Numerical test of the Cardy-Jacobsen conjecture in the site-diluted Potts model in three dimensions.

L.A. Fernandez, 1,2 A. Gordillo-Guerrero, 3,2 V. Martin-Mayor, 1,2 and J.J. Ruiz-Lorenzo 4,2

1 Departamento de Física Teórica I, Universidad Complutense, 28040 Madrid, Spain.
2 Instituto de Biocomputación and Física de Sistemas Complejos (BIFI), 50009 Zaragoza, Spain.
3 Departamento de Ingeniería Eléctrica, Electrónica y Automática, Universidad de Extremadura, 10071 Cáceres, Spain.
4 Departamento de Física, Universidad de Extremadura, 06071 Badajoz, Spain.

(Dated: May 1, 2014)

We present a microcanonical Monte Carlo simulation of the site-diluted Potts model in three dimensions with eight internal states, partly carried out in the citizen supercomputer Ibercivis. Upon dilution, the pure model’s first-order transition becomes of the second-order at a tricritical point. We compute accurately the critical exponents at the tricritical point. As expected from the Cardy-Jacobsen conjecture, they are compatible with their Random Field Ising Model counterpart. The conclusion is further reinforced by comparison with older data for the Potts model with four states.

PACS numbers: 05.50.+q, 64.60.De, 75.40.Mg

I. INTRODUCTION

When two ordered phases compete, even a tiny amount of disorder is significant. Consider, for instance, the antiferromagnetic insulator LaCuO4. A small La ↔ Sr substitution turns it into a high-temperature superconductor. Also for colossal magnetoresistance oxides the importance of the combination of phase coexistence and chemical disorder has been emphasized.

These examples suggest a simple, yet general question: which are the effects of quenched disorder on systems that undergo a first-order phase transition in the ideal limit of a pure sample? (quenched disorder models impurities that remain static over experimental time-scales). In fact, this question has been relevant in a large number of physical contexts. A non-exhaustive list includes nanoscale ferroelectricity, 11 tilt ordering, 12 ferroelectric thin films, 13 random block copolymers, 14 ferroelectric nanodisks, 15 topological phases in correlated electron systems 16 effects of multiplicative noise on electronic RLC circuits 17 and surface waves. 18

Unfortunately, only for two spatial dimensions (D = 2) we have a good understanding of the effects of quenched disorder on phase coexistence: the slightest concentration of impurities switches the transition from first-order to second-order. 19,20

In D = 3 we lack a general description. One should consider two different possibilities: disorder may couple either to the order parameter, as in the Random Field Ising Model (RFIM) 21,22 or it may couple to the energy, as in the disordered Potts model. 23 In both cases, quenched disorder is unreasonably efficient at softening the transition. It has been surprisingly difficult to show that the transition actually remains of the first-order for some range of impurity concentration. 24,25

Actually, the Cardy and Jacobsen conjecture relates the two types of disorder by means of a mapping between the RFIM and the disordered Potts model. 26 The conjecture reads as follows. Consider a ferromagnetic system undergoing a first-order phase transition for a pure sample. 27 Let T be the temperature and p be the concentration of magnetic sites (see the generic phase diagram in Fig. 1). A transition line, Tc(p) separates the ferromagnetic and the paramagnetic phases in the (T, p) plane. In D = 3 a critical concentration is expected to exist, 1 > p* > 0, such that the phase transition is of the first-order for p > p* and of the second order for p < p* (at p* one has a tricritical point). When p approaches p* from above, the latent-heat and the surface tension vanish while the correlation-length diverges. The corresponding critical exponents can be obtained from those of the RFIM (see below).

However, the Cardy-Jacobsen mapping relates two problems unsolved in D = 3. In particular, the RFIM (the supposedly well-known partner in the conjecture) suffers from severe inconsistencies between analytical, experimental and numerical work. On the experimental side, mutually inconsistent results for the correlation-length exponent 28 were obtained, 29 due to the uncertainties in the parameterization of the scattering line shape. Also, the estimate of the anomalous dimension = 3 violates hyperscaling bounds. 23,24 Numerical deter-
minations of exponent $\nu$ are scattered on a wide range and hyperscaling-violating results have been reported. The order-parameter’s critical exponent $\beta \sim 0.01$ is so small (yet, see Ref. [25], that it has even been conjectured that the transition could be of the first order [26].

