Renormalization group study of the two-dimensional random transverse-field Ising model

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The infinite disorder fixed point of the random transverse-field Ising model is expected to control the critical behavior of a large class of random quantum and stochastic systems having an order parameter with discrete symmetry. Here we study the model on the square lattice with a very efficient numerical implementation of the strong disorder renormalization group method, which makes us possible to treat finite samples of linear size up to \(L = 2048\). We have calculated sample dependent pseudo-critical points and studied their distribution, which is found to be characterized by the same shift and width exponent: \(\nu = 1.24(2)\). For different types of disorder the infinite disorder fixed point is shown to be characterized by the same set of critical exponents, for which we have obtained improved estimates: \(x = 0.982(15)\) and \(\psi = 0.48(2)\). We have also studied the scaling behavior of the magnetization in the vicinity of the critical point as well as dynamical scaling in the ordered and disordered Griffiths phases.

I. INTRODUCTION

The random transverse-field Ising (RTFI) model is the prototype of random quantum systems having a quantum critical point at zero temperature. This model has experimental realizations and there is a large amount of theoretical work, which aims to clarify the properties of the random critical point. It is expected that basic features of the critical behavior are demonstrated in the onedimensional (1d) model and therefore most of the theoretical studies are performed in 1d. After early works by McCoy and others, \(^{4,5}\) Fisher \(^{6}\) has used a renormalization group (RG) framework to obtain several presumably exact results about its critical properties. It has been shown that the critical properties of the 1d model are governed by a so called infinite disorder fixed point (IDFP), in which the strength of disorder grows without limit during renormalization\(^{8,9}\). As a consequence disorder fluctuations are dominated over quantum fluctuations and the approximations used in the RG approach become exact at the critical point. The IDFP of the RTFI model is shown to govern the critical properties of another random quantum systems having an order parameter with discrete symmetry\(^{8,9}\). Furthermore this fixed point is found to be isomorphic with that of several 1d stochastic models, such as the Sinai walk\(^{10}\), the random contact process\(^{11}\) or the random exclusion process\(^{12}\).

Comparatively less results are available about the critical behavior of the RTFI model in higher dimensions, which are almost exclusively restricted to 2d. By now different numerical studies are in favor of the conclusion, that also in 2d the critical behavior is controlled by an IDFP. In this respect we mention different numerical implementations\(^{13,12}\) of the strong disorder renormalization group (SDRG) method, as well quantum Monte Carlo simulations\(^{13}\). These results are in agreement with recent simulation studies of the 2d random contact process\(^{15}\), which is expected to belong to the same universality class. Also the 2d random walk in a self-affine random potential\(^{16}\) could be related to the 2d RTFI model. As far as the numerical estimates of the critical exponents in 2d are concerned, these contain quite large errors, for a summary of the estimates see Ref.\(^{18}\). In simulation studies these are connected to the logarithmically slow critical relaxation, whereas in the SDRG method the errors has mainly finite-size origin the typical linear size of the largest systems being about \(L = 128 – 160\). Also the type of disorder used in the calculations has an influence on the error of the results. Due to these numerical limitations there are no quantitative results in 3d, although it is very probable that the random critical point is an IDFP in this case, too\(^{13}\).

In this paper we are going to revisit the problem of the critical behavior of the 2d RTFI model. Like other studies we use numerical implementation of the SDRG method, however we have developed a very efficient algorithm, which make us possible to treat systems as large as \(L = 2048\). In this way the number of sites in our samples are several hundred larger, than in previous studies. Comparing with earlier SDRG investigations our study has several different features. i) We define and calculate finite-size pseudo-critical points and study their distribution. ii) We obtain accurate estimates for the true critical point of the model, calculate effective, size-dependent critical exponents and study their extrapolation. iii) We consider different forms of the disorder and study the universality of the critical exponents as well as the scaling functions. iv) We also study scaling outside the critical point, as well as dynamical scaling in the disordered and ordered Griffiths phases.

The structure of the rest of the paper is the following. The model and the SDRG method is presented in Sec. \(\text{II}\). The basic features of the computer algorithm are dis-
cussed in Sec. III. In Sec. IV we describe how finite-size transition points are defined and calculated within the frame of the SDRG method. We study their distribution and analyze the shift of the mean value as well as the width as a function of the size of the system. In Sec. V critical exponents are extracted through finite-size scaling and the scaling behavior outside the critical point is analyzed. We also study dynamical scaling in the disordered and in the ordered Griffith phases. Our results are discussed in the final section, whereas some details of the computer algorithm are presented in the Appendix.

II. MODEL AND THE SDRG METHOD

A. Random transverse-field Ising model

We consider the RTFI model defined by the Hamiltonian:

\[ \mathcal{H} = - \sum_{(i,j)} J_{ij} \sigma_i^x \sigma_j^z - \sum_i h_i \sigma_i^z \]  

(1)

in terms of the Pauli matrices, \( \sigma^x, \sigma^z \). Here \( i \) and \( j \) are sites of a square lattice and the first sum runs over nearest neighbors. The \( J_{ij} \) couplings and the \( h_i \) transverse fields are independent random numbers, which are taken from the distributions, \( p(J) \) and \( q(h) \), respectively.

Here we use two different type of distributions, which have both the same uniform distribution of the couplings:

\[ p(J) = \Theta(J) \Theta(1-J) \]  

(2)

\( \Theta(x) \) being the Heaviside step-function. For the 'box-h' disorder we have:

\[ q(h) = \frac{1}{h_b} \Theta(h) \Theta(h_b - h) \]  

(3)

whereas for the 'fixed-h' model we have a constant transverse field:

\[ q(h) = \delta(h_f - h) \]  

(4)

In the following we use the logarithmic transverse-field \( \theta = \ln h_b \) or \( \theta = \ln h_f \) to characterize the system. In the thermodynamic limit, \( L \to \infty \), the system in Eq. (1) displays a paramagnetic phase, for \( \theta > \theta_c \), and a ferromagnetic phase, for \( \theta < \theta_c \). In between there is a random quantum critical point at \( \theta = \theta_c \). The quantum control parameter is defined as \( \delta = \theta - \theta_c \).

