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Ising-like transitions in the O(n) loop model on the square lattice

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We explore the phase diagram of the O(n) loop model on the square lattice in the (x,n) plane, where x is the weight of a lattice edge covered by a loop. These results are based on transfer-matrix calculations and finite-size scaling. We express the correlation length associated with the staggered loop density in the transfer-matrix eigenvalues. The finite-size data for this correlation length, combined with the scaling formula, reveal the location of critical lines in the diagram. For $n \gg 2$ we find Ising-like phase transitions associated with the onset of a checkerboardlike ordering of the elementary loops, i.e., the smallest possible loops, with the size of an elementary face, which cover precisely one-half of the faces of the square lattice at the maximum loop density. In this respect, the ordered state resembles that of the hard-square lattice gas with nearest-neighbor exclusion, and the finiteness of n represents a softening of its particle-particle potentials. We also determine critical points in the range $-2 \le n \le 2$. It is found that the topology of the phase diagram depends on the set of allowed vertices of the loop model. Depending on the choice of this set, the n > 2 transition may continue into the dense phase of the $n \le 2$ loop model, or continue as a line of $n \le 2$ O(n) multicritical points.

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I. INTRODUCTION

The O(n) loop model is a highly useful tool for the analysis of O(n) symmetric *n*-component spin models [1,2], and also for that of polymers [3-5]. A number of such loop models in two dimensions is exactly solvable [6–15]. The present work investigates the nonintersecting loop model described by the partition sum

$$Z_{\text{loop}} = \sum_{\text{all } \mathcal{G}} x^{N_x} y^{N_y} z^{N_z} n^{N_l}, \tag{1}$$

where \mathcal{G} is a graph consisting of any number of N_l closed, nonintersecting loops. Each lattice edge may be covered by at most one loop segment, and there can be 0, 2, or 4 incoming loop segments at a vertex. In the latter case, they can be connected in two different ways without having intersections. The allowed four kinds of vertex configurations are shown in Fig. 1, together with their weights denoted x, y, and z. The numbers of vertices with these weights are denoted N_x , N_y , N_7 respectively.

This loop model is equivalent with an O(n) spin model, as described by Ref. [16]. The n-component spins are sitting in the middle of the edges of the square lattice. The Boltzmann weight for each spin configuration is the product over all vertices of the lattice of the local weights w,

$$w = 1 + x(\vec{s}_1 \cdot \vec{s}_2 + \vec{s}_2 \cdot \vec{s}_3 + \vec{s}_3 \cdot \vec{s}_4 + \vec{s}_4 \cdot \vec{s}_1) + y(\vec{s}_1 \cdot \vec{s}_3 + \vec{s}_2 \cdot \vec{s}_4) + z[(\vec{s}_1 \cdot \vec{s}_2)(\vec{s}_3 \cdot \vec{s}_4) + (\vec{s}_2 \cdot \vec{s}_3)(\vec{s}_4 \cdot \vec{s}_1)],$$
(2)

where the spins \vec{s}_1 to \vec{s}_4 sit on the four edges incident to the vertex and are labeled anticlockwise. The spins are subject to a measure and normalization

$$\int d\vec{s}_k = 1, \quad \int d\vec{s}_k (\vec{s}_k \cdot \vec{s}_k) = n. \tag{3}$$

Expansion of the partition integral in powers of the coupling constants x, y, and z turns the spin model into the loop model of Eq. (1).

Although the spin dimensionality n can assume only integer values n = 1, 2, ... in the original O(n) spin model, n can also have noninteger and even negative values in the loop model of Eq. (1), while the partition sum remains well defined. Whereas the Boltzmann weight of Eq. (2) can become negative when x, y, and/or \sqrt{z} exceeds values of order 1/n, the Boltzmann weights of the loop model remain physical for all non-negative values of x, y, z, and n.

