are used to drive the system to a phase transition point. Both of these rules require that some cluster is barely
the size of the system. We refer to these and related stopping rules as *spanning* rules and to the unique cluster which satisfies the rule as the *spanning cluster*.

The mass rule involves an input parameter, $m$ and permits us to simulate an arbitrary temperature and also to explore the coexistence region of discontinuous transitions. The mass rule is an example of a *fixed parameter* rule, some alternative fixed parameter rules that we have not yet studied numerically are:

- **Magnetization rule**

  This is closely related to the mass rule and is defined in systems where the spins at the boundary of the sample are all set to 1. Thus bonds connecting the boundary sites to internal sites may only be occupied if the latter also have spin value 1. The stopping condition is fulfilled when the number of sites connected to the boundary first exceeds $mN$.

- **Susceptibility rule**

  After the $k^{th}$ bond has been occupied, let us suppose that there are $r = r(k)$ clusters, $C_1, \ldots C_r$ containing $|C_1|, \ldots, |C_r|$ sites. The stopping condition is fulfilled when $|C_1|^2 + \ldots + |C_r|^2$ first exceeds $\chi N$.

- **Energy rule**

  Here one counts the number of bonds whose endpoints are in the same connected cluster (regardless of whether the bond itself has actually been occupied) and stops when the tally exceeds $\varepsilon |E|$.

- **Density rule**

  Essentially equivalent to the energy rule, we simply count the number of bonds that have been selected and stop when the total is $\rho |E|$.

The magnetization, susceptibility and energy rules are derived from their thermodynamic namesakes via the FK representation. In a system with symmetry breaking boundary condi-
tions as described above, the average magnetization is precisely the number of sites connected to the boundary. Similarly, the susceptibility is the average size of the bond cluster containing the origin. In periodic boundary conditions this is the same as the average of the sum of the squares of the cluster sizes divided by $N$. Finally, (cf. below) the energy per bond is related to the probability that both ends of a bond are in the same cluster.

The mass rule and density rule both require some additional explanation. Let us start with the mass rule: in free or periodic boundary conditions, the finite-volume magnetization will always vanish by symmetry. Presumably, this is because all of the $q$ distinct low temperature states are equally represented. Below the bulk transition temperature, this is manifested in the FK picture by the appearance of one large cluster and a multitude of small clusters. The $q$ different values that could be assigned to the single large cluster yields the above mentioned convex combination. However, because of the a priori symmetry between these states, if we agree in advance to always assign a fixed value to the large cluster, we get (a finite volume approximation to) the corresponding pure state. Such a picture can be easily established with complete rigor if the temperature is low. For $q = 2$ in $d = 2$ this picture holds up to the critical point as can be derived from the main theorem in [24,25] and for $q \gg 1$ it is a consequence of the result in [26]. Thus, we will suppose that fixing the fraction of sites in the largest cluster is equivalent to fixing the magnetization.

Turning attention to the fixed bond density rule, let us assume that we are in a finite box $\Lambda$ e.g. with periodic boundary conditions. The total energy divided by the number of bonds is given by

$$\varepsilon = \frac{1}{|E|} \sum_{<i,j>} <1 - \delta_{\sigma_i,\sigma_j}>_{\beta,\Lambda}$$  \hspace{1cm} (5)$$

where $<>_{\beta,\Lambda}$ denotes the thermal average at temperature $\beta^{-1}$ in the box $\Lambda$. However, the quantity $<1 - \delta_{\sigma_i,\sigma_j}>$ is easily evaluated in the FK representation as

$$<1 - \delta_{\sigma_i,\sigma_j}>_{\beta,\Lambda} = (1 - \frac{1}{q})(1 - a)$$  \hspace{1cm} (6)$$

where $a(q, p, \Lambda)$ is the probability that two neighboring sites belong to the same cluster. On the other hand, differentiating the expression Eq. (2) with respect to $\beta$ we get
\[
\varepsilon = -\frac{1}{|E|} \frac{\partial \log Z}{\partial \beta} = -\frac{1}{|E|} (1-p) \frac{1}{Z} \frac{\partial Z}{\partial p} = \left[ 1 - \frac{b}{p} \right]
\]

where \( b = b(p, q, \Lambda) \) is the probability (in a translation and reflection invariant system) that a given bond is present. Evidently, there is a simple relationship between the quantities \( a \) and \( b \). We thus argue that fixing the density of bonds has an equivalent effect as fixing the density of neighboring pairs in the same cluster. The mass and density rules have clear computational advantages over their counterparts, the magnetization and energy rules respectively. For this reason (and certain others) these rules will supersede the energy and magnetization rules in the remainder of this paper.

Later we will argue that the fixed parameter rules produce, in the thermodynamic limit, the equilibrium system at the temperature corresponding to the chosen value of the parameter. This temperature is therefore an output of the simulation. For first-order transitions, the goal is to find parameter values in the “forbidden region”; which is in principal easy for the magnetization and the susceptibility. For continuous transitions, we may probe the critical region by considering a sequence of magnetizations tending to zero or susceptibilities tending to infinity. For example, choosing \( \chi \sim N^s \) with \( s < 1 \) the transition temperature and the critical behavior of local observables will again, presumably, emerge from the output.

The fixed parameter stopping rules are related to “constrained” random cluster ensembles. Consider, for example, the constrained mass ensemble defined by restricting attention exclusively to bond configurations \( \omega \) that satisfy the mass condition (that the largest cluster in \( \omega \) has mass \( M = mN \)) and are otherwise weighted by \( q \) raised to the number of components of \( \omega \):

\[
W_M(\omega) = \mathbb{I}_M(\omega) q^{C(\omega)}
\]

3If \( b_{i,j} \) is the probability that the bond \( < i, j > \) is occupied and \( a_{i,j} \) is the probability that \( i \) and \( j \) belong to the same cluster, the relation \( (q - 1)(1 - a_{i,j}) = q(1 - b_{i,j}/p) \) holds even without the assumption of translation invariance.
where $\mathbb{1}_M(\omega)$ is one if $\omega$ satisfies the stated mass condition and is zero otherwise. Similarly, we may construct the \textit{constrained density ensemble} and the \textit{constrained susceptibility ensemble} using weights as in Eq. 8 with $\mathbb{1}_M$ replaced by the appropriate restricting functions.

Although we have not yet attempted a derivation, it would seem that standard “equivalence of ensembles” arguments could be developed to show that in the thermodynamic limit, these measures are identical to the usual random cluster measures at an appropriate value of $p$. In this case, according to the ideas in [19], the associated measure in the spin system converges to the corresponding Gibbs distribution.