B. Strong disorder renormalization group method

Here we use the SDRG method\(^{22}\), which has been introduced by Ma, Dasgupta and Hu\(^{23}\) and later developed by D. Fisher\(^6\) and others. During the method, the largest local term in the Hamiltonian (which defines the energy-scale, \( \Omega \), at the given RG step) is successively eliminated and at the same time new terms are generated between remaining sites by second-order perturbation method. The procedure is sketched in Fig. 1 for a higher dimensional system. If the largest term is a coupling (see the right panel of Fig. 1), say \( \Omega = J_{ij} \) connecting sites \( i \) and \( j \), then the two sites involved form a spin cluster with an effective moment \( \mu' = \mu_i + \mu_j \) (initially \( \mu_i = 1 \forall i \)) in the presence of an effective transverse field: \( h' \approx h_i h_j / J_{ij} \). The renormalized value of the coupling of the cluster to a site \( a \) is given by the 'maximum rule' \[ \max \{ J_{a,i}, J_{a,j} \} \]. On the other hand, if the largest local term is a transverse-field (see the left panel of Fig. 1), say \( \Omega = h_i \), then site \( i \) is eliminated and effective couplings are generated between each pairs of spins, among the neighbors of \( i \). If \( a \) and \( b \) are neighboring spins to \( i \), then the generated coupling is given by: \( J'_{a,b} \approx J_{a,i} J_{b,i} / h_i \). If the sites \( a \) and \( b \) are already connected by a coupling, \( J_{a,b} \neq 0 \), than the renormalized coupling is given by the 'maximum rule' as \( \max \{ J_{a,b}, J'_{a,b} \} \). The use of the maximum rule is justified if the renormalized couplings have a very broad distribution, which is indeed the case at the IDFP. We shall see that with the maximum rule the numerical algorithm can be simplified. At each step of the renormalization one site is eliminated and the energy scale is continuously lowered. For a finite system the renormalization is stopped at the last site, where we keep the energy-scale and the total moment as well as the structure of the clusters.

C. Scaling at the infinite disorder fixed point

At the IDFP the distribution of the effective couplings and that of the transverse fields becomes broader and broader during the renormalization\(^{7,22}\). Considering the ratio of two effective terms in the Hamiltonian at a given stage of the SDRG it will tend either to infinity or to zero. This indicates that the disorder is infinitely strong.
and the perturbative results during the RG are exact. Qualitatively the log-energy scale, $\ln \Omega$, scales with the linear size of the system size, $L$:

$$\ln (\Omega_0/\Omega) \sim L^\phi,$$

(5)

where $\Omega_0$ denotes a reference energy scale. The average spin-spin correlation function is defined as $G(r) = \langle \sigma_i^x \sigma_j^x \rangle / \langle \sigma_i^x \sigma_i^x \rangle$, where $\langle \ldots \rangle$ denotes the ground-state average and $\langle \ldots \rangle$ stands for the averaging over quenched disorder. The asymptotic value of the correlation function defines the magnetization, $m$, in the system:

$$\lim_{r \to \infty} G(r) = m^2,$$

(6)

and $m > 0$ in the ferromagnetic phase and $m = 0$ in the paramagnetic phase. The connected correlation function, $\tilde{G}(r) = G(r) - m^2$, in the vicinity of the critical point behaves as:

$$\tilde{G}(r) \sim r^{-2x} \exp(-r/\xi),$$

(7)

where the correlation length, $\xi$, is divergent at the critical point as:

$$\xi \sim |\delta|^{-\nu}.$$  

(8)

Thus at $\delta = 0$ there is a power-law decay of the correlations, which is related to the fractal structure of the spin clusters. Indeed, the average cluster moment, $\mu$, is related to the energy-scale, $\Omega$ as:

$$\mu \sim [\ln(\Omega_0/\Omega)]^{\phi},$$

(9)

and can be expressed also with the size:

$$\mu \sim L^{df}.$$  

(10)

Here the fractal dimension of the cluster, $df$, is related to the other exponents as:

$$df = \phi \psi = d - x.$$  

(11)

In 1d the critical exponents are exactly known:

$$\psi = \frac{1}{2}, \quad \phi = \frac{\sqrt{5} + 1}{2}, \quad \nu = 2, \quad x = \frac{3 - \sqrt{5}}{4}.$$  

(12)

III. IMPLEMENTATION OF THE SDRG METHOD

A. Problems with the naïve implementation

The SDRG decimation rules are very simple and it is straightforward to implement the method numerically. In higher dimensions, however, the topology of the lattice is changing during renormalization, which could result in considerable increase of the computational time. More dangerous steps in this respect are the $h$-decimations, during which numerous new bonds are generated and as a result sites with large number of links are formed. In this way, after a naïve implementation of the method, the system is transformed into an almost complete graph and the subsequent $h$-decimations, generating new links between practically all pair of sites, will be very slow. For a system with $N$ sites this algorithm would need $O(N^3)$ time.

Using the maximum rule in the approach, however, offers two ways to speed up the procedure. First, one can notice that the renormalization trajectory is not unique in this case. There are terms in the Hamiltonian, which are called as “local maxima” and which can be decimated independently. Thus one should not follow the “decimation of the largest term in each step” principle, instead we are going to optimize the time of the renormalization trajectory, which goes over in some order of the local maxima. The second consequence of the maximum rule is, that a large number of bonds will never be participating in the renormalization process. These latent bonds can be deleted from the list of edges without consequences. The latent bonds are in such a local environment, that after decimating a nearby site or bond a stronger new coupling is generated to the same edge, thus the original bond disappears without participating in the renormalization. Filtering out these irrelevant bonds will result in a considerable improvement of the algorithm. In the following we discuss the properties of the local maxima and the optimal RG trajectory as well as the main features of the filtering process.

B. Local maxima and the optimal RG trajectory

A local maximum in the set of couplings and transverse fields is such a term, which is larger (not smaller) than any of its neighboring terms. Considering a coupling $J_{ij}$ is a local maximum, provided $J_{ij} \geq h_i$, $J_{ij} \geq h_j$, and $J_{ij} \geq J_{ik}$, $\forall k$, as well as $J_{ij} \geq J_{kj}$, $\forall l$. Similarly a transverse field, $h_i$, is a local maximum, if $h_i \geq J_{ij}$, $\forall j$. It can be shown that the local maxima can be decimated independently, the renormalization performed in any sequence gives the same final result.

C. Filtering out irrelevant bonds

The principle is, that some bonds are irrelevant and does not modify the renormalization procedure has been first realized by Kawashima. He has also suggested a criterion to identify the irrelevant bonds which are then deleted from the graph. This filtering procedure as illustrated in Fig. 2 is used in a few $2d$ numerical works. In Appendix A we give the proper definition of the filtering criterion and prove it.