The parameter space of Eq. (1) contains several exactly solved "branches" [8,9]. These solutions appear as special lines parametrized by n in the (x, y, z, n) parameter space. They have shown the existence of a richness of "nonuniversal" behavior as a function of n in the range n < 2. Branches 1, 2, 3, and 4 are defined by the weights

$$z = \{2 - [1 - 2\sin\alpha][1 + 2\sin\alpha]^2\}^{-1},\tag{4}$$

$$x = \pm 4z \sin \alpha \cos(\pi/4 - \alpha/2), \tag{5}$$

$$y = \pm z(1 + 2\sin\alpha),\tag{6}$$

$$n = -2\cos(4\alpha), \quad 2 - k \le 4\alpha/\pi \le 3 - k, \quad k = 1,2,3,4,$$
(7)

where k is the branch number. Branch 1 describes, in the spin language, the O(n) critical point separating the paramagnetic phase from the low-temperature O(n) phase. In the language of the loop model, this critical point separates the dilute loop phase from the dense phase at larger x, y, z. The size of the largest loops diverges at the O(n) critical point.

Analytic continuation of branch 1 into the range $\alpha < \pi/4$ leads to branch 2, which describes the universal properties

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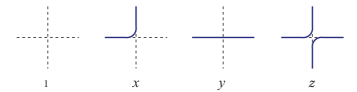


FIG. 1. (Color online) The four kinds of vertices of the O(n) loop model on the square lattice, together with their weights. Rotated versions have the same weights. The present work is restricted to two subspaces of (x, y, z), namely y = x, $z = x^2$ and y = 0, $z = x^2$.

of the dense phase of nonintersecting loop models in two dimensions. Indeed, the universal properties of branches 1 and 2 match with similar branches of the O(n) loop model on the honeycomb [6] and on the triangular [10] lattices. For general n, the dense O(n) phase is still critical in the sense that the correlation functions decay algebraically. It is noteworthy that both branch 1 and branch 2 represent points in the parameter space where the leading temperature field vanishes. It is irrelevant on branch 2, but its absence greatly suppresses corrections to scaling, which enables accurate numerical analyses [16].

To explain the physics of branches 3 and 4, it is helpful to introduce auxiliary dual Ising variables on the faces of the square lattice. Two neighboring Ising spins receive the same sign if and only if they are separated by a loop segment. Then, dense loop configurations dominated by z-type vertices will display ferromagnetic Ising order, while more dilute loop configurations may be Ising disordered. Since it has been found that the universal properties of branch 4 display a combination of Ising critical exponents and dense O(n) exponents, branch 4 has been identified as a locus of Ising transitions in the dense O(n) phase [9,16]. The phase diagram conjectured in Ref. [16] predicts that, for constant x/y and n, these Ising transitions form a critical line in the (x,z) diagram. At its small-x end, this line is believed to merge with the line of O(n) critical transitions, in a multicritical point with the universal behavior represented by branch 3.

In addition, there is a fifth exactly solvable branch, called branch 0, with weights x = z = 1/2, y = 0. Its universal properties are different from branches 1 to 4. The vanishing of y imposes Ising order on the dual Ising spins along each loop, but does not determine the sign of the spins along other loops. For this reason the y = 0 plane represents a special locus of partial Ising order in the parameter space. Branch 0 has been interpreted [16] as the y = 0 special point on the line of higher critical points represented by branch 3 in the (x, y, z) space.

Whereas these exact solutions have yielded a considerable insight in O(n)-type universality classes, they cover only a small part of the parameter space, so that further exploration is desirable. In particular, the questions arise of what phase transitions may occur for n > 2, and whether the phase diagram conjectured in Ref. [16] applies to other parts of the parameter space as well.

Therefore, our present aim is to supplement the results of the exact solutions with a numerical analysis of loop models in other parts of the parameter space. We pay particular attention to the range n > 2, where we expect an Ising-like phase transition takes place when the vertex weights are increased.