Somewhat to our surprise, it is readily shown that the IC density rule algorithm simulates the constrained density ensemble. This constitutes the current high water mark as far as rigorous results are concerned. We present the argument below:

\textbf{Theorem} In finite volume, the IC algorithm with the density rule with $\rho|E| = P$ samples the joint bond-spin distribution defined by the weights

\[ W(\sigma, \omega) = \mathbb{1}_P(\omega)\Delta(\sigma, \omega). \quad (9) \]

In particular, the random cluster marginal is the constrained density ensemble defined as in Eq. 8.

\textit{Proof} For a fixed spin configuration with at least $P$ satisfied bonds, consider the set $\Omega_P(\sigma)$ of all bond configurations $\omega$ consistent with the spin configuration $\sigma$ and with $|\omega| = P$:

\[ \Omega_P(\sigma) = \{ \omega \mid \mathbb{1}_P(\omega) = 1 \text{ and } \Delta(\sigma, \omega) = 1 \}. \quad (10) \]

We define an “algorithm” as follows: starting from an $(\omega, \sigma)$ with $\mathbb{1}_P(\omega)\Delta(\sigma, \omega) = 1$, the spin moves are identical to the usual SW or IC spin moves while the bond moves are defined by selecting, without bias, any $\omega$ out of $\Omega_P(\sigma)$. It is manifestly apparent that this “algorithm” samples the above joint bond-spin distribution.

We claim that for the density rule, the bond moves are equivalent to this unbiased selection from $\Omega_P(\sigma)$. Indeed let $\sigma$ denote a spin configuration. For the purpose of this
theorem, let us implement the IC algorithm by assigning random numbers to the bonds and selecting the lowest $P$ satisfied bonds. We may write $\Omega_P(\sigma) = \{\omega_1, \ldots, \omega_R\}$ with each $\omega_k$ a subset of the satisfied bonds of the $\sigma$ and having $|\omega_k| = P$. It is clear that under IC dynamics, the criterion for selecting $\omega_k$ is that the highest bond in $\omega_k$ is lower than the highest bond in any other $\omega \in \Omega_P(\sigma)$. Configurations in $\Omega_P(\sigma)$ all have the same number of bonds, hence the probabilities are unbiased $\square$.

III. VALIDITY OF THE IC METHOD

A. Comparison to the Swendsen-Wang algorithm

The IC algorithm is in fact very similar to the SW algorithm and this, to the greatest extent, is the basis of our intuition concerning the former. In both cases, occupied bond clusters are grown on top of a spin configuration by randomly selecting some of the satisfied bonds. When the growth process is stopped, both algorithms generate the updated spin configuration from the bond configuration by the same procedure (described in the second paragraph of the introduction). Thus the “only” difference lies in how the bonds are selected.

In the SW algorithm, satisfied bonds are independently accepted with probability $p$. If $p$ is identified with a temperature as in the FK representation (Eq. 3) one can show [2,20] that detailed balance is satisfied for both the spins and the bonds: For the spins, this is with respect to the canonical Gibbs measure and for the bonds, with respect to the random cluster measure. The key observation [20] is that the SW algorithm simulates a joint measure on spin and bond configurations that is defined by the weights

$$W(\sigma, \omega) = p^{\omega} (1 - p)^{|E | - |\omega|} \Delta(\sigma, \omega).$$

(11)

In the above, $\Delta(\sigma, \omega)$ insures consistency between the bond and spin configurations: $\Delta(\sigma, \omega)$ is one if all occupied bonds in the configuration are satisfied, otherwise $\Delta(\sigma, \omega)$ is zero.

It is worth pointing out that the SW algorithm can in fact be described in the framework of an IC algorithm. Let $S = S(\sigma)$ denote the number of satisfied bonds in a given spin
configuration. Let $A_p(\sigma)$ denote the random variable that is chosen according to binomial statistics:

$$P(A_p(\sigma) = A) = \binom{S}{A} p^A (1 - p)^{S-A}$$

(12)

The SW algorithm is defined by the stopping rule that on each round, an $A_p(\sigma)$ is drawn, and then the growth stops as soon as the first $A_p(\sigma)$ satisfied bonds are accepted. Viewed in this light, the SW algorithm is seen to be quite similar to some of the IC algorithms under consideration. However, unlike the SW algorithm, we are unable to write down a joint distribution similar to Eq. (11) for any of the IC ensembles except for the case of the density rule.

**B. Away from criticality: fixed parameter algorithms**

As the title of this subsection indicates, we will provide separate discussions of the critical and non-critical algorithms. In part, this is because the latter require fewer assumptions about the equilibrium state of the Potts/FK system. For the fixed parameter stopping rules, the fundamental assumption is that the ensemble sampled by the IC algorithm has the same local properties as the canonical ensemble at a temperature that yields the chosen value for the thermodynamic function (e.g. magnetization or susceptibility) that defines the stopping rule. We present the argument for the susceptibility rule though similar arguments are possible for other rules.

Consider a large Potts/FK system in equilibrium at $p < p(\beta_c)$ (i.e. $T > T_c$). The “system wide” average cluster sizes will be very close to the mean value which, in turn, is close to the limiting infinite volume susceptibility, $\chi(p)$. Explicitly, for bond configuration $\omega_L$ on a lattice of scale $L$, we may compute

$$C(\omega_L) = \frac{C_1^2 + \ldots + C_N^2}{N}$$

(13)

and, as $L \to \infty$, the distribution of $C(\omega_L)$ will be sharply peaked about $\chi(p)$.
Now let us contrast the behavior of the SW and IC algorithms on a given spin configuration. Suppose that both the SW and IC algorithms run by assigning a random number uniformly in $[0, 1]$ to each bond of the lattice. For the IC algorithm, this provides us with the ordering while for the SW algorithm at parameter $p$, the instructions are to occupy all satisfied bonds with random number less than $p$. For the SW algorithm we may occupy bonds as a function of continuous time $t$, $0 \leq t \leq 1$. At time $t$, all satisfied bonds with value less than $t$ are occupied. Clearly, if we were to stop too soon, e.g. at $t = p - \epsilon$, the value of $C$ will be, with high probability, strictly smaller than the equilibrium value. Similarly if we were to go beyond $p$ to $t = p + \epsilon$, we will get a value of $C$ that is too large. On the other hand, stopping, at $t = p$ as we are supposed to, yields a value of $C$ that, by definition, is typical of the equilibrium distribution.

Now, starting from the same spin configuration, let us do a step of the IC algorithm (with the susceptibility rule) stopping when $C = \chi = \chi(p)$. We may also envision this operation taking place as a function of continuous time. Now at time $t$, the same bonds have been collected in both Monte Carlo schemes and we can reiterate the previous discussion to conclude that the algorithm will not stop significantly before or after $t = p$. If the system is in equilibrium initially then the IC algorithm chooses nearly the same stopping time as the SW algorithm. Since the latter was in equilibrium, evidently under IC, we stay in equilibrium.