The use of the filtering criterion for a site with $k$ neighbors needs typically $O(k^3)$ time, since $O(k^2)$ triangles have to be checked. However, the application of the fil-
tering makes not too much reduction of the computational time, if the bond is in a general position, where not too many new terms are generated after consecutive RG steps. There are, however, bonds in “dangerous positions”, having at least one of the neighboring transverse fields as a local maximum. Decimating this local maximum many new terms are generated and therefore the use of prefiltering, which checks the bonds of the decimated spin just before its decimation is very effective, which typically needs only $O(k^2)$ time.

D. Basic steps in the numerical SDRG method

In order to obtain an efficient implementation of the SDRG method we combine the optimal selection of the RG trajectory with the filtering algorithm. In our method we use the following steps.

1. We check all terms of the Hamiltonian and select the local maxima. We make two lists, one for the couplings and one for the transverse fields.

2. We decimate every coupling, which is on the list of local maxima. In this step the order of the decimation is arbitrary. After decimating the original couplings new terms are generated, among which there are new local maxima. We include those into the list of local maxima and at the same time we decimate the new couplings from this list. We repeat this step, until the list of local maximum couplings is empty.

3. If all or all but one the transverse fields are local maxima we select the smallest one, delete all the rest sites and END the iteration. Otherwise we select a transverse field from the list of local maxima having a small (or the smallest) degree, which is defined as the number of edges to the given site and filter for the neighboring bonds. We decimate this transverse field and check the generated new terms for local maxima. If the list of local maximum couplings is not empty, we go to step 2. Otherwise we go to step 3.

Using the selection rule in the second part of step 3. will ensure that the average degree of the sites will not be large. In this way we prevent the formation of too connected clusters, the decimation of which being time consuming. In step 3. making the filtering before decimation (prefiltering) will ensure that the dangerous bonds are deleted.

This algorithm in 2d works typically in $O(L^3 \ln L)$ time in a $L \times L$ system near the critical point. With our method an $L = 128$ sample is renormalized in $\sim 1$ second, whereas for $L = 1024$ the typical time is $\sim 1.5$ minutes (in a 2.4GHz processor).

IV. FINITE-SIZE CRITICAL POINTS

A. Scaling of pseudo-critical points

In the study of the critical behavior of random systems it is very important to find an accurate estimate of the location of the critical point. The quality of the estimate of $\theta_c$ will influence the error of the calculated critical exponents and scaling functions. In a random sample of linear size, $L$, one can generally define finite-size pseudo-critical points, $\theta_c(L)$, which are usually given as the position of the maximum of some physical quantity, which is divergent in the thermodynamic limit (c.f. susceptibility) at $\theta_c$. The distribution of $\theta_c(L)$ provides important information about the scaling behavior at the fixed point of the system. In particular one studies the shift of the average value, $\bar{\theta}_c(L)$, which is expected to scale as:

$$|\theta_c - \bar{\theta}_c(L)| \sim L^{-1/\nu_s},$$

with the shift exponent, $\nu_s$. Similarly, one measures the width of the distribution, $\Delta \theta_c(L)$, which behaves for large-$L$ as:

$$\Delta \theta_c(L) \sim L^{-1/\nu_w},$$

where $\nu_w$ denotes the width exponent. According to renormalization group theory for a classical random system with relevant disorder the critical behavior is controlled by a conventional random fixed point, with $\nu_s = \nu_w = \nu$, where $\nu \geq 2/3$ is the correlation length critical exponent of the system.

B. Identification of pseudo-critical points

In a random quantum system one has to use another methods to locate pseudo-critical points. One method, which is well suited to the SDRG approach is the doubling method, which has been used for chains and for ladders of the RTFI model. In the doubling procedure in 1d (or quasi-1d) geometry one considers a random sample $(\alpha)$ of length, $L$, and makes a duplicated sample $(2\alpha)$ of length $2L$ by joining two copies of $(\alpha)$. Using the SDRG method one calculates some physical quantity (magnetization or gap) in the original and in the replicated sample, which is denoted by $f(\alpha, L)$ and $f(2\alpha, 2L)$, respectively, and study their ratio, $r(\alpha, L) = f(2\alpha, 2L)/f(\alpha, L)$, as a function of the

FIG. 2: Illustration of the filtering criterion.
control parameter, $\theta$. At $\theta = \theta_c(\alpha, L)$ this ratio has a sudden jump, which is identified with the pseudo-critical point of the sample. We note that the actual value of $\theta_c(\alpha, L)$ is practically independent of the physical quantity we considered, since this singularity is connected to the topology of clusters produced by the SDRG approach.

Having this observation in mind we can generalize the doubling method for two (and higher) dimensions. In $2d$ we glue together two identical square-shaped samples at the boundaries as indicated in Fig. 3. Then we renormalize the duplicated sample and calculate the structure of the connected clusters, among which there might be such, which have sites (in equivalent positions) in both replicas. These sites are correlated and the fraction of these correlated sites defines the correlation function between the replicas. By increasing the control parameter, $\theta$, the replica correlation function is decreasing and at a well defined value, $\theta_c(L)$, it suddenly jumps to zero. We consider $\theta_c(L)$ as the pseudo-critical point of the given sample. It is easy to see that this definition is equivalent to the previously used criterion in $1d$, furthermore it is straightforward to generalize it to three- and higher dimensions.

![Fig. 3: Illustration of the boundary conditions used in the doubling procedure in higher dimensions.](image)

C. Numerical results

1. Distribution of pseudo-critical points

We have calculated pseudo-critical points of square-shaped samples for various linear sizes, $L$, which are expressed as $L = 2^l$ and $L = 3 \times 2^{l-1}$ up to $l = 10$. Generally we have considered $4 \times 10^4$ realizations except for the largest system, when we had at least $10^3$ samples. The distribution of the pseudo-critical points is shown in Fig. 4 for both type of disorder. The mean value of the critical points is considerably larger for box-$h$ randomness and also the width of the distribution - for the same value of $L$ - is larger in this case. (We note that the same trend is seen in $1d$, where $\theta_c^{(b)} = 0$ and $\theta_c^{(f)} = 1$.) Taking into account the result in Eq. (13) the appropriate scaling combination is $y = (\theta_c(L) - \theta_c) L^{1/\nu}$ in terms of which the scaled distributions, $\bar{p}(y)$, are shown in the insets of Fig. 4 Here using our final estimates in Eqs. (22) and (21) we obtain excellent scaling collapse of the data points for both type of randomness. The scaling curves for the two different disorder approach the same standardized master curve, which indicates that the fixed point of the RG transformation is unique and (at least for strong enough disorder) strongly attractive. The master curve is different from that in $1d$, which is Gaussian in this case. In $2d$ the maximum of the curve is shifted to negative values and the distribution is non-symmetric. We have calculated the percolation (or spanning) probability, $P_{pr}$, at the critical point, which is given by the fraction of samples having finite replica correlation function at $\theta_c$. It can be expressed with the scaled distribution function as $P_{pr} = \int_0^{\infty} \bar{p}(y) dy$. Our estimate is:

$$P_{pr} = 0.149(2),$$

which is much smaller, than for standard $2d$ percolation. We have also calculated the skewness, $s$, of the distribution, which has the value $s = 0.19(3)$ for both type of disorder. The asymmetric form of the distribution indicates that in $2d$ the topology of the renormalized model is different from that in $1d$. Samples having more strongly connected clusters, thus a higher $\theta_c(L)$, have a somewhat larger weight than the less strongly connected clusters.