This expectation is based on the observation that, in the limit of large n, the local weights are maximal for configurations with small loops on the elementary faces of the lattice. At most one-half of these faces can be covered by a loop. There exist two checkerboardlike configurations at maximum covering, similar to the ordered phase of the hard-square model with nearest-neighbor exclusion. Thus, at a sufficient density of these loops, they will condense on one of the two sublattices.

At this point, we wish to specialize to the simplest nontrivial case, which is the case parametrized by a single bond weight. Since the x-type vertex covers one lattice edge, we denote the bond weight x. The z-type vertices cover two edges, we thus have $z=x^2$. For this choice, there is, except for the nearestneighbor exclusion, no further interaction between the hard squares in the limit $n \to \infty$. This avoids the complications that may arise when such further-neighbor interactions lead to different critical behavior. Also the y-type vertex covers one edge. Thus, in the present work we will study the case y=x, $z=x^2$ in the (x,n) plane.

As explained in Ref. [16], the y-type vertices are responsible for the flipping of the above-mentioned Ising-like degree of freedom along a loop. This leads to the possibility of different behavior when the y-type vertices are suppressed. For this reason the present work will also focus on a second case parametrized by n and a single bond weight x, namely the case y = 0, $z = x^2$.

For large n, we expect only small loops, and similar behavior in both cases. However, for small n larger loops exist with, if $y \neq 0$, straight segments due to the y-type vertices. Thus, for small n we may expect qualitative differences between the cases y = x and y = 0.

In Sec. II we sketch our numerical procedures used to locate the phase transition lines in the phase diagram. Section III presents the analysis of these numerical results, as well as the resulting phase diagram. The conclusions are summarized and discussed in Sec. IV.

II. TRANSFER-MATRIX ANALYSIS

Our analysis is based on the numerical transfer matrix (TM) calculation of Z_{loop} for $L \times \infty$ square lattices wrapped on a cylinder with circumference L. The transfer matrix keeps track of the change of the numbers of loops and the four kinds of vertices when a new layer of L sites is added. The TM techniques for the O(n) loop model and the procedure for the sparse-matrix decomposition are already described in the literature, e.g., see Ref. [16].

The largest eigenvalue Λ_0 of the TM determines the freeenergy density f(L) by

$$f(L) = \frac{\ln(\Lambda_0)}{L}.$$
 (8)

Its finite-size-scaling behavior at the critical point determines the conformal anomaly c according to [17,18]

$$f(L) \simeq f(\infty) + \frac{\pi c}{6L^2} + \cdots$$
 (9)

The magnetic correlation function of the O(n) spin model over a distance r can be expressed in terms of the probability that two vertices at this distance are connected by a single loop

segment [19]. Thus one may write

$$g_m(r) = \frac{Z'}{Z},\tag{10}$$

where Z' is the same as in Eq. (1), but with the sum on all loop configurations \mathcal{G}' that contain one additional single loop segment that runs from position 0 to r.

The exponential decay of $g_m(r)$ at large distances is determined by the magnetic correlation length $\xi_h(x,n,L)$, which can be obtained numerically as

$$\xi_h^{-1}(x, n, L) = \ln\left(\frac{\Lambda_0}{\Lambda_1}\right),\tag{11}$$

where Λ_1 is the largest eigenvalues in the "magnetic sector," which refers to the TM for Z', which is based on loop configurations with an additional single loop segment running along the cylinder. The scaled magnetic gap X_h is defined as

$$X_h(x, n, L) = \frac{L}{2\pi \xi_h(x, n, L)}.$$
 (12)

Its finite-size-scaling behavior [20,21] near a critical point x_c is given by

$$X_h(x,n,L) = X_h + a(x - x_c)L^{y_t} + bL^{y_u} + \cdots,$$
 (13)

where X_h is the magnetic scaling dimension, $y_t = 2 - X_t$ is the temperature exponent, and y_u is the leading irrelevant exponent. The amplitudes a and b are nonuniversal quantities.