The opposite perspective provides us with an equally valid argument: Suppose it is the case (as is observed) that the fraction $f$ of satisfied bonds that are occupied has a distribution that is sharply peaked. Then, each step of the IC algorithm amounts to an iteration of the SW algorithm with parameter value equal to $f$. If the stopping rule demands that $C = \chi(p)$, it follows that $f = p$.

Of course for an actual simulation in finite volume, the above arguments are by no means a rigorous proof: To achieve $C = \chi(p)$, the algorithm stops at $t = p + \eta_L$ where $\eta_L$ is a random variable depending on the spin configuration. Even if we know that $|\eta_L| \to 0$ as $L \to \infty$, we cannot, as of yet, control the effects that these fluctuations have on the limiting
distribution for the IC algorithm. Indeed, the IC distribution will differ from the canonical distribution in finite volume. However, it is hard to believe that these objects do not tend to the same distribution in the infinite volume limit, all we lack is a proof. Nevertheless, to summarize our argument, the assumption that the bond/spin configurations typical of the Gibbs distribution are close to the ones of the IC ensemble is self-consistent in and of the fact that iterations of the IC algorithm keep us in the vicinity of the Gibbs distribution.

Let us now turn our attention to a discussion of the situation when the spin configuration is not typical of the desired equilibrium distribution; here the arguments will be somewhat less complete. We will again consider a given spin configuration and compare what happens under SW vs. IC dynamics. Suppose, for example, that the spin configuration is at too high a temperature. We may imagine that we have a configuration that is typical of the Gibbs distribution at an inverse temperature $\tilde{\beta}$ such that $1 - e^{-\tilde{\beta}} \equiv \tilde{p} < p \equiv 1 - e^{-\beta}$. If we do a single iteration of the SW algorithm, again collecting our bonds according to the time parameter $t$, as always, we stop when $t = p$. Because the temperature of the spin configuration was higher than $1/\beta$, the percentage of satisfied bonds will be relatively lower than typical for configurations that are at the right temperature. Evidently, when we stop, with large probability the average bond cluster size will be smaller than $\chi(p)$ but larger than $\chi(\tilde{p})$. The resulting spin configuration will therefore be of intermediate character between those that are typical of temperatures $1/\beta$ and temperatures $1/\tilde{\beta}$. By contrast, when $t = p$, the invaded cluster algorithm does not stop. Hence, the fraction $f$ of satisfied bonds that get occupied under IC dynamics is in excess of $p$ and the new configuration (also of intermediate character) is further towards the low temperature side. It is thus apparent that away from equilibrium, we are pushed in the right direction and, as this example illustrates, we are pushed harder in the direction of equilibrium under IC dynamics than under SW dynamics. Similar arguments apply to the case $p < \tilde{p}$ and show that now $f < p$. We believe, this “negative feedback mechanism” is ultimately responsible for the immense reduction (or complete absence) of critical slowing down that results from using IC dynamics.

Unfortunately, we now leave terra firma in order to discuss the case where the configu-
rations cannot be characterized in terms of a single temperature-like parameter. In general, dynamically generated configurations are out of equilibrium, however, we can consider the following proposal for the definition of an effective temperature: For a given configuration $\sigma_L$ let $C(\sigma_L, t)$ be the mean square cluster size (as defined in Eq. [3]) observed when we have occupied all satisfied bonds with value less than $t$. Let $\overline{C}(\sigma_L, t)$ be the average of this quantity over all realizations of random numbers on the bonds. A measure of the effective temperature of the configuration $\sigma_L$ is to compare $\overline{C}(\sigma_L, t)$ with $\chi(p)$, the actual (equilibrium) susceptibility as a function of the temperature parameter $p$. If there is a single (non–zero) point where these curves cross, this may be identified as an effective temperature. In any case, if $\overline{C}(\sigma_L, t = p)$ is smaller than $\chi(p)$ we can assert that the temperature is higher than that corresponding to the parameter $p$ while if $\overline{C}(\sigma_L, t = p) > \chi(p)$ it is lower. Obviously the colder configurations should be heated up and the hotter configurations should be cooled down. For spin configuration $\sigma$ and target susceptibility $\chi$, the typical simulation temperature of a move will be given by $t(\chi)$ such that $\overline{C}(\sigma_L, t(\chi)) = \chi(p)$. Since the curve $\overline{C}(\sigma_L, t)$ is monotone we again see the negative feedback mechanism–if the spin configuration is colder than that corresponding to $p$ the simulation temperature will be higher than that corresponding to $p$ and vice versa.

Of course these considerations apply as well to the other fixed parameter IC algorithms where we would argue–just as persuasively–that the magnetization vs. $t$ or energy vs. $t$ profiles of a spin configuration can be used to measure an effective temperature. It is worth remarking that there is one system where these suppositions are exact, namely the long-ranged mean-field Ising model. Here, for an $N$ site system, each site interacts with every other site via a coupling that scales inversely with $N$. At non-critical temperatures, the SW algorithm drives this system to equilibrium exponentially quickly (as expected) while, by contrast, the IC algorithm achieves equilibrium in at most two Monte Carlo steps [16].
C. At criticality: spanning algorithms

The general philosophy that underpins our belief in the validity of the critical algorithms is quite similar to the non–critical cases. The important differences lie in the implicit assumptions we have made concerning the behavior of the graphical representation at criticality—in particular in finite volume. Indeed, for a Potts system with a continuous transition, if we ask for the value of the parameter where the susceptibility is equal to a certain value, the answer is unambiguous. Furthermore, provided that \( L \) is large compared to the typical size of bond clusters, the statistics in a finite lattice of side \( L \) should represent an excellent approximation to the infinite volume behavior.

For the critical algorithms, the entire premise begins with non-trivial questions about the equilibrium critical behavior of the random cluster model in finite volume. For example, in a large system is there a single cluster with the following two properties?

1. The extent of the cluster is the scale of the system.

2. The cluster does not contain two (or more) disjoint subsets each of which satisfy condition (1).

Under the standard assumptions concerning the nature of the critical point the following picture, in the graphical representation, emerges: Above \( T_c \), the probability that there is any such cluster goes to zero exponentially in the scale of the system. Below \( T_c \), there is a single large cluster that exhausts a fraction—equal to the spontaneous magnetization—of the system. Within this cluster, there are many separate paths that are the scale of the system.