2. Shift of the finite-size critical points

For a fixed linear size, $L$, we have calculated the mean value, $\bar{\theta}_c(L)$, as well as the median $\bar{\theta}_c^{med}(L)$ of the pseudo-critical points. Since the distribution is non-symmetric these two characteristic values are not identical, however both are expected to follow the scaling form in Eq. (13urdy with the same value of the shift exponent, $\nu_s$. This is illustrated in Fig. 5, in which $\theta_c - \bar{\theta}_c(L)$ as well as $\theta_c - \bar{\theta}_c^{med}(L)$ is shown as a function of $L$ in a log-log scale for both type of disorder. Having appropriate limiting values for $\theta_c$, see Eq. (22), the points in Fig. 5 are asymptotically very well on straight lines (both for the mean value and for the median) having approximately the same slopes: $-1/\nu_s \approx -0.8$.

To get more quantitative estimates we have calculated effective, size dependent shift exponents which are defined as:

$$\frac{1}{\nu_s(L)} = -\frac{1}{\ln 2} \ln \left[ \frac{\bar{\theta}_c(2L) - \bar{\theta}_c(L)}{\bar{\theta}_c(L) - \bar{\theta}_c(L/2)} \right],$$

and similarly for the median values. These are shown in the two insets of Fig. 5 as a function of $1/L$ (upper inset for the mean) and $\ln L$ (lower inset for the median), respectively. The exponents calculated from the mean values show $1/L$ correction terms for both type of disorder, although with different signs. The extrapolated values are $1/\nu_s^{(b)} = 0.79(2)$ and $1/\nu_s^{(f)} = 0.81(2)$, which agree within the error of the calculation and leads the estimate: $1/\nu_s = 0.80(2)$. As seen in the lower inset of
for ‘f’), as well as from the median (× for ‘b’ and □ for ‘f’) of the distribution, as a function of the linear size, \( L \), in log-log scale. In agreement with the scaling prediction in Eq. (14) the points in Fig. 5 are asymptotically on straight lines, the slope of which is approximately the same for both types of disorder and can be well fitted as \(-1/\nu_w \approx -0.8\). As for the shift exponent in Sec. IV C 2 we have measured effective, size-dependent critical exponents, which are defined as:

\[
\nu_w = \sinh^{-1}\left[ -\frac{\Delta \theta_c(2L) - \Delta \theta_c(L/2)}{2\Delta \theta_c(L)} \right] \frac{1}{\ln 2},
\]

and plotted in the inset of Fig. 6 as a function of \( 1/L \). Extrapolating the effective exponents yields \( 1/\nu_w^{(b)} = 0.805(10) \) and \( 1/\nu_w^{(f)} = 0.811(10) \) for the box-h and the fixed-h randomness, respectively. These values indeed agree within the error of the calculation, thus we can conclude that \( 1/\nu_w = 0.808(10) \) and thus

\[
\nu_w = 1.24(2).
\]

Comparing our estimates for the shift-exponent in Eq. (17) with that of the width-exponent in Eq. (19) we notice that they agree within the error of the method, which corresponds to the renormalization group result for a classical conventional random critical point.\( ^{25} \) In order to make a direct check of the equivalence of the two exponents we have formed the ratio:

\[
\alpha(L) = \frac{\theta_c - \overline{\theta}_c(L)}{\Delta \theta_c(L)},
\]

which should approach an \( L \)-independent constant value at the ‘true’ critical point, \( \theta_c \), provided \( \nu_s = \nu_w \). In Fig. 7 we have plotted the \( \alpha(L) \) ratios as a function of

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FIG. 4: (Color online) Distribution of the pseudo-critical points, \( \theta_c(L) \), for various lengths for box-h randomness (upper panel) and for fixed-h randomness (lower panel). In the insets the scaled distributions are shown as a function of \( y = (\theta_c(L) - \overline{\theta}_c)L^{1/\nu_s} \), see the text.

FIG. 5: (Color online) Scaling of the shift of the finite-size transition points, calculated from the mean value (× for ‘b’ and * for ‘f’), as well as from the median (× for ‘b’ and □ for ‘f’) of the distribution, as a function of \( L \) in log-log scale for both type of disorder. Estimates for \( \theta_c \) are taken from Eq. (12) and the straight lines indicating the asymptotic behavior have the same slope: \(-1/\nu_s \approx -0.8\). The effective critical exponents calculated from Eq. (16) are shown in the upper inset (for the mean) and in the lower inset (for the median).

3. Scaling of the width of the pseudo-critical points

We have measured the standard deviation of the distribution of the pseudo-critical points, \( \Delta \theta_c(L) \), which are shown in Fig. 4 as a function of the linear size, \( L \), in a log-log scale. In agreement with the scaling prediction in Eq. (14) the points in Fig 4 are asymptotically on straight lines, the slope of which is approximately the
V. SCALING AT THE CRITICAL POINT

Having accurate estimates of the critical points for both types of randomness we are ready to study the critical behavior of the system. In this respect we have concentrated our effort at the critical point, where we have studied the distribution function of the magnetization, as well as that of the (log-)gaps and calculated critical exponents by finite-size scaling. At the critical point we have considered finite systems up to a linear length $L = 2048$ and studied $4 \times 10^4$ realizations for each sizes.

These investigations are supplemented with numerical studies outside the critical point, both in the disordered and in the ordered phases. For the magnetization we have studied the scaling regime, defined as $\delta L^{1/\nu} = O(1)$. To obtain information about the dynamics of the system we have studied the scaling behavior of the excitation energies both in the ordered and in the disordered Griffiths phases and in this investigation we have not restricted ourselves to the vicinity of the critical point. In the off-critical region we studied random samples with fixed-$h$ randomness up to a linear length $L = 512$ and for $10^4$ realizations.