On the other hand, the investigation of the disordered Potts model has been mostly numerical up to now. In the conventional approach, one averages over disorder the free-energy at fixed temperature [29]. It works nicely for the second-order part of the critical line $T_c(p)$ [19,28] but the first-order piece is plagued by huge sample-to-sample fluctuations of the specific-heat or the magnetic susceptibility [23]. Fortunately, these wild fluctuations can be avoided by averaging over disorder the entropy obtained from microcanonical Monte Carlo [24] at fixed energy [19]. We investigated in this way the site-diluted Potts model with $Q = 4$ states. A delicate system size showed that $p_i < 1$. Unfortunately, the relevance of the RFIM universality class for the tricritical point (the core of Cardy and Jacobsen conjecture) could not be addressed up to now.

Here we show that the Cardy-Jacobsen conjecture is verified to high numerical accuracy in the site-diluted Potts model with $Q = 4$ and 8 states. This results follows from a finite size scaling analysis of old $Q = 4$ data [19] and new, extensive Monte Carlo simulations for $Q = 8$, partly carried out in the Ibercivis citizen supercomputer [25]. Our analysis benefits from a recent computation of the RFIM critical exponents [18] that also exploits the redefinition of the disorder average [19].

In Section II, we summarize the main implications of the Cardy-Jacobsen conjecture and define our specific model. Our methodology, including details on simulation and statistical analysis, is presented in Section III. In Section IV, we present our main numerical evidences for the validity of the conjecture. We give our conclusions in Section V. Finally in the appendix, we describe how the Control Variates technique improves the determination of some important quantities.

II. THE CARDY-JACOBSEN CONJECTURE

Specifically, we consider the $D = 3$ site-diluted Potts model with $Q$ internal states [17]. The spins, $\sigma_i = 1, \ldots , Q$, occupy the nodes of a cubic lattice of linear size $L$, with periodic boundary conditions. Each spin interacts with its nearest neighbors through the Hamiltonian

$$\mathcal{H}^{\text{spin}} = - \sum_{\langle i,j \rangle} \epsilon_i \epsilon_j \delta_{\sigma_i \sigma_j}.$$  

The quenched randomness is represented by the occupation variables $\epsilon_i = 0, 1$ ($\epsilon_i = 1$ means that the $i$-th spin is present). We choose each $\epsilon_i$ independently, setting $\epsilon_i = 1$ with probability $p$. Each specific disorder realization is called a sample. The pure system is recovered for $p = 1$, where it undergoes a, generally regarded as very strong, first-order phase transition for $Q \geq 3$ [20,24]. We show in Fig. I the full phase diagram of this model.

The Cardy and Jacobsen mapping relates the large-$Q$ limit of the disordered Potts model to the RFIM [18]. At the tricritical point $p_i$ of the Potts model, we encounter three relevant scaling fields (see, e.g., Ref. [46]). The dilution field lies along the critical line $T_c(p)$. We name its scaling dimension $y_p$. The thermal scaling field has dimension $y_T$, and is responsible for the ferromagnetic transition when varying the temperature. Finally, the magnetic scaling field is related to an external magnetic field in Eq. (1). The mapping to the RFIM is

$$y_p = y_T^{\text{RFIM}} = - \frac{1}{\nu} \frac{\gamma_2}{\gamma_2 - 1},$$

$$y_T = y_T^{\text{RFIM}} = \frac{1}{2} (D - \theta + 2 - \eta),$$

where $\gamma$ is the correlation-length exponent [27], $\eta$ is the anomalous dimension, while $\theta$ is the hyperscaling-violations exponent [18]. Furthermore, the exponent of the surface tension $\mu$ verifies a modified Widom law: $\mu = D - \theta - 1$. Cardy and Jacobsen predicted as well that, upon approaching the tricritical point $p_i$, the latent heat in the diluted Potts model vanishes with the same exponent $\beta$ that rules the vanishing of the order parameter in the RFIM.

III. METHODOLOGY

A. The Microcanonical Ensemble

For the simulation of the model described by Eq. (1), we have used an extended microcanical method which is suitable to study the first order part of the transition line [24].