These include absence of an intermediate phase, uniqueness of the phase above and at \( T_c \), uniqueness of the \( q \) magnetized phases among the translationally invariant states below \( T_c \), continuously diverging correlation lengths using several definitions of the correlation length and positivity of the surface tension below \( T_c \). Nearly all of these properties have been established for independent percolation and a significant (but not sufficient) fraction in the Ising case.
To prevent this large cluster from happening, requires a fluctuation presumably as rare as \( \exp[-\text{const.}L^{d-1}] \). By process of elimination, the only place that a cluster with the above properties could exist on all scales is the critical point.

This is not to say that the above line of reasoning proves that these or other kinds of spanning clusters are indeed typically observed at the critical point (e.g. they could be power law rare) however, as we will see, the full validity of such an assertion is not essential for the broader features of our argument.

Indeed, the basic reasoning is now pretty much the same as in the non-critical case: If the distribution of \( f \) (the fraction of satisfied bonds that are occupied) is sharply peaked, the central value must correspond to that of the critical parameter, \( p(\beta_c) \). If this central value were too small then the correlation length would be a small fraction of the system and the clusters would not get big enough. If the peak value is so large that the (low temperature) correlation length is small compared to the system size, the biggest cluster would be too big. Thus, the value of \( f \) has to be close enough to the critical point to ensure that the correlation length is at least a scaling fraction of the system size.

If the required spanning cluster is itself, somehow, atypical of criticality, the above argument is still valid. As a concrete example, consider a stopping rule that terminates the cluster growth when there is a cluster of size \( \sqrt{L} \). Such a bad choice of a spanning cluster will nonetheless heavily favor the critical value of \( f \) over any non–critical value. The worst that could happen is that the scaling of the spanning cluster itself might be of the wrong type but in any case all local observables will still take on their critical values. Finally, starting at non–critical spin configurations, the negative feedback mechanism discussed in the previous subsection applies to these algorithms as well.

The weaker point in our argument concerns our reasoning as to why the distribution of the \( f \) values should be sharply peaked. (First and foremost, this has been observed in every critical system.) Let us imagine the problem in an infinite volume setting and consider a critical spin configuration. We again regard the process of growing the clusters as an independent percolation problem defined on the random graph that is provided by
the satisfied bonds of the spin configuration. Let us first assume that, in the usual sense, this problem has a sharp percolation threshold, $t_c$. It then follows easily that $t_c$ corresponds to the critical value of the FK parameter; $t_c = 1 - e^{\beta_c} \equiv p(\beta_c)$. Indeed, at $t = p(\beta_c)$, we have achieved the critical FK bond configuration and our assumption of a sharp threshold rules out the possibility of any other value. Going back to finite volume and starting from a critical configuration, the argument in this case is finished: If we stop at an $f$ significantly different from $p(\beta_c)$, we will get the wrong sort of clusters and stopping at $f \approx p(\beta_c)$ we keep the spin configuration critical.

Unfortunately, in $d = 2$, it is not the case that the underlying percolation problem has a sharp transition. Specifically, in the Ising model on the two-dimensional square lattice it was shown in [27,28] that percolation of one spin type is necessary and sufficient for the existence of a positively magnetized phase. (The analog of this result for the general $q$–state Potts models was proved in [29].) In particular, this means that in a critical configuration, there is no infinite cluster of satisfied bonds and thus, even if $t = 1$, there is no percolation in our secondary process. Evidently the percolation clusters on the critical spin configuration will themselves look critical for all $t$ between $p(T_c)$ and 1. It is easy to believe (but hard to prove so we will spare the reader the details of the heuristics) that the clusters will not go critical until $t = p(T_c)$. Thus, the algorithm will not stop collecting bonds until at least this point. However, in finite volume, there are, undoubtedly, typical critical spin configurations that forbid the existence of a spanning cluster. Of course this kind of disaster is ruled out by the mechanics of the algorithm: whatever the stopping condition, if it was satisfied on the

\textsuperscript{5} For example, if there are star-connected chains (meaning that neighbors and next nearest neighbors both count as connected) of plus spins and of minus spins winding both ways around the torus, the topological condition cannot be satisfied. At the critical point, such configurations presumably have uniformly positive probability on all scales.
last iteration, it is satisfiable on the next one. However, near disasters can occur causing “bottlenecks”–situations where one of a relatively few bonds must be occupied in order to achieve a spanning cluster. This would have a tendency to drive us to higher values of $f$.

We believe that these bottlenecks do occur and, in fact, are responsible for the relatively broad tails in the distribution of $f$ in the region $p(\beta_c) < f < 1$ that have been observed in our two-dimensional simulations. However, we also believe that these events affect only the details of how the $L \to \infty$ limit is achieved, not the limit itself since there are alternate routes circumventing bottlenecks occurring on all scales. Nevertheless, the finite-size scaling is sometimes quite complicated and, in certain instances, we must resort to semi-empirical fitting of the data.

An interesting feature of the IC algorithm (using spanning rules) is that the approach to equilibrium is along a critical trajectory. For example, if the starting configuration is characteristic of zero temperature, the initial bond configuration is typical of ordinary bond percolation at threshold. Thus some sort of power law correlations are actually established on the first step.

IV. NUMERICAL METHODS AND RESULTS

A. Implementation of the algorithm

The most difficult part of the IC algorithm is the construction of a cluster configuration from a spin configuration. The first step is to produce a random permutation of the set, $E$ of bonds of the lattice ($|E| = dL^d$ here). This is accomplished through $|E|$ random pairwise permutations. Initially, let $\pi : \{1, \ldots, |E|\} \to E$ be some conventional initial order on $E$. For $j = 1$ to $|E|$, $\pi$ is updated by choosing a random number, $r$ in the range $j$ to $|E|$; then the $j^{th}$ and $r^{th}$ elements of the permutation are interchanged, $\pi(j) \leftrightarrow \pi(r)$. It is well known that after $|E|$ steps, $\pi$ is a random permutation. The computational work involved in making the random permutation is nearly linear in the number of bonds.
Bonds are explored in the order given by the random permutation. If a bond is satisfied it is added to the cluster configuration. The data structure for the cluster configuration and its updating is done in the same general way as for other cluster and percolation algorithms using the Hoshen-Kopelman (or “disjoint set forests”) method. Each cluster of sites is described as a rooted tree and when two clusters are combined, the root of the smaller cluster becomes a son of the root of the larger cluster. When two clusters of the same size are combined the conventional direction associated with the bond that joined the clusters determines which site is to be the root. Information concerning the current state of the cluster as a whole, such as its mass, is stored with the root.