In general, one expects that two phase transitions may occur in the two-dimensional O(n) model with $n \leq 2$ on the square lattice when the bond weight is increased [16]. The first one is the transition from the dilute loop phase to the low-temperature O(n) phase, where the loops are densely packed. The size of the longest loop diverges at this transition point. The universal properties of this transition follow from the exact solution [8,9] for branch 1, and from the Coulomb gas analysis [19]. The results for the conformal anomaly and the magnetic exponent are

$$c = 1 - \frac{6(g-1)^2}{g}, \quad X_h = 1 - \frac{1}{2g} - \frac{3g}{8},$$
 (14)

where g is the Coulomb gas coupling, which is related to n by $n=-2\cos(\pi g)$ and $1\leqslant g\leqslant 2$. The low-temperature phase is still critical in the sense that the magnetic correlation function decays algebraically in the infinite system. The universal properties of the low-temperature phase are characterized by a conformal anomaly $c^{(LT)}$ and a magnetic scaling dimension $X_h^{(LT)}$, which can be obtained from the results [8,9,19] for branch 2 of the square-lattice loop model. They are still given by Eq. (14) and $n=-2\cos(\pi g)$, but with $0\leqslant g\leqslant 1$.

A second transition may occur inside the low-temperature phase, when the loops enter an even denser phase which breaks the Ising-like symmetry of the loop configurations [16]. Its universal properties [8,16] were derived from the solvable case denoted as branch 4. The magnetic scaling dimension $X_h^{(\text{LTI})}$ and the conformal anomaly $c^{(\text{LTI})}$ at this Ising-like transition correspond with a combination of low-temperature O(n) and Ising-like critical behavior, namely

$$X_h^{(LTI)} = X_h^{(LT)} + 1/8$$
 (15)

and

$$c^{(LTI)} = c^{(LT)} + 1/2.$$
 (16)

To analyze the expected transition for n > 2, which drives the loop gas into a loop "solid" phase with a checkerboard pattern, we introduce the staggered loop density and interpret it as the order parameter. First, we define a face as "occupied" by a loop if it is surrounded by a loop or any odd number of loops. In analogy with the hard-square lattice gas, we also divide the faces of the lattice into "odd" and "even" ones. Then one defines the staggered loop density as the density of the occupied odd faces minus that for the even faces. The staggered lattice gas correlation function is thus

$$g_s(r) = \langle \rho_s(0)\rho_s(r)\rangle,$$
 (17)

where $\rho_s(0)$ and $\rho_s(r)$ are the staggered densities at positions 0 and r, respectively. For large n, we expect that the dense phase is dominated by configurations of elementary loops covering either the even or the odd faces. Therefore, the staggered correlation function is associated with the leading eigenvector \vec{v} of the TM that is antisymmetric under the operation \mathbf{R} , i.e.,

$$\vec{v} = -\mathbf{R}\vec{v}.\tag{18}$$

where **R** is the operator that rotates the lattice by one lattice unit about the axis of the cylinder. As a consequence of the Perron-Frobenius theorem, the absolute value of the corresponding TM eigenvalue Λ_2 cannot exceed Λ_0 which is associated with a symmetric eigenvector, at least for n > 0. We expect that the staggered correlation function scales in a similar way as the magnetic correlation function. Thus we describe the exponential decay of the staggered correlation function along the cylinder by means of the staggered scaled gap, defined as

$$X_s(x,n,L) = \frac{L}{2\pi} \ln\left(\frac{\Lambda_0}{\Lambda_2}\right). \tag{19}$$

The scaled gap $X_s(x,n,L)$ is expected to behave according to Eq. (13), with X_h replaced by the staggered lattice gas scaling dimension X_s . This transition breaks the Z_2 symmetry of odd and even lattice faces, and is thus expected in the Ising universality class: c = 1/2 and $X_s = 1/8$.