We will briefly review the main facts of this simulation approach. Using a mechanical analogy, each spin is complemented with a conjugated momenta. The total energy is the sum of a kinetic term, $\mathcal{K}$ (the halved sum of the squared momenta) and the potential energy, namely the spin Hamiltonian of Eq. (1).

We consider the microcanonical ensemble, where the energy (kinetic plus potential) is kept fixed to the total value $Ne$, where $N = \sum \epsilon_i$ is the total number of spins. The momenta can be explicitly integrated out. The entropy density $s(e)$ and the microcanonical weight $\omega(e,N;\{\sigma_i\})$ turn out to be

$$\exp[Ns(e,N)] = \frac{(2\pi N)^{\frac{D}{2}}}{\Gamma(N/2)} \sum_{\{\sigma_i\}} \omega(e,N;\{\sigma_i\}),$$

$$\omega(e,N;\{\sigma_i\}) = \left( \frac{\mathcal{K}}{N} \right)^{\frac{D-1}{2}} \theta^{\gamma} \mathcal{K},$$

The role of the Heaviside step function in Eq. (5) is preventing the kinetic energy from becoming negative.

The Monte Carlo simulation of the weight in Eq. (4) is straightforward. Both Metropolis and cluster methods are feasible and efficient [24]. In the present work, we have used the Swendsen-Wang algorithm [24] (see Ref. [19] for implementation details). One obtains in this way mean-values at fixed $e$ that will be denoted $\langle \cdots \rangle_e$. 


A particularly important mean-value comes from the Fluctuation-Dissipation relation

$$\frac{ds}{de} = \langle \hat{\beta} \rangle_e,$$  \hspace{1cm} (7)

where

$$\hat{\beta} = \frac{N - 2}{Ne - \mathcal{H}_{\text{spin}}}.$$  \hspace{1cm} (8)

On the view of Eq. (7), it might be inspiring to think of $\langle \hat{\beta} \rangle_e$ as the inverse-temperature corresponding to energy density $e$. The connection between the canonical and the microcanonical ensembles is discussed in Ref. 48. Finally, our main observable will be $\beta(e)$, defined as

$$\beta(e) = \langle \hat{\beta} \rangle_e.$$  \hspace{1cm} (9)

where the overline stands for the disorder-average as computed at fixed $e$.

**B. The Maxwell construction**

A standard way of studying phase-coexistence in a microcanonical setting is the Maxwell construction. This allows to compute from the curve $\beta(e)$ several important magnitudes: the critical inverse temperature $\beta_c$, the energies of the two coexisting phases and the surface tension. Furthermore, one may apply the very same method to the sample dependent $\langle \hat{\beta} \rangle_e$, as shown in Fig. 2. We follow the numerical methods described in Refs. 44 and 19. We briefly summarize them now, for the sake of completeness.

Consider the equation

$$\beta(e) = \beta_c \text{, or (single sample) } \langle \hat{\beta}_e \rangle_e = \beta_c.$$  \hspace{1cm} (10)

In normal situations, $\beta(e)$ is monotonically decreasing with $e$, so that Eq. (10) has a unique solution. However, at phase-coexistence $\beta(e)$ is no longer monotonically decreasing, see Fig. 2. Therefore, Eq. (10) has three important solutions, named $e^o$, $e^s$, and $e^d$ ($e^o < e^s < e^d$):

- The rightmost root of (10), $e^d_{L,\beta}$, corresponds to the “disordered phase”.
- The leftmost root of (10), $e^o_{L,\beta}$, corresponds to the “ordered phase”.
- The second rightmost root of (10), $e^s_{L,\beta}$ is a saddle-point among the two phases.

Note that these three solutions do depend on $L$, although we shall not explicitly indicate it unless necessary.

We compute the inverse critical temperature $\beta_c$ from the equal-area rule:

$$0 = \int_{\beta_c^d}^{\beta_c^o} de \ (\beta(e) - \beta_c),$$  \hspace{1cm} (11)

see Fig. 2. Note that the $\beta_c$ computed from Eq. (11) does depend on the system size. In fact, in the thermodynamic limit, Eq. (11) is a mere consequence of the continuity of the free-energy density (as a function of temperature) at the phase transition. In fact, recall that the free energy density can be expressed in terms of the inverse temperature and of the internal energy and entropy densities: $f = e - s/\beta$. Now, if we recall Eq. (7), we see that the equality of the free-energy densities of the ordered and the disordered phases at the critical temperature can be recast as

$$\beta_c (e^d_{\beta_c} - e^o_{\beta_c}) = s(e^d_{\beta_c}) - s(e^o_{\beta_c})$$  \hspace{1cm} (12)

$$= \int_{\beta_c^o}^{\beta_c^d} de \ (\beta(e) - \beta_c).$$  \hspace{1cm} (13)