After the cluster configuration is updated by the addition of a bond it is necessary to check whether the current cluster satisfies the stopping rule. For the fixed parameter rules this is straightforward. For the other spanning rules, it is necessary to associate with each site a vector from the site to the root of its cluster. The set of distance vectors, \( \{v_i\} \) is updated in the natural way when two clusters are combined. The sites in the larger cluster retain their previous coordinates relative to the root. The sites in the smaller cluster take new coordinates, \( v' \),

\[
v'_i \leftarrow v_i - v_k - e_{jk} + v_j
\]

where \( e_{jk} \) is the unit vector of the new bond added to the lattice, \( j \) is the site in the larger cluster which connects to \( k \) in the smaller cluster. For the topological rule, stopping can only occur if the new bond is added as an internal bond. If the new bond \( \langle j, k \rangle \) is an internal bond we evaluate,

\[
v_k^* \leftarrow v_j - e_{jk}.
\]

If \( v_k^* \neq v_k \) the current cluster is multiply connected and the topological rule is satisfied.

For the extension rule, each cluster must have associated with it the coordinates of the \( 2d \) sites which are the most distant from root along the \( d \) axes in the positive and negative directions. Updating these coordinates after two clusters are combined is somewhat tedious due to periodic boundary conditions and is described in \[30\].
The invaded cluster algorithm requires $1.8 \times 10^{-5}$ seconds per update per spin on a DEC Alpha 2100 workstation. The running speed is about a factor of two slower than for the SW algorithm.

For the results reported here we start with an initial ordered ($T = 0$) configuration. Unless otherwise stated, the system is allowed to equilibrate for two hundred Monte Carlo steps (MCs) before data collection. If no error bars are shown in a figure, the error is smaller than the symbol size.

**B. Continuous transitions**

1. **Three-dimensional Ising model**

Figure 1 shows data for the mean value of the ratio of occupied to satisfied bonds, $\langle f \rangle$ for the three-dimensional Ising model as a function of $L^{-1.59}$. The power of $L$ is chosen to approximate the inverse of the three-dimensional Ising correlation length exponent $1/\nu \approx 1.59$. Results for both the topological and extension spanning rules are shown. Finite-size corrections are smaller for the topological stopping rule. The best linear fit to the data for the topological rule yields $0.35803$ in comparison with a recent value $p(T_c) = 0.358098(7)$ from Ref. [31].

Using ideas made plausible by finite-size scaling theory we can obtain two independent critical exponents. Figure 2 shows $\log(\langle f \rangle_t - \langle f \rangle_e)$ plotted against $\log(L)$ for the three-dimensional Ising model where $\langle f \rangle_t$ is measured using the topological rule and $\langle f \rangle_e$ is measured using the extension rule. A fit to the data yields $\nu = .63$, which is in agreement with the value $0.6289(8)$ from [31]. A second independent exponent can be obtained either from the cluster size distribution of the scaling of the largest cluster. Figure 3 shows $n(s)$, the number of clusters per site of size $s$ with the data binned in octaves. This figure shows that the cluster size distribution is indeed self-similar and allows us to estimate the exponent $\tau$, defined by $n(s) \sim s^{-\tau}$. A straight line fit yields $\tau = 2.19$ compared to the accepted value.
of 2.21. A more efficient way to obtain a second independent exponent is via the fractal dimension of the spanning cluster. The average size of the spanning cluster is plotted against the system size in the inset of Fig. 3 from which we obtain $\beta/\nu = 2.45$ compared to the accepted value 2.47.

2. Critical two-dimensional Potts models

Figure 4 shows results for the extension rule applied to the two-dimensional Ising model. Both the mean and median value of $f$ are plotted against $L^{-1}$ (in accord with finite-size scaling since $\nu = 1$ for the two-dimensional Ising model) and are seen to converge to the exactly known value of $p(T_c)$. The fact that median lies below the mean shows that the distribution is skewed toward larger values of $f$. This is presumably caused by the simultaneous percolation of the spins discussed in Sec. III C. This is in contrast to the three-dimensional Ising model for which the $f$ distribution is very symmetrical. The inset to Figure 4 shows var($f$)$^{1/2}$ plotted against $1/L$. The solid line is a fit to the data whose leading behavior is $L^{-1/2}$. This curve supports the hypothesis that the distribution of $f$ becomes sharp as $L \to \infty$.

Figure 5 is a plot of the average energy per spin vs. $L^{-1}$ with the exact value plotted on the vertical axis. A fit to the data of the form, $\varepsilon_0 + \varepsilon_1 L^{-1} + \varepsilon_2 L^{-2}$ yields, $\varepsilon_0 = -1.706$ which is reasonably close to the exact value $-1.7071 \ldots$ Energy fluctuations are shown in the inset of Fig. 5. The quantity var($\varepsilon$)$N$ is seen to increase roughly linearly in $L$. This is in contrast to the canonical ensemble where var($\varepsilon$)$N$ is the specific heat and diverges logarithmically in $L$ for the two-dimensional Ising model. The behavior of energy fluctuations underscores the difference between the IC ensemble and the canonical ensemble.

The top panel of Fig. 6 shows the fraction of occupied bonds vs. the mass of the largest cluster for the mass rule. Data collapse for a range of system sizes predicted by the finite-size

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In this section, $\varepsilon$ refers to the energy per spin rather than the energy per bond.
scaling ansatz,
\[ \langle f \rangle - p(T_c) L^{1/\nu} \sim G(mL^{\beta/\nu}). \] (16)

is confirmed in the lower panel. These results demonstrate that the IC algorithm can be used to extract quantitative results for the critical temperatures and critical exponents using systems of modest size.

Figure 7 is a log-log plot of the deviation of \( f \) from its exact value versus the system size for two-dimensional Potts models with continuous transitions, \( q = 1, 2, 3, 4 \). Except for the Ising case we have used the topological rule. The extension rule is used for the Ising case. Figure 8 is a log-log plot of \( \text{var}(f)^{1/2} \) vs. the system size for the two-dimensional Potts models with continuous transitions. The figure shows that the \( f \) distribution becomes narrow as a power of the system size \( L \). Fitting the last five data points for each \( q \) to the form, \( \text{var}(f)^{1/2} \sim L^{-b(q)} \) yields \( b(1) = .71(1), b(2) = .46(2), b(3) = .30(2) \) and \( b(4) = .23(1) \). For percolation this result is close to the expected scaling, \( b(1) = 1/\nu(1) \). For invasion percolation it is believed that the full \( f \) distribution scales with \( L^{-1/\nu(1)} \). For the other values of \( q \), \( b(q) \) is much smaller than \( 1/\nu(q) \) and decreasing with \( q \). We do not yet understand the finite-size scaling of the \( f \) distribution.