A. Magnetization

1. Spin clusters

During renormalization effective spin clusters are formed, which are than decimated at different energy scales. We illustrate the cluster structure of a given sample at the critical point in the insets of Fig. 8 in which clusters having the same size are marked with the same greyscale (color). Note that most of the clusters consist of only one site and the large clusters are generally geometrical.
The magnetic properties of a given finite sample are related to the magnetic moment of some effective spin cluster, which appears at the last stages of the RG procedure. In principle one can define different types of such spin clusters. i) The magnetization cluster has the smallest effective transverse field, thus decimated at the lowest energy scale. The moment of magnetization clusters are denoted by $\tilde{\mu}$. By definition the smallest possible value of the magnetization cluster is $\tilde{\mu}_{\min} = 1$, thus the corresponding minimal magnetization at the disordered phase is given by $1/L^d$, i.e. it varies as a power-law of $L$. ii) The correlation cluster with a moment $\tilde{\mu}_{\text{corr}}$ is present in the duplicated sample and involved in the replica correlation function. By definition a correlation cluster exists only for $\theta \leq \theta_c(L)$, i.e. below the pseudo-critical point of the given sample. We have checked, that for large $L$ if a correlation cluster exists, it is almost always the magnetization cluster. iii) Finally we define also an energy cluster in a sample, which is the magnetization cluster for $\theta > \theta_c(L)$, whereas for $\theta \leq \theta_c(L)$ it is the cluster decimated before the correlation cluster. These clusters are involved in the low-temperature or the small longitudinal field properties of the system and will be considered in dynamical scaling in Sec IV.B.

2. Moment of magnetization clusters

We have calculated the distribution functions of the moments of magnetization clusters for different lengths, $R_L(\tilde{\mu})$, which are shown in Fig 9 for both types of randomness. According to scaling theory $R_L(\tilde{\mu}) = L^{d_f} \tilde{R}(\tilde{\mu} L^{-d_f})$ which is illustrated in the insets of Fig 9. Up to a multiplicative constant the scaling functions, $\tilde{R}(\omega)$, are identical for the two different randomness and can be approximated with an exponential function: $\tilde{R}(\omega) \sim \exp(-\omega/\omega^*)$, $\omega^*$ being some randomness dependent value.

3. Fractal dimension and critical exponent

In order to obtain an accurate estimate for the fractal dimension, $d_f$, and for the magnetization exponent, $x$, we have calculated average moments of the magnetization clusters, $\overline{\mu}_L$, which are plotted in the inset of Fig. 10 as a function of $L$ in a log-log scale. For both type of randomness the points are asymptotically on straight lines having the same slope, which is in agreement with the scaling relation in Eq. (10). We have calculated effective, size-dependent fractal dimensions through:

$$d_f(L) = \sinh^{-1} \left( \frac{\overline{\mu}_L - \overline{\mu}_{L/2}}{2 \overline{\mu}_L} \right) \frac{1}{\ln 2},$$

which are plotted in Fig. 10. As seen in this figure the effective fractal dimensions show no systematic trend with $L$ and the $d_f(L)$ values spread around the same mean value for both form of disorder. This mean value is taken to our estimate for the fractal dimension:

$$d_f = 1.018(15)$$

and from Eq. (11) we have for the magnetization exponent:

$$x = 0.982(15).$$

4. Scaling of the magnetization

The magnetization is given by the asymptotic value of the correlation function in Eq. (11), which definition can be extended in a finite system in terms of the replica correlation function as introduced in Sec IV.B. In a given
sample of linear size $L$ two spins at a distance $r \sim L$ are correlated if both are in the same correlation cluster. Consequently (the average value of) the magnetization is given by: $m(\delta, L) = \frac{\sum \mu_{\text{corr}}(L/d)}{L/d}$, where $\mu_{\text{corr}}$ is the average value of the mass of the correlation cluster as defined in Sec. V A 1. The magnetization as a function of $\delta$ is plotted in Fig. 11 for different finite systems. Using scaling theory the magnetization in the vicinity of the critical point is expected to behave as $m(\delta, L) L^{x} \sim (\delta L^{1/\nu})$. To test this assumption in the inset of Fig. 11 we have plotted $m(\delta, L) L^{x}$ as a function of $\delta L^{1/\nu}$. Using the estimates for the critical exponents in Eqs. 21 and 25 an excellent scaling collapse of the data is obtained.

B. Dynamical scaling

Here we study the properties of the low-energy excitations, which are responsible for the dynamical behavior of the system, such as the auto-correlation function or the low-temperature and small-field behavior of the susceptibility, specific heat, magnetization, etc. Such a low-energy collective excitation of a random sample of linear length $L$ in the SDRG method is represented by
a large spin cluster which is formed at the last steps of the renormalization process and its excitation energy, $\epsilon_L$, is measured by the effective transverse field of the cluster. These clusters are the so called energy clusters as defined in Sec. V A 1. According to the scaling relation in Eq. (3) at the critical point it is convenient to use the log-variable: $\gamma_L = -\ln(\epsilon_L)$. In the following we study the distribution of $\gamma_L$ at the critical point, as well as in disordered and ordered Griffiths phases, and investigate its scaling behavior with $L$. As far as dynamical properties are considered the SDRG method gives asymptotically exact results also in the off-critical region, where the relaxation time is divergent. This is not the case for static quantities due to the finite correlation length.

1. Critical point

At the critical point the energy clusters are decimated either at the last step of the renormalization process, which happens if the pseudo-critical point of the sample, $\theta_c(L)$, is smaller than $\theta_c$, or at the last but one step, if $\theta_c(L) > \theta_c$. The distribution of the log-excitation energies at the critical point for different finite sizes are shown in Fig. 12 for the two type of randomness. As seen in this figure the distributions broaden with increasing $L$, which is a clear signal of infinite-disorder scaling. Referring to Eq. (5) we introduce the scaling combination: $\tilde{\gamma} = (\gamma_L - \gamma_0)L^{-\psi}$, in terms of which the distributions collapse to the same curve provided the exponent is $\psi \approx 0.48$. This is illustrated in the insets of Fig. 12. The constant term, $\gamma_0$, used in the fitting process is found to be $O(1)$ and has only a little influence on the value of the exponent $\psi$. The scaling functions in the insets of Fig. 12 have the same form for the two types of applied randomness and have a heavier tail than in 1d. The skewness values at $L = 1024$ are $s = 0.82(1)$ for 2d to be compared with $s = 0.64(1)$ in 1d.