This textbook reasoning can be extended to the more complicated case of a finite-system. In fact, it is easy to show, see Refs. 44 and 49 that Eq. (11) is identical to the criterion of equal-height in the energy histogram. Such a finite-system indicator of the critical temperature suffers from finite-size corrections of order $\sim 1/L^D$.

Once we know $\beta_c$, we may compute the latent-heat as

$$\Delta e = e^d_{\beta_c} - e^o_{\beta_c}.$$  \hspace{1cm} (14)

Finally, the surface-tension, $\Sigma$, is calculated as

$$\Sigma(L) = \frac{N}{2L^{D-1}} \int_{\beta_c^o}^{\beta_c^d} de \ (\beta(e) - \beta_c^o).$$  \hspace{1cm} (15)

Note that, in order to compute integrals such as the one in Eq. (11), we interpolate $\beta(e)$ (which is numerically computed
FIG. 3. (Color online) Maxwell construction, see Eq. (11), as obtained for the sample-averaged $\beta(e)$. Data for $L = 48$ and several values of the spin concentration. The transition becomes smoother as $p$ decreases (from bottom to top). In fact, for $p = 0.8$ the Maxwell construction can no longer be done (because the corresponding $\beta(e)$ is monotonically decreasing with $e$).

over a grid in the $e$-line), through a cubic spline. Statistical errors are computed using a jackknife method (see e.g. Ref. 44). In the case of the sample-averaged $\beta(e)$, the jackknife blocks are formed from the microcanonical mean-values obtained on the different samples. On the other hand, when one performs the Maxwell construction for a single sample as in Fig. 2 the jackknife blocks are formed from the Monte Carlo history. It is interesting to compare the curves $\beta(e)$ for fixed $L = 48$, as the disorder increases (i.e. as $p$ decreases), see Fig. 3. In the limit of a pure system, $p = 1$, $\beta(e)$ displays the expected cusps and steps for a system with well developed geometric and condensation transitions.22 As soon as the system becomes disordered, the transition becomes smoother: both the latent heat, see Eq. 14, and the surface tension, Eq. 15, are sizably smaller for $p = 0.95$ than for $p = 1$. This trend is maintained for decreasing $p$, to the point that the phase transition is clearly of the second order at $p = 0.8$ (for that dilution, $\beta(e)$ is monotonically decreasing with $e$). We note as well that the curve $\beta(e)$ for $p < 1$ is remarkably featureless, specially if compared to its $p = 1$ counterpart. Actually, geometric transitions are also found for individual samples at $p = 0.95$. However, the energies at which this singular behavior arise depend on the considered sample, which results in a smooth averaged $\beta(e)$.

C. Finite Size Scaling near a Tricritical Point

In the following we will discuss some relevant facts about the scaling near a tricritical point, see, e.g., Ref. 45 Consider some quantity $O$, that, in the thermodynamic limit, scales as $O_{L=\infty} \sim x^\xi$, where $\xi$ is the correlation length. The Finite Size Scaling (FSS) ansatz, tells us how the same quantity behaves in a finite system of size $L$. Close to the tricritical point at $(p_t, T_t = T_c(p_t))$

$$O(L, p_t + \delta p, T_t + \delta T) = L^\delta G(L^\gamma u_T, L^\rho u_p),$$

where $G$ is a scaling function, and we have neglected scaling corrections. As stated in Eqs. 2 and 3, there are two relevant scaling fields, the thermal field $u_T$ and the disorder field $u_p$. Both $u_T$ and $u_p$ are functions of $\delta p$ and $\delta T$, the deviations from the tricritical point. If we work at $u_T = 0$, we should expect that, at linear order, $u_p|_{u_T=0} \propto \delta p$. Then the phase transition is of the second order if $\delta p < 0$, and of the first order if $\delta p > 0$.