The critical point can be estimated by numerically solving *x* in the scaling equation involving two different system sizes,

$$X_i(x,n,L) = X_i(x,n,L'), \quad (i = h,s),$$
 (20)

of which the solution $x_c(L)$ scales as

$$x_c(L) = x_c + a' L^{y_u - y_t} + \cdots,$$
 (21)

where a' is an unknown constant. Because $y_u < 0$ and $y_t > 0$, $x_c(L)$ converges to the critical point x_c for a sequence of increasing system sizes L. At $x_c(L)$, the scaled gap in Eq. (13) converges to the magnetic or the staggered lattice gas scaling dimension $X_i(n)$ according to the scaling equation,

$$X_i(x_c(L), n, L) = X_i(n) + b'L^{y_u} + \cdots,$$
 (22)

with an unknown amplitude b'. An alternative way to obtain estimates $x_c(L)$ of the critical point is to neglect the correction term and thus to solve for x in the equation

$$X_i(x,n,L) = X_i(n), \tag{23}$$

where $X_i(n)$ is the theoretical prediction for the pertinent scaling dimension. Such predictions can follow the assumption that Eq. (14) or the Ising magnetic scaling dimension 1/8 applies. If this assumption is correct, the solutions of Eq. (23) converge to the critical point x_c as described by Eq. (21). If the assumption is not correct, the finite-size dependence of the solutions behaves as L^{-y_t} , so that they still converge to the critical point for $y_t > 0$, but relatively slowly. The finite-size dependence of the solutions of Eq. (23) may thus reveal if the assumed value of X_h is right.

In the present work, we shall make use of both Eqs. (20) and (23) to determine the critical points. In most of these calculations we restrict the system size to even values, because dense loop configurations do not fit well in odd systems, which thus leads to an even-odd alternation effect. Here we have given only a short summary of the relevant finite-size-scaling procedures; for more details, see Ref. [21].

III. RESULTS

In this section we explore the phase diagram as a function of n, for loop models described by Eq. (1) and a single bond weight x. The two cases y = x and y = 0 are treated in different subsections. We hope to find additional information beyond what is already known from exact solutions and earlier numerical work [8,9,16,22,23] for the square-lattice O(n) loop model, which have mostly been restricted to the range $n \le 2$.

A. Subspace $y = x, z = x^2$

In this subsection we explore the phase diagram for the case that there are no further conditions on the set of allowed vertices, thus with vertex weights x, y = x, and $z = x^2$.

We first consider the range $n \le 2$. Since we can compute the left-hand side of Eq. (23) numerically, we can solve that equation when we substitute the expected magnetic scaling dimensions X_h of the O(n) critical branch or of the low-temperature branch on the right-hand side. In this way, we obtain finite-size estimates of the critical point that can be extrapolated to the thermodynamic limit on the basis of the scaling behavior given in Eq. (21). That equation predicts convergence with increasing L if $y_u - y_t < 0$. Thus, the procedure still leads to convergent results in the case that the temperature field associated with x is irrelevant, but less irrelevant than the other nonzero scaling fields. The latter situation is expected for the low-temperature phase of the O(n) model with not too small n, on the basis of the existing results for branch 2 [9,16].

These transfer-matrix calculations were performed for system sizes up to L=16 or, in some cases, 18. Making use of Eq. (8), we then calculated the free-energy densities f(L) at the estimated critical point x_c , and obtained the conformal anomaly c by fitting these data according to Eq. (9). The numerical results for x_c and c are listed in Tables I and II respectively. Our numerical estimations of c agree well with the theoretical predictions, except for n=-2 where the finite-size data display poor convergence.

For the Ising-like transition in the LT dense phase, the numerical results for x_c at n = 2 and n = 1.5 are extrapolated from the solutions of the finite-size scaling equation (20) for

TABLE I. Numerical results for the critical points $x_c(n)$ and conformal anomaly c(n) for the O(n) critical branch in the range $-2 \le n \le 2$ in the subspace $y = x, z = x^2$. The theoretical values for branch 1 of the conformal anomaly are also listed. Estimated numerical uncertainties in the last decimal place are shown between parentheses.