In order to obtain more accurate estimate for the exponent $\psi$ we have considered the mean value, $\overline{\gamma}(L)$, as well as the standard deviation, $\Delta\gamma(L)$, of the distributions. We have formed the differences, $d_1(L)$, which are $d_1(L) = \overline{\gamma}(L) - \overline{\gamma}(L/2)$ for the mean and $d_1(L) = \Delta\gamma(L) - \Delta\gamma(L/2)$ for the width of the distribution, respectively and which are plotted in the inset of Fig. 13 as a function of $L$ in a log-log plot. As seen in this figure the points for the two quantities and for the two-type of randomness are on parallel straight lines the slope of which is compatible with $\psi \approx 0.48$. In the next step we have calculated finite-size effective exponents through the definition:

$$\psi(L) = \frac{1}{\ln 2} \ln \left[ \frac{d_1(2L)}{d_1(L)} \right] , \quad (26)$$

which are plotted in Fig. 13 as a function of $L$. As seen in this figure there is some systematic trend of the points up to $L = 128 - 192$, but for larger $L$-s there are only fluctuations around a mean value what we use as an estimate for the exponent:

$$\psi = 0.48(2) . \quad (27)$$

This limiting value is the same for both type of randomness and describes well the scaling behavior both the mean value and the width of the distribution.

2. Disordered Griffiths phase

In the disordered Griffiths phase the energy clusters, which are related to the low-energy excitations in a large system are decimated at the last step of the renormalization process. The scaling behavior of the low-energy excitations here is different from that at the critical point. We illustrate the scaling behavior of the distribution of the log-excitation energies for different sizes in Fig. 14 for
FIG. 13: (Color online) Effective exponents $\psi(L)$ as a function $\ln(L)$ calculated from the mean or from the width of the distribution of the log-excitation energies for the two type of randomness. Inset: finite differences of the mean and the width of the log-excitation energy distribution as a function of $L$ in a log-log scale, see text. The slope of the dotted straight line is given by $\psi = 0.48$.

the fixed-$h$ randomness at a distance $\delta = 9.64 \times 10^{-3}$ from the critical point. As seen in this figure the shape of the (logarithm of the) distribution functions is very similar for different $L$-s and the curves are merely shifted with $\ln(L)$. This follows from the assumption, that the typical value of the excitation energy in a system of size $L$ scales as a power-law:

$$\epsilon_L \sim L^{-\delta},$$

$\delta$ being the dynamical exponent, which is a continuous function of the control parameter, $\delta > 0$. Consequently the appropriate scaling combination is:

$$\tilde{\gamma} = \gamma_L - \delta \ln(L) - \gamma_0,$$

in terms of which the distribution functions have a scaling collapse, provided the appropriate value of the dynamical exponent is used. This is illustrated in the inset of Fig.13. An estimate for the dynamical exponent from the shift of the distributions, $z_{sh}(L)$, can be obtained from the optimal collapse of the data points for sizes $L/2$ and $L$. In Fig.13 we have $d/z_{sh}(512) = 2.3(2)$. If the low-energy excitations in the system are localized then the scaled distribution function is suggested to be given by the Fréchet distribution known from extreme value statistics, to be written in the form:

$$\ln p(\tilde{\gamma} - \gamma_0) = -\frac{d}{z} \tilde{\gamma} - \exp \left( -\frac{d}{z} \tilde{\gamma} \right) + \ln(d/z).$$

Indeed the scaled distribution function in the inset of Fig.13 is well described by the function in Eq. (33), where only one fitting parameter, $\gamma_0$ in Eq. (29) is used. We note that for large $\tilde{\gamma}$ the tail of $\ln p(\tilde{\gamma})$ is linear and its slope, $-d/z_{sh}(L)$, can be used to obtain an independent estimate for the dynamical exponent. The measured values of $d/z_{sh}(L)$ are given in the caption of Fig.13 and these are compatible with those calculated from the shift of the distributions. There are, however finite-size corrections for $L < \xi$, where the correlation length at the studied value of $\delta$ is of the order of $\xi = O(10^2)$.

FIG. 14: (Color online) Distribution of the log-excitation energies in the disordered Griffiths-phase (at $\delta = 9.64 \times 10^{-3}$) in a log-log scale for different sizes. The slopes of the straight lines indicating the tail of the curves are $d/z = 2.5(2)$, 2.1(2), 2.1(2) and 2.1(2), for $L = 64, 128, 256$ and 512, respectively. In the inset the scaled distributions are shown in terms of the variable in Eq. (30), which is well described by the Fréchet distribution (full line) in Eq. (30).

We have repeated the previous calculation for several values of $\delta$ in the disordered Griffiths phase and calculated estimates for the dynamical exponent both from the shift of the distributions and from the slope of the tail. These estimates obtained at different $L$-s are shown in a log-log plot in Fig. 15. One can notice that the finite-size corrections are stronger for small $\delta$-s, where the correlation length is comparatively larger. According to scaling theory the dynamical exponent for small $\delta$ behaves as

$$\frac{d}{z} \approx c\delta^{\psi_{d/z}}$$

and divergent at $\delta = 0$. We have checked the relation in Eq. (34) and indeed in Fig. 10 one can identify an approximately linear part for $\delta \leq 0.02$ having a slope $\approx 0.6$. This value is compatible with our previous estimates $\psi_{d/z} = 0.60(6)$ using results in Eqs. (31) and (27).

Closing this section we note that the dynamical exponent enters into the singularities of different physical quantities. For example at low-temperature the susceptibility behaves as: $\chi(T) \sim T^{d/z-1}$ and the specific heat has the form $C_v \sim T^{d/z}$. More details about the scaling relations in the Griffiths phase can be found in [22,49].
In the disordered Griffiths lapse, as illustrated in the inset of Fig.16, the distributions are shown in terms of the variable in Eq. (32). In the inset data for the largest system, $L = 512$, compared with the scaling curve in Eq. (31) with $c = 36.5$ in a linear asymptotics from above, on the contrary in the ordered Griffiths phase in the inset of Fig.16 the points bend below a straight line. This is compatible with the scaling result, that asymptotically:

$$\ln p(\tilde{\gamma}) \sim -\tilde{\gamma}^d.$$  

Our data in the inset of Fig.16 are still not in the asymptotic regime but the tail of the distribution clearly decreases faster than linear for the large sizes.

### VI. Discussion

The concept of infinite disorder fixed point has been introduced quite recently and its basic properties have been demonstrated in partially exact calculations in different one-dimensional random quantum systems and stochastic systems. In higher dimensions, in particular in two dimensions the calculations are numerical and have only limited accuracy. In the present paper we have considered the prototypical model in $2d$ having an IDFP the random transverse-field Ising model and studied its critical properties by a numerical implementation of the SDRG approach. As follows from the concept of IDFP our numerical results are expected to be asymptotically exact.