Our main assumption will be that the Maxwell construction, see Ref. 44 and the previous subsection, enforces the constraint $u_T = 0$ to an accuracy of order $O(L^{-D})$ (this expectation is well founded in the first-order part of the critical line). Hence, Eq. (16) simplifies to

$$O(L, p, \text{Maxwell}) = L^\delta G(L^\gamma (p - p_t))(1 + O(L^{-D})).$$

(17)

So, the Maxwell construction allows us to employ standard FSS with an effective scaling-corrections exponent $\omega = D - \gamma T$. The combination of Eqs. 2 and 3, standard RFIM scaling relations and the numerical estimates in Ref. 24 yield $\omega = \theta + \beta_{\text{RFIM}}/\nu_{\text{RFIM}} = 1.48(2)$. A further irrelevant scaling field $u_Q = 1/\log Q$ with exponent $-\theta$ is also present.24 Numerically, $\theta = 1.468(2)$, while we expect $\omega = 1.48(2)$. These two exponents are so similar that, given our limited numerical accuracy, we shall not attempt to distinguish them. However, we remark that one expects a larger amplitude of the scaling corrections for $Q = 4$, which is confirmed by our data (see Fig. 6).

D. Numerical Simulations and Thermalization Checks

We considered concentration values $0.65 \leq p \leq 1$ and lattice sizes $12 \leq L \leq 64$. The precise values are indicated in Table 1. The $p$-resolution becomes denser close to the $L$-dependent position of the tricritical point. For all pairs $(p, L)$ we simulated 500 samples, with the obvious exception of $p = 1$.

Each sample was simulated on a $e$-grid fine enough to allow for a correct spline interpolation, see Fig. 4. The simulations at the different $e$ values were mutually independent. Hence, we faced an embarrassingly parallel computational problem, suitable for Ibercivis (with a caveat, see below).

All samples were simulated for the same number of Monte Carlo steps, at every $e$ value. However, the number of Monte Carlo steps did depend on $e$, as we explain now. First, we ran all samples at a given $e$-value for a fixed amount of Swendsen-Wang steps (e.g. $3 \times 10^5$ for $L = 64$, or $2 \times 10^5$ for $L = 48$), then we assessed thermalization.

The thermalization check was the standard logarithmic data-binning: for any given value of $e$, we computed different estimates of the sample-averaged $\beta(e)$, using disjoint pieces of the Monte Carlo history. On the first bin, we included only the second-half of the Monte Carlo history (i.e., our safest data
from the point of view of thermalization). The second bin contained only the second quarter of the Monte Carlo history, etc. We checked for statistical compatibility, at least, among the first and second bins, see Fig. [4] If for a given value of $e$ the thermalization criterion was not met, the total simulation time was doubled. The procedure was cycled until convergence was achieved. We note that, for the concentrations nearest to $p = 1$, we encountered strong metastabilities, that prevented us from simulating $L = 128$ (that could instead be simulated for $Q = 4$ in Ref. [19]).

The thermalization protocol is not well suited for Ibercivis, because the simulation of a given sample at some difficult energy may last up to some days. Yet, Ibercivis relies on volunteers’ computers that frequently switch from on-line to off-line. To minimize the number of unfinished simulations, we have implemented a continuity system. It divides every simulation, no matter how long it is, in small time steps (typically 30 minutes). After every step, consistency checks are performed and the current system configuration is sent again to the simulation queue. This solved the problem for relatively long (5-6 hours) simulations but the few more demanding simulations were completed on local clusters.

Altogether, this work has consumed (the equivalent of) $3 \times 10^6$ hours of a single Intel Core2 duo at 2.5 GHz.

We should also mention that we have performed some new, short simulations for $Q = 4$ at $p = 0.95$, complementary to those reported in Ref. [19]. The simulated sizes were $L = 24$ and $L = 48$ (128 samples each). Our goal was to improve the accuracy of the interpolations described below.

### IV. RESULTS

To check the Cardy-Jacobsmen conjecture we have performed numerical simulations for $Q = 8$, hence further approaching the large-$Q$ limit where the mapping becomes exact [23].