C. First order transitions in Potts models

1. Three-dimensional 3-state Potts model

We first discuss results for the topological and extension stopping rules for the three-dimensional 3-state Potts model. This model has a weak first order transition. Fig. 9 shows the mean and median values of \( f \) vs. \( L^{-1} \) for system sizes between 10 and 70 for the topological stopping rule and between 10 and 110 for the extension stopping rule. Data was taken from samples of \( 10^4 \) MCs for each lattice size up to \( L = 70 \) and between 3000 and 6000 MCs for the larger sizes. In the case of the topological stopping rule, a fit of the mean to a function of the form \( c_0 + c_1 L^{-1/2} + c_2 L^{-1} + c_3 L^{-2} \) gives \( c_0 = 0.4232 \). This
is in good agreement with results obtained with other methods. With the topological stopping rule, the median shows no finite-size effects within the error, as is the case for the three-dimensional Ising model. It can be fitted to a practically horizontal line, which extrapolates to 0.4228. Again the median seems to be the better choice if predictions about the infinite system are to be made, although it tends to be more noisy. As in the case of the two-dimensional Ising model, the difference between the mean and the median results from a tail in the distribution of \( f \) towards \( f = 1 \).

For the extension stopping rule, the median enters a flat region starting at about \( L = 40 \). The arithmetic mean of the last seven data points is \( 0.42336 \). Current values of \( T_c \) agree in the first four digits, and so does the value that we obtained from the median in this way. Again, the mean is above the median and starts to approach it only for very large system sizes (\( L \geq 80 \)).

2. Mass rule for first-order transitions

The above results show that spanning rules may be used to accurately locate a weak first-order transition however, they perform poorly for strong first-order transitions (large values of \( q \)). The difficulty is that the \( f \) distribution becomes increasingly broad and asymmetric with a tail extending toward \( f = 1 \). We believe that the tail in the \( f \) distribution is related to the way the spanning condition is met for strong first-order transitions. The spanning cluster is a nearly linear object which extends across the system in a background of small clusters whose size is presumably the correlation length. For large values of \( q \) the spanning cluster is very narrow (somewhat like a river running through a terrain of small clusters). This observation is consistent with the increase in \( p(T_c) \) with increasing \( q \). In addition, for large \( q \), there are severe bottlenecks; one of only a few bonds must be occupied to meet the spanning condition. This leads to a broad \( f \) distribution with a tail toward \( f = 1 \). Although we believe the \( f \) distribution becomes sharp for large \( L \) the convergence is very slow. An additional difficulty in using either of the spanning rules for strong first-order transitions
arises from the fact that the spanning cluster is nearly reproduced in successive Monte Carlo steps so that the autocorrelation time is large.

These problems can be avoided with the mass stopping rule. At the phase coexistence temperature $T_c$, the magnetization may take any value from 0 to $m_l$ where $m_l$ is the magnetization of the pure low temperature phase. Thus we expect $f$ to approach $p(T_c)$ for every $m$ between 0 and $m_l$. At $m_l$ we expect the derivative of $f(m)$ to jump to a finite value as the systems enters the ordered phase with $T < T_c$.

Figure 10 shows a plot $\langle f \rangle$ vs. $m$ for the two-dimensional 10-state Potts model, obtained from the mass rule for system sizes 50, 100 and 200. Data was obtained after an equilibration of 1000 MCs from a sample of $2 \times 10^4$ to $5 \times 10^4$ MCs. The dashed line denotes the exact value of $p(T_c) = \sqrt{q}/(1 + \sqrt{q})$. It is clear that the crossing point of these curves (near $m = 0.6$) for different system sizes provides an accurate estimate of $p(T_c)$. Furthermore, the curves become increasing flat for large $L$ and presumably converge (non-uniformly) to $p(T_c)$. The value of $m_l$ can be estimated from the largest value of $m$ for which $\langle f \rangle = p(T_c)$. From data for $L = 50, 100, 150, 200$ and 500 (the data for 150 and 500 is omitted from the plot for clarity) we find convergence to the value, $m_l = 0.8544$.

For small values of $m$ we find a region which moves increasingly toward zero where $f$ is significantly greater than $p(T_c)$. The non-monotonicity of $f$ as a function of $m$ occurs only for those Potts models with first-order transition (compare Fig. 12).

In the inset of Fig. 10 the standard deviation of $f$ is plotted vs. $1/L$. This quantity appears to vanish as $L \to \infty$ though at a rate that depends on $m$. The data is consistent with our belief that the $f$ distribution approaches a delta function at $p(T_c)$ for any $m < m_l$. Any mixture of low and high temperature phases in the coexistence region can be sampled by fixing the ratio, $m/m_l$. This argument can be confirmed by looking at the energy per spin. Let $n_l$ be the fraction of the system that is in the low temperature phase and assume that it is proportional to $m$. If $\varepsilon_l$ ($\varepsilon_h$) is the energy per spin of the pure low (high) temperature phase, we expect the energy per spin $\varepsilon$ to behave like
\[ \langle \varepsilon \rangle = (m/m_l)\varepsilon_l + (1 - m/m_l)\varepsilon_h + \varepsilon_{sf}A/N \]  \hspace{1cm} (17)

at the transition point. The third term includes the interfacial energy per unit area \( \varepsilon_{sf} \) and the area \( A \) and should vanish with system size like \( 1/L \).

Fig. 11 shows \( \langle \varepsilon \rangle \) vs. \( m \) for \( L = 50, 100 \) and 200 from the same runs as the data of Fig. 10. Error bars obtained with the jack-knife method are smaller than the symbols. There is a large region from small \( m \) to about \( m_l = 0.855 \) where the energy is described by a line with negative slope plus a correction that vanishes as \( L \) becomes large. Again, this statement is also based on additional data for \( L = 150 \) and 500. This behavior is in good agreement with Eq. 17. At \( m_l \) the energies for different \( L \) collapse, as there is no interfacial energy left, and the systems enter the ordered phase below \( T_c \), causing the energy to drop. Only for small \( m \) the energies leave the presumed curve due to finite size effects. In order to estimate the energy of the high temperature phase, we fitted the \( L = 200 \) data points in the range \( 0.05 \leq m \leq 0.5 \) to a function of the form \( c_0 + c_1m + c_2m^2 \), that gives \( \varepsilon_h = c_0 = -0.969 \), which agrees well with the exact result \( -0.9682 \ldots \). Evaluating the data for the biggest system used, \( L = 500 \) at \( m_l \) gives \( e_l = -1.661 \), which is close to the exact value \( -1.6642 \ldots \) as well.