In our approach we have used a very efficient computational algorithm which made us possible to treat samples which are ten-times larger in linear size, compared with previous calculations. In this way we could reduce the effect of finite-size corrections and could also study off-critical properties, such as scaling functions and dynamical scaling in the disordered and ordered Griffiths phases.
The main results of our investigations are the following. We have extended the finite-size scaling study for pseudo-critical points and from their distribution we have obtained precise estimates for the correlation length critical exponent, which has been shown to govern both the shift and the width of the distribution. Using different types of randomness we have demonstrated that the IDFP is universal, the critical exponents as well as the critical scaling functions are independent of the randomness used in the calculation. We have also studied cross-over phenomena with respect of the linear size of the system as well as the type of randomness used in the calculation. The sometimes large errors and deviations between the results of previous numerical studies are presumably due to cross-over effects. This can be seen, e.g. in Fig. 13 where the estimates of the exponent $\psi$ have strong finite-size as well as randomness-type dependence.

As a result of the larger samples and the good statistics of the numerical data we have obtained accurate estimates for the critical exponents and studied - at the first time - the behavior of scaling functions, both at the critical point and in the finite-size scaling limit in the vicinity of the critical point. We have also extended our investigations to the disordered and the ordered Griffiths phases and have checked various predictions of phenomenological scaling theory.

Comparing the critical behavior in 1d to that in 2d we have qualitatively similar results, but there are also important differences. First of all the actual values of the critical exponents as well as the form of the scaling functions are different. Scaling in the ordered Griffiths-phase, which involves powers of $d$ in Eqs. (32) and (33) however, is also qualitatively different. One particular feature of the model in 1d is self-duality, which could be the reason why in 1d the distribution of the pseudo-critical points involves different shift and mean exponents. Our calculations can be extended to several directions. One possibility is to study the entanglement properties of the model, which can be well performed within the frame of the SDRG approach. At present conflicting theoretical predictions are available about the finite-size dependence of the critical entanglement entropy, which could be possibly clarified by using larger systems in the calculation. A second promising direction of application of our approach is to consider higher dimensional systems. At present even in three dimensions only the existence of infinite disorder scaling is demonstrated, but no estimates are known about the critical exponents. In four and higher dimensions no numerical studies have been performed so far. Studies in these directions are in progress.

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Appendix A: Filtering out irrelevant bonds

In the SDRG procedure if the maximum rule is applied several bonds are irrelevant, which means that they can be deleted without modifying the final result of the renormalization.

1. Condition for a bond to be irrelevant

Let us consider a bond with a log-coupling $\kappa_{ij} = \ln J_{ij}$ between sites $i$ and $j$ and consider such nearest neighbor points, one of those is denoted by $k$ in Fig. 17 which has bonds both to $i$ and $j$ of strength $\kappa_{ik}$ and $\kappa_{jk}$, respectively. The triangle $(i,j,k)$ is called the 'majorating triangle' of the $(i,j)$ bond if $\kappa_{ij}$ is the smallest bond in the triangle, and it is also smaller, than the potentially generated new bond: $\kappa'_{ij} = \kappa_{ik} + \kappa_{jk} - \theta_k$, where $\theta_k = \ln h_k$ is the log-transverse field at site $k$. If such a majorating triangle exists then the bond is irrelevant.

2. Proof of the filtering criterion

For the proof it is enough to consider such a decimation step during which the majorating triangle collapses and we follow also the evolution of the parameters in another triangle, $(i,j,l)$, which is not necessarily a majorating one. The maximal log-parameter in the four-site system in Fig. 17 is denoted by $\omega$. We note that during renormalization the log-transverse fields can not increase, thus the change of their values are unimportant in the proof.

a. Bond decimation

The $(i,j)$ bond is not the largest one, since it has a majorating triangle, so that for the position of the largest bond we have three different cases. i) Largest bond in the majorating triangle: $\omega = \kappa_{ik}$ (or $\kappa_{jk}$). The spins $i$ and $k$ fuse into a new effective spin, which is connected to the spin $j$ with a coupling of $\kappa'_{ij} = \max(\kappa_{ij}, \kappa_{jk}) = \kappa_{jk}$, thus, $\kappa_{ij}$ simply disappears. ii) Largest bond in the another triangle: $\omega = \kappa_{jl}$ (or $\kappa_{kl}$). The $j$ and $l$ spins fuse into a new effective spin. If $\kappa_{ij} < \kappa_{jl}$, than the value of $\kappa_{ij}$ simply disappears, otherwise its value does not change. Similarly, the value of $\kappa_{kl}$ does not change, while $\kappa_{jk}$ is replaced by $\kappa'_{jk} = \max(\kappa_{jk}, \kappa_{kl}) \geq \kappa_{jk}$. Taken all round, the new $(i,j,k)$ triangle is a majorating triangle of the $(i,j)$ bond. iii) Largest bond between the two triangles: $\omega = \kappa_{kl}$. The spins $k$ and $l$ fuse and there is one triangle
left. Here the value of $\kappa_{ij}$ does not change and the new triangle is a majorating one of the $(i,j)$ bond.

b. Site decimation

We should consider two different cases: i) $\omega = \theta_k$: $\kappa'_{ij} = \kappa_{ik} + \kappa_{jk} - \theta_k > \kappa_{ij}$ due to our assumptions, thus $\kappa_{ij}$ disappears without affecting the results. ii) $\omega = \theta_i$ (or $\theta_j$ analogously): $\kappa'_{jk} = \kappa_{ij} + \kappa_{ik} - \theta_i \leq \kappa_{ij} < \kappa_{jk}$, thus this coupling is unaffected by the $\kappa_{ij}$ coupling. However, in principle the other newly generated bonds are not irrelevant, e.g. $\kappa'_{jl} < \kappa_{jl}$ is not always fulfilled. Due to this, the effect of the majorated $(i,j)$ edge will not certainly disappear after one single decimation step, in which it is involved. Moreover, a lot of new couplings can be generated occasionally, if the bond lies in a ’dangerous position’.