Consider the $p$ and $L$ evolution of the latent-heat and the surface tension in Fig. [5]. If $p < p_t$, i.e. if we are in the second-order piece of the critical line, both $\Delta e$ and $\Sigma$ vanish in the large-$L$ limit (the two are positive for $p > p_t$). However, for small lattices, both $\Delta e$ and $\Sigma$ decrease gently upon decreasing $p$ which suggests that dilution merely smoothed the first-order transition. However, the curve for $\Delta e$ becomes sharper as we approach $p_t$ (for instance, the point of sharpest drop of $\Delta e$ in Fig. [5]—top) grows quickly with $L$. On the view of the $D = 2$ no-go theorems [12] one could be afraid that $p_t^L \to 1$ for large $L$ also in $D = 3$. We know that this is not the case [15], but it is clear that a careful scaling analysis is needed.

Eq. (17) tells us that $L^{2-\theta} \Sigma$ is scale-invariant, and thus allows to locate the tricritical point (because $x_t^L = \theta-D+1 = \theta-2$, $\theta = 1.469(20)$ [24]. Indeed, in Fig. [5]—bottom, we see

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**TABLE I**

| $L$ | Simulated $p$ values |
|-----|----------------------|
| 12  | 0.65, 0.675, 0.7, 0.725, 0.75, 0.775, 0.8, 0.825, 0.832, 0.85, 0.875, 0.9, 0.925, 0.9375, 0.95 |
| 16  | 0.65, 0.675, 0.7, 0.725, 0.75, 0.775, 0.8, 0.825, 0.85, 0.854, 0.875, 0.9, 0.925, 0.9375, 0.95 |
| 24  | 0.7, 0.725, 0.75, 0.775, 0.8, 0.825, 0.832, 0.845, 0.85, 0.875, 0.9, 0.925 |
| 32  | 0.75, 0.775, 0.8, 0.825, 0.85, 0.854, 0.8625, 0.875, 0.886, 0.8875, 0.9, 0.925, 0.9375, 0.95 |
| 48  | 0.75, 0.775, 0.8, 0.825, 0.85, 0.8625, 0.875, 0.877, 0.8875, 0.9, 0.925, 0.9375, 0.95 |
| 64  | 0.8, 0.825, 0.85, 0.86875, 0.875, 0.8875, 0.9, 0.925, 0.9375, 0.95 |

For each of the lattice sizes $L$, we indicate the values of $p$ (the concentration of magnetic sites) for which we carried out simulations. We shall need to regard the various quantities defined, as continuous functions of the density of magnetic sites, $p$. We shall need as well the corresponding $p$-derivatives. As a rule, we have obtained these functions of $p$ through a cubic-spline interpolation of the data computed at these $p$-values. In fact, some of them were chosen in order to minimize the interpolation errors at some particularly important values of $p$, see Tables [I] and [III]. Derivatives with respect to $p$ were computed simply by derivating the cubic-spline interpolating function. The error estimates where obtained through a jack-knife (see for instance Ref. [46] over the sample-averages.
that the curves for system sizes $L_1 < L_2$ cross at $p^{L_1,L_2}_t$:

$$L_1^{2 - \theta} \Sigma(L_1, p^{L_1,L_2}_t) = L_2^{2 - \theta} \Sigma(L_2, p^{L_1,L_2}_t),$$  \hspace{1cm}  (18)

($p^{L_1,L_2}_t \to p_t$ when $L_1$ diverges). We recall that a similar method was used recently in a spin-glass context. There are two main consequences of choosing a wrong estimate of exponent $\theta$ in Fig. 5—bottom and Eq. (18). First, in the limit of large lattice sizes, the height of the crossing point diverges (or goes to zero) if $\theta$ is underestimated (overestimated). Second, the size corrections to the crossing points are larger for a wrong $\theta$. Specifically, $p_t^{L_1,L_2} - p_t = O(L_1^{-\gamma_p})$. The amplitude for these scaling corrections cancels only for the exact choice of $\theta$.

The critical exponent for a quantity $O$ is obtained from its quotients at $p^{L_1,L_2}_t$:

$$\left. \frac{O(L_2)}{O(L_1)} \right|_{p^{L_1,L_2}_t} = \left( \frac{L_2}{L_1} \right)^{\alpha_o} \left[ 1 + A_O \left( \frac{1}{L_2^{\omega}} - \frac{1}{L_1^{\omega}} \right) \right].$$ \hspace{1cm} (19)

Above, we included only the leading scaling-corrections ($A_O$ is an amplitude). We use Eq. (19) for the logarithmic $p$-derivative of $\Sigma$ (scaling dimension $x = \gamma_p$), and for the latent heat (scaling dimension $x = \gamma_p$), which should be $\beta^{RFIM}/\nu^{RFIM}$, according to Cardy and Jacobsen. Our results are in Table III ($Q = 8$), and Table III ($Q = 4$). In both cases we see that the convergence of $p^{L_1,L_2}_t$ to the thermodynamic limit is very fast. The height of the crossing point seems also stable with growing sizes.