\overline{n}	$x_c(n)$	$c_{ m num}$	$c_{\rm br1}$
-2	0.33732317(2)	-1.69(1)	-2
-1.5	0.3444544(1)	-1.009(1)	-1.00961
-1.0	0.35259515(1)	-0.60000(2)	-3/5
-0.5	0.3620756(1)	-0.27901(1)	-0.279017
0.0	0.3734237(1)	0	0
0.5	0.38757234(1)	0.25594(1)	0.255949876
1.0	0.406446(2)	0.50000(1)	1/2
1.5	0.4353496(1)	0.74183(2)	0.74184247
1.8	0.4664502(1)	0.89185(1)	0.89185788
1.9	0.484688(1)	0.94432(1)	0.9443219
1.95	0.4988697(1)	0.97151(1)	0.971508
1.98	0.512488(1)	0.98835(1)	0.988346
1.99	0.519766(2)	0.99411(2)	0.994103
2.0	0.5386256(2)	1.00000(1)	1

 $X_h(L)$ with even L. For other values of n, the critical points x_c are extrapolated from the solution of the scaling equation (23), in which X_h is taken as $X_h^{(LTI)}$, for even system sizes up to L = 16. These results are listed in Table III.

The free-energy densities f(L) at the estimated critical points were calculated for even system sizes up to L=16. Using a fit of these data according to Eq. (9), we then estimated the conformal anomaly c. The numerical results for c are included in Table III. They appear to be in a good agreement with the expected values $c^{(\rm LTI)}$ as given in Eq. (16).

Next, we explore the phase diagram for n > 2. For relatively large values $(n \ge 8)$, we numerically solved for $x_c(L)$ in Eq. (23), making use of the expected Ising value $X_s = 1/8$. For smaller values of n, the corrections to scaling become more prominent, and for n = 3,4,5 they are so large that solutions of Eq. (23) are absent. We then solved for $x_c(L)$ in the scaling equation (20). On the basis of these finite-size data, we obtained the critical point x_c , using extrapolation according to the finite-size scaling behavior given in Eq. (21).

TABLE II. Numerical results for the location $x_c(n)$ and the conformal anomaly c(n) for the low-temperature branch (branch 2) in the range $-2 \le n < 2$ in the subspace $y = x, z = x^2$. The theoretical predictions of the conformal anomaly are also listed. Estimated numerical uncertainties in the last decimal place are shown between parentheses.

n	$x_c(n)$	c (numerical)	c (theory)
1.99	0.559583(1)	0.99372(2)	0.993716
1.98	0.568975(1)	0.98725(2)	0.987247
1.95	0.58904(1)	0.96714(3)	0.967132
1.9	0.6145(1)	0.93180(1)	0.93179998
1.8	0.657(1)	0.8557(1)	0.855602
1.5	0.72(1)	0.588(1)	0.587572
1.4	0.74(1)	0.485(1)	0.4849998

TABLE III. Numerical results for the critical points $x_c(n)$ and the conformal anomaly c(n) for the Ising-like transition inside the LT dense phase in the subspace $y = x, z = x^2$. Estimated numerical uncertainties in the last decimal place are shown in parentheses. The results for c(n) agree well with the theoretical predictions $c_{\rm br4}$ for branch 4 of the square O(n) model.

\overline{n}	$x_c(n)$	c(n)	C _{br4}
-1.5	1.580(2)	-14.4(4)	-13.9612
-1.0	1.463(1)	-6.51(1)	-6.5
-0.5	1.38398(1)	-3.318(1)	-3.31779
0.0	1.3229(1)	-1.501(1)	-1.5
0.5	1.27287(2)	-0.318(3)	-0.319736
1.0	1.23019(2)	0.50000(4)	0.5
1.2	1.21468(1)	0.759(2)	0.758346
1.5	1.19282(2)	1.088(1)	1.08757
2.0	1.15943(2)	1.502(2)	1.5

We then calculated the free