There are few computational methods for reliably distinguishing the order of a phase transition 34. The 3-state Potts model in three dimensions for example at the phase transition point in many respects behaves just like a second-order transition. If the magnetization rule is used, however, even weak first-order transitions seem to behave differently than continuous ones. We observed that \( f \) is a non-monotonic function of \( m \) for first-order transitions. Figure 12 shows the median value of \( f \) vs. \( m \) for Potts models for several values of \( q \). The known values of \( p(T_c) \) are marked by dashed lines. Note that the curves are monotone increasing for models with second-order transitions, \( (q = 2, 4) \) and non-monotonic for models with first-order transitions \( (q = 5, 6 \, \text{and} \, 10) \). Non-monotonicity is also found for the three-dimensional 3-state Potts model, see Fig. 13. It should be noted that both the three-dimensional 3-state and two-dimensional 5-state Potts models have extremely weak
first-order transitions so that this criterion is quite sensitive. It is useful even if the correlation length is larger than the system size.

D. Dynamics of the IC algorithm

In this section we study the dynamic properties of the IC algorithm. The normalized autocorrelation function of an observable $A$ is defined by,

$$
\Gamma_A(t) = \frac{\langle A_0 A_t \rangle - \langle A \rangle^2}{\langle A^2 \rangle - \langle A \rangle^2},
$$

where $t$ is time in Monte Carlo steps. The three sets of points in Fig. [3] are the normalized autocorrelation functions of the absolute value of the magnetization $m$, energy $\varepsilon$, and fraction of occupied bonds $f$ for the two-dimensional Ising model. Numerical data were collected for the topological stopping rule from a run of $10^4$ MCs which was divided into 10 groups with errors estimated by the jack-knife method. In a few steps all three autocorrelation functions have nearly vanished. The autocorrelation functions of $f$ and $\varepsilon$ display a negative overshoot on the first step which becomes larger for larger system sizes.

The integrated autocorrelation time $\tau_A$, which is required for estimating the errors in measuring the observable $A$, is defined by,

$$
\tau_A(w) = \frac{1}{2} + \sum_{t=1}^{w} \Gamma_A(t),
$$

$$
\tau_A = \lim_{w \to \infty} \tau_A(w).
$$

The integrated autocorrelation time determines the size of the standard error in measuring $A$ according to,

$$
\delta A = \left(2\text{var}(A)\tau_A/N_{MC}\right)^{1/2}
$$

with $\text{var}(A)$ the variance in $A$ and $N_{MC}$ the number of Monte Carlo steps. Figure [15] is a plot of $\tau_A(w)$ with $A = m$, $\varepsilon$ and $f$ as a function of $w$ for the two-dimensional Ising model. Similar behavior was found for the three-dimensional Ising model. The error for $\tau_A(w)$ was
estimated by taking the square root of the sum of the variances of $\Gamma_A(t)$’s for $t \leq w$. For all three observables, $\tau_A(w)$ reaches a plateau in a few steps.

Table I is a summary of the integrated autocorrelation time for the two and three-dimensional Ising model at $w = 6$, where all the $\tau$’s have saturated but still have relatively small errors. The values for $\tau_\varepsilon$ are compared with results [4] for the SW and Wolff single cluster algorithm. The energy autocorrelation time is markedly smaller for the IC algorithm than the other two cluster algorithm. Furthermore, $\tau_\varepsilon$ decreases for larger systems while for the other cluster algorithms it increases. $\tau_m$ appears to be independent of system size suggesting the possibility that the dynamic exponent for IC dynamics is zero for both two and three-dimensional Ising models.

For the largest systems, $\Gamma_f(1)$ is close to $-0.5$ and, as a result, $\tau_f$ is very small. Although the data is not good enough to draw clear conclusions, it appears that $\tau_f$ approaches zero as a power, $L^{z_f}$ with $z_f \approx -1$. The anticorrelation in $f$ and $\varepsilon$ means that these quantities can be accurately estimated in a small number of Monte Carlo steps. Indeed, for averaging these variables, the IC algorithm is better than performing independent sampling from the invariant IC measure. For example, Eq. 20 and the behavior of $\text{var}(f)$, implies that $\delta f \sim L^{-a/\sqrt{N_{MC}}}$ where $a \approx 1.3$.

Table II shows results for the integrated autocorrelation time for the 3 and 4-state Potts models. We find again that the IC algorithm is much faster than the SW algorithm though for $q = 3$ and 4 there appears to be some critical slowing. Based on Table II we can obtain estimates for the dynamic exponent for the magnetization, $z_m(3) = .28$ and $z_m(4) = .63$. Note however that these values are less than the Li and Sokal [35] bound for the dynamic exponent for the SW algorithm ($z \geq \alpha/\nu$).

V. SUMMARY

The invaded cluster method comprises a class of algorithms for sampling equilibrium spin systems. Because cluster growth is controlled by a spanning rule rather than the
temperature, the method is able to simulate the phase transition point without \textit{a priori} knowledge of the phase transition temperature. The transition temperature is, instead, an output of the algorithm. We have demonstrated this numerically for Ising/Potts models in two and three dimensions.

We may also use parameterized stopping rules to explore either the coexistence region of discontinuous transitions or the critical region near a continuous transition. For these rules we specify a quantity such as the energy or susceptibility and learn the corresponding temperature. Using the mass rule we have been able to sweep through the coexistence region of first-order transitions and to obtain quantities such as the energy of the high and low temperature coexisting phases. The behavior of the effective transition temperature with the mass parameter apparently yields a very sensitive method to distinguish continuous from discontinuous transitions.

The invaded cluster algorithm is very similar to the Swendsen-Wang algorithm except that the occupied bonds are determined by a stopping rule rather than the temperature. We argued that this leads to a feedback mechanism that forces the system to the desired equilibrium state much faster than is the case for the Swendsen-Wang algorithm. The consequence is that the algorithm is extremely fast. Measured autocorrelation times are less than unity and decrease with system size for the energy and estimated critical temperature. The magnetization integrated autocorrelation time is constant for the two and three-dimensional Ising models but grows slowly for the 3 and 4-state two-dimensional Potts models. We speculate that the invaded cluster algorithm applied to Ising critical points has no critical slowing down. For this reason and because there is no need to know the transition temperature, we believe the IC method will prove to be the most efficient approach for high precision measurements of critical properties.

Although we have tested the algorithm in a number of setting and supplied non-rigorous arguments for its validity much work remains to be done in understanding the method and putting it on a firm footing. It is important to prove that the IC ensemble is equivalent to the usual statistical mechanics ensembles. We would also like to understand the finite-size
scaling properties of the IC ensemble since these differs from our naive expectation in some cases.

In this paper we have confined our attention to Potts models however the method is much more broadly applicable. In a future paper \cite{16} we will show how to use the approach for a variety of discrete spin systems such as the Ashkin-Teller model. Similarly, the embedding approach described by Wolff can be used to generalize the method to $O(n)$ models.