The results in Tables II and III need to be extrapolated to the limit of infinite system sizes. This can be done by considering leading order scaling corrections, as in Eq. (19). The extrapolation greatly improves by imposing to $Q = 4$ and 8 a common extrapolation and the same scaling-corrections exponent $\omega$, as required by the University predicted in Ref. 13.

In this way, we obtain $\beta^{y_p} = 0.0022(48)(3)$ and $\omega = 1.36(8)(1)$ where the second parenthesis indicates the uncertainty induced by the error in $\theta$ [4]. The fit quality is assessed through the $\chi^2$ test. We obtain $\chi^2/dof = 8.5/14$, which is almost too good (dof stands for the number of degrees of freedom of the fit). Indeed the probability of getting such a low value of $\chi^2$ with 14 degrees of freedom is only 14%. We note as well that $\beta^{y_p} = 0.0022(48)(3)$ is only barely compatible with the best RFIM estimate $\beta^{y_p} = 0.0119(4)$ [34] (since the discrepancy is as large as two standard deviations).
At this point, we can try to disproof universality. We make the assumption that $\beta y_p$ takes exactly the RFIM value, redo the fit and see the outcome of the $\chi^2$ test. This second fit, with $\beta y_p$ fixed to 0.0119, turns out to be perfectly reasonable ($\chi^2/\text{dof} = 14/15$, see Fig. 6). Hence we conclude that our data set is statistically compatible with universality.

A second, unexpected bonus of fixing $\beta y_p$ in the fit to the RFIM value, is a remarkable increase in the accuracy of the fit and see the outcome of the $\chi^2$ test. This second fit, with $\beta y_p$ fixed to 0.0119, turns out to be perfectly reasonable ($\chi^2/\text{dof} = 14/15$, see Fig. 6). Hence we conclude that our data set is statistically compatible with universality.

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Our $y_p$ is in the lower range of previous numerical and analytical estimates: $0.73 \leq 1/\sqrt{\text{RFIM}} \leq 1.12$.[14][55] Hyperscaling and our $y_p$ implies a slightly positive specific heat exponent $\alpha = (2y_p - D + \theta)/y_p = 0.03(10)$, in agreement with experimental claims of a (possibly logarithmic) divergence.[56] We warn however that severe hyperscaling violations (namely $\alpha = -0.63(7)$) have been reported in numerical work.[55]

One may compute as well the exponent $\theta$, by fitting $\Sigma(L, p_i^{L/2L}) = A_Q L^\theta(1 + B_Q L^{-\Delta})$ (only the amplitudes $A_Q$ and $B_Q$ are $Q$-dependent on the fit). Taking $\omega = 0.15(1)$, we obtain $\theta = 1.52(11)/2$, with an acceptable fit ($\chi^2/\text{dof} = 4.9/3$). The result is compatible with, but less accurate than, the latest RFIM result $\theta = 1.469(20)$.[55]

In summary, we have presented a finite-size scaling analysis of the tricritical point of the site-diluted Potts model in three dimensions for $Q = 4$ and 8 internal states. By considering leading-order scaling corrections we were able to show that the relevant Universality class for the tricritical point is the one of the RFIM. To our knowledge, this is the first verification of the Cardy-Jacobsen conjecture.[13]

Three technical ingredients were crucial to obtain this achievement: the use of the microcanonical Monte Carlo,[54] a new definition of the disorder average,[10] and the use of the citizen supercomputer Ibercivis.[53]

VI. ACKNOWLEDGEMENTS

We have been partly supported through Research Contracts Nos. FIS2009-12648-C03 and FIS2010-16587 (MICINN), GR10158 (Junta de Extremadura), ACCVII-08 (UEX), and from UCM-Banco de Santander. We thank Ibercivis for the equivalent of $3 \times 10^6$ CPU hours. The simulations were completed in the clusters Terminator (BIIF) and Horus (U. Extremadura). We also thank N. G. Fytas for a careful reading of the manuscript.