c. Completing the proof

Here we show, that the newly generated coupling in the $(i,j,k)$ triangle, $\kappa'_{ij}$, is always majorated by this triangle. The two neighboring couplings of $\kappa'_{ij}$ are $\kappa_{jk}$ (due to the fact, that $\kappa'_{jk} < \kappa_{jk}$), and $\max(\kappa_{kl}, \kappa'_{kl})$, where $\kappa'_{kl} = \kappa_{ik} + \kappa_{il} - \theta_i$. $\kappa'_{jl}$ is smaller than these neighboring couplings, namely $\kappa'_{jl} < \kappa_{ij} < \kappa_{jk}$ and $\kappa'_{jl} - \kappa'_{ij} = \kappa_{ik} - \kappa_{ij} > 0$. From the latter follows, that $\kappa'_{jl} < \kappa'_{kl}$, which can not be greater, than $\max(\kappa_{kl}, \kappa'_{kl})$ corresponding to the new value of the coupling between the $k$ and $l$ spins. Now we see, that the $\kappa'_{ij}$ coupling is majorated by the $(i,j,k)$ triangle, if $\kappa'_{ij}$ is smaller, than the generated coupling $\kappa''_{ij} \equiv \max(\kappa_{kl}, \kappa'_{kl})+\kappa_{ik} - \theta_k$, which is obtained by decimating the transverse field at $k$. Let us consider their difference: $\kappa''_{ij} - \kappa'_{ij} \geq \kappa_{kl} + \kappa_{ik} - \theta_k - \kappa_{ij} + \theta_l = \kappa_{ik} + \kappa_{ij} - \theta_k - \kappa_{ij} = \kappa'_{ij} - \kappa_{ij} > 0$. Thus the $\kappa'_{ij}$ coupling is always majorated by the $(i,j,k)$ triangle.

Let us summarize our findings. If the bond $(i,j)$ has a majorating triangle, it can not be decimated directly. Decimating in its neighborhood either this bond disappears or new couplings are generated, the value of which involves $\kappa_{ij}$. These new couplings, however, are always majorated by a triangle, thus they are never decimated during the renormalization process. Consequently the value of $\kappa_{ij}$ does not influence the result of the renormalization.

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1. Electronic address: ikovacs@szfki.hu
2. Electronic address: igloi@szfki.hu
3. For reviews, see: H. Rieger and A. P. Young, in Complex Behavior of Glassy Systems, ed. M. Rubi and C. Perez-Vicente, Lecture Notes in Physics 492, p. 256, Springer-Verlag, Heidelberg, 1997; R. N. Bhatt, in Spin glasses and random fields A. P. Young Ed., World Scientific (Singapore, 1998).
4. S. Sachdev, Quantum Phase Transitions (Cambridge University Press, 1999)
5. D.H. Reich, B. Eliman, J. Yang, T. F. Rosenbaum, G. Aeppli and D. P. Belanger, Phys. Rev. B42, 4631 (1990); W. Wu, B. Eliman, T.F. Rosenbaum, G. Aeppli and D.H. Reich, Phys. Rev. Lett. 67, 2076 (1991); W. Wu, D. Bitko, T.H. Rosenbaum and G. Aeppli, Phys. Rev. Lett. 71, 1919 (1993); J. Brooke, D. Bitko, T.F. Rosenbaum and G. Aeppli, Science 284, 779 (1999).
6. B. M. McCoy and T. T. Wu, Phys. Rev. 176, 631 (1968); Phys. Rev. 188, 982 (1960); B. M. McCoy, Phys. Rev. 188, 1014 (1969); Phys. Rev. B 2, 2795 (1970).
7. D. S. Fisher, Phys. Rev. Lett. 69, 534 (1992); Phys. Rev. B 51, 6411 (1995).
8. D. S. Fisher, Physica A 263, 222 (1999)
9. T. Senthil and S. N. Majumdar Phys. Rev. Lett. 76, 3001 (1996)
10. E. Carlon, P. Lajkó, and F. Iglói, Phys. Rev. Lett. 87, 277201 (2001)
11. D. S. Fisher, Le Doussal and C. Monthus, Phys. Rev. E 59 (1999) 4795
12. J. Hooyberghs, F. Iglói and C. Vanderzande, Phys. Rev. Lett. 90 100601, (2003); Phys. Rev. E 69, 066140 (2004).
13. R. Juhász, L. Santen and F. Iglói, Phys. Rev. Lett. 94, 010601 (2005); Phys. Rev. E 72, 046129 (2005); Phys. Rev. E 74, 061101 (2006).
14. O. Motrunich, S.-C. Mau, D.A. Huse and D.S. Fisher, Phys. Rev. B 61, 1160 (2000).
15. Y.-C. Lin, N. Kawashima, F. Iglói and H. Rieger, Progress in Theor. Phys. 138, (Suppl.) 479 (2000).
Iglói, Eur. Phys. J. B 20 267 (2001).

Y.-C. Lin, F. Iglói and H. Rieger, Phys. Rev. Lett. 99, 147202 (2007).

R. Yu, H. Saleur and S. Haas, Phys. Rev. B 77, 140402 (2008).

I. A. Kovács and F. Iglói, Phys. Rev. B 80, 214416 (2009).

C. Pich, A.P. Young, H. Rieger and N. Kawashima, Phys. Rev. Lett. 81, 5916 (1998).

R. Yu, H. Saleur and S. Haas, Phys. Rev. B 77, 140402 (2008).

I. A. Kovács and F. Iglói, Phys. Rev. B 80, 214416 (2009).

C. Monthus and T. Garel, arXiv:1003.1627.

For a review, see: F. Iglói and C. Monthus, Physics Reports 412, 277, (2005).

S.K. Ma, C. Dasgupta and C.-K. Hu, Phys. Rev. Lett. 43, 1434 (1979); C. Dasgupta and S.K. Ma, Phys. Rev. B 22, 1305 (1980).

There is a third advantage of the maximum rule, which follows from the fact that in this case the average couplings are weaker than for the summing rule. As a consequence the critical point of the system is considerably lower than for the summing rule and therefore less h-decimations have to be performed in the vicinity of the critical point, which speeds up the calculation.

N. Kawashima, unpublished.

F. Pázmándi, R.T. Scalettar, and G.T. Zimányi, Phys. Rev. Lett. 81, 5130 (1998).

S. Wiseman and E. Domany, Phys. Rev. Lett. 81 (1998) 2938.

A. Aharony, A.B. Harris and S. Wiseman, Phys. Rev. Lett. 81 (1998) 252.

K. Bernardet, F. Pazmandi and G. Batrouni Phys. Rev. Lett. 84 4477 (2000).

M. T. Mercaldo, J-Ch. Anglès d’Auriac, and F. Iglói Phys. Rev. E 69, 056112 (2004).

C. Monthus and T. Garel, Eur. Phys. J. B 48, 393-403 (2005).

A. B. Harris, J. Phys. C 7 , 1671 (1974)

J. T. Chayes et al., Phys. Rev. Lett. 57, 299 (1986).