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**NOTE ADDED**

After completing this research we received an interesting preprint \cite{36} that describes a “fixed cluster” algorithm. This algorithm uses a stopping rule based on the extent $l$ of the largest cluster. However, in contrast to the fixed parameter rules used here, $l$ does not correspond to a thermodynamic quantity.
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TABLE I. Integrated autocorrelation times for two- and three-dimensional Ising models for the SW, Wolff and IC algorithms. Results for the IC algorithm are measured at time step $w = 6$.

| d | L | $\tau_{\varepsilon,SW}^a$ | $\tau_{\varepsilon,Wolff}^a$ | $\tau_{\varepsilon,IC}$ | $\tau_{m,IC}$ | $\tau_{f,IC}$ |
|---|---|----------------|----------------|----------------|----------------|----------------|
| 2 | 32 | 4.13(4) | 1.80(1) | .51(3) | .88(3) | .19(3) |
| 2 | 64 | 4.92(8) | 2.23(3) | .42(2) | .78(3) | .11(2) |
| 2 | 128 | 6.00(8) | 2.69(4) | .42(3) | .80(3) | .07(3) |
| 2 | 256 | 3.17(8) | .37(3) | .81(3) | .06(3) | |
| 3 | 16 | 5.6(1) | 1.36(2) | .35(2) | .65(2) | .09(2) |
| 3 | 24 | 6.8(1) | 1.50(3) | .27(2) | .62(3) | .07(2) |
| 3 | 32 | 7.8(3) | 1.72(4) | .25(2) | .65(2) | .05(2) |
| 3 | 48 | 9.9(4) | 1.90(6) | .19(2) | .66(3) | .02(3) |

$^a$Ref. [37].
TABLE II. Integrated autocorrelation times for two-dimensional, 3 and 4-state Potts models for the SW and IC algorithms. Results for $\tau_{\varepsilon,IC}$ and $\tau_{f,IC}$ for IC dynamics are measured at time step $w = 6$ while the time step $w_m$ for $\tau_{m,IC}$ is shown in the last column. Results for SW dynamics are for sizes 128 and 256 rather than 120 and 250.

| q | L  | $\tau_{\varepsilon,SW}^b$ | $\tau_{\varepsilon,IC}$ | $\tau_{f,IC}$ | $\tau_{m,IC}$ | $w_m$ |
|---|----|---------------------|---------------------|-------------|-------------|--------|
| 3 | 120| 30.3(1.2)           | .73(3)              | .08(3)      | 1.40(4)     | 6      |
| 3 | 250| 39.6(1.7)           | .59(2)              | .06(2)      | 1.73(5)     | 11     |
| 3 | 500| .52(2)              | .06(3)              |             | 2.08(6)     | 15     |
| 4 | 120| 115.7(6.1)          | 1.23(2)             | .11(3)      | 2.97(7)     | 16     |
| 4 | 250| 232.0(24.6)         | 1.10(3)             | .09(2)      | 4.61(10)    | 27     |
| 4 | 500| 0.88(3)             | .05(2)              |             | 7.31(16)    | 57     |

$^b$Ref. [35].
FIGURES

FIG. 1. $\langle f \rangle$ vs. $L^{-1/\nu}$ for the three-dimensional Ising model using different stopping rules. The infinite volume estimate of $p(T_c)$ from Ref. [31] is shown on the vertical axis.

FIG. 2. Double logarithmic plot of $\langle f \rangle_t - \langle f \rangle_c$ vs. $L$ for the three-dimensional Ising model.

FIG. 3. Double logarithmic plot of the distribution of cluster sizes $n(s)$ for the three-dimensional Ising model. The inset shows a double logarithmic plot of the average size of the largest cluster.

FIG. 4. The mean and the median of $f$ vs. $1/L$ for the two-dimensional Ising model. The solid line shows a linear fit to the median and the exact infinite volume value is shown on the vertical axis. The inset shows the standard deviation of the $f$ vs. $1/L$. A least squares fit to the form $c_0 + c_1 L^{-1/2} + c_2 L^{-1}$ (solid line) suggests that the distribution becomes sharp in the infinite volume limit.

FIG. 5. $\langle \varepsilon \rangle$ vs. $1/L$ for the two-dimensional Ising model. The solid line is a fit to the form $\varepsilon_0 + \varepsilon_1 L^{-1} + \varepsilon_2 L^{-2}$ and the exact infinite volume result is shown on the vertical axis. The inset shows $\text{var}(\varepsilon)N$ vs. $L$ with a linear fit through the data.

FIG. 6. The mass rule applied to the two-dimensional Ising model. The upper graph shows $\langle f \rangle$ vs. $m$. In the lower graph the same data is scaled as described in the text.

FIG. 7. Double logarithmic plots of $|f - p(T_c)|$ vs. system size $L$ for the two-dimensional $q$-state Potts models. Exact values of $p(T_c)$ are used.

FIG. 8. Double logarithmic plots of the standard deviation of $f$, $\text{var}(f)^{1/2}$ vs. system size $L$ for the two-dimensional $q$-state Potts models.

FIG. 9. The mean and median of $f$ for the three-dimensional 3-state Potts model. The upper graph shows results from the topological stopping rule, the solid line is a linear fit to the median. The lower graph shows results from the extension stopping rule.
FIG. 10. $\langle f \rangle$ vs. $m$ for the two-dimensional 10-state Potts model using the mass rule. The dashed line marks the exact value, $p(T_c)$. The inset shows the standard deviation of $f$ vs. $1/L$ for $m = 0.4$ (squares) and $m = 0.85$ (triangles).

FIG. 11. $\langle \varepsilon \rangle$ vs. $m$ for the two-dimensional 10-state Potts model using the mass rule for several lattice sizes $L$. The solid line is a fit to the form $c_0 + c_1 m + c_2 m^2$ for the $L = 200$ data points with $0.05 \leq m \leq 0.5$, the intercept with the $m = 0$ axis is our estimate for $\varepsilon_h$, the energy of the high temperature phase at the transition.

FIG. 12. The median of $f$ vs. $m$ for the two-dimensional Potts models using the mass rule with $q = 2, 4, 5, 6,$ and $10$ and $L = 200$. The exact value of $p(T_c)$ for each $q$ is shown by a dashed line.

FIG. 13. $f$ vs. $m$ for the three-dimensional 3-state Potts model.

FIG. 14. Autocorrelation functions of the magnetization $m$, energy $\varepsilon$ and occupation fraction $f$ vs. time step $t$ for the two-dimensional Ising model for size, $L = 256$.

FIG. 15. Integrated autocorrelation times for $m$, $\varepsilon$ and $f$ vs. integration time, $w$ for the two-dimensional Ising model for size, $L = 256$. 