Appendix A: Control variates

The statistical quality of data may sometimes be significantly increased by means of a very simple trick, named control variates (see e.g. Ref. 58).

In short, we want to improve our estimation of a stochastic variable $A$ through its correlations with another random variable $B$ ($B$ is named a control variate). If $\overline{B} = 0$ and $\overline{A} = A + \alpha B$, then the expectation value does not change: $\overline{A} = \overline{A}$. How-
without CV

ever, depending on the arbitrary election of $\alpha$, we can get $\text{var}(\hat{A}) < \text{var}(\hat{A})$. The $\alpha$ election minimizing the variance $\text{var}(\hat{A})$ is

$$\hat{\alpha} = \frac{\text{cov}(A, B)}{\sqrt{\text{var}(A)\text{var}(\hat{B})}},$$  \hspace{0.5cm} (A1)$$

which coincides with the correlation coefficient $r_{AB}$. The optimal variance is

$$\text{var}(\hat{A}) = \text{var}(A)\left(1 - r_{AB}^2\right).$$  \hspace{0.5cm} (A2)$$

Hence, the stronger the statistical correlation (or anticorrelation) between $A$ and $B$, the more effective the control variate is.

In our case, a rather obvious control variate is

$$B = \frac{1}{V}\sum_i \epsilon_i - p,$$  \hspace{0.5cm} (A3)$$

namely the difference among the real and the nominal concentrations of magnetic sites. It is clear that the disorder average $\bar{B}$ vanishes. We will employ $B$ to improve the determination of the sample-averaged $\hat{\beta}(\epsilon)$. Note that, although the value of $B$ does not depend on the considered energy (it is fixed by the $\{\epsilon_i\}$), its correlation coefficient with $\langle \hat{\beta} \rangle_{\epsilon}$ needs to be computed for all energies in the $e$-grid.

$B$ is extremely effective as a control variate for the computation of the inverse critical temperature $\beta_c$, as suggested from Fig. 7. The correlation coefficient in that plot is so high, 0.956, that the expected error reduction factor is 3.4. However, the alert reader will note that this is a hasty conclusion. In fact, the $\beta_c$ obtained from $\hat{\beta}(\epsilon)$ is not exactly the average of the inverse critical temperatures found for each sample. The reason for this non-linearity in the Maxwell rule, see Eq. (11), is that the energies $\epsilon^{d,o}$ are not the same for $\beta(\epsilon)$ and for the $\langle \hat{\beta} \rangle_{\epsilon}$ in a given sample. Yet, the dependency on $\epsilon^{d,o}$ of the integral in Eq. (11) is extremely weak [recall the stationarity condition with respect to $\epsilon$ in Eq. (10)].

In fact, the correct computation with $\hat{\beta}(\epsilon)$ does show a significant error reduction, see Fig. 8 close to the factor 3.4 anticipated by the naive analysis in Fig. 7. We note in Fig. 9 an equally significant reduction of the statistical errors for the latent heat. Therefore, our computation of these quantities, obtained with only 500 samples, has been made equivalent to a 5000-samples computation. This is a remarkable reward for such a simple analysis.

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One may fit spin-glass interactions into the same scheme, see Ref. 21 and 42. In the Cardy-Jacobsen mapping, the disorder strength in the Potts model, w, corresponds exactly with τ. Hence, \( y_p = 1/\sqrt{Nd} \). However, keep in mind that this relation was obtained by reasoning in the limit \( Q \gg 1 \).

In a system with periodic boundary conditions, geometrical transitions arise when the energy \( e \) varies from the ordered to the disordered phase. In fact, the system struggles to minimize the surface energy while respecting the global constraint for \( e \). Depending on the fraction of ordered phase, which is fixed by \( e \), the minimizing geometry can be either a bubble, a cylinder or a slab of ordered phase in a disordered matrix (or vice versa). As \( e \) varies, the minimizing geometry changes at definite \( e \) values. This phenomenon is named geometric transition and it is the reason underlying the steps and the flat central region for \( \phi \). In the Cardy-Jacobsen mapping, the disorder strength in the Potts model, w, corresponds exactly with \( \tau \). However, keep in mind that this relation was obtained by reasoning in the limit \( Q \gg 1 \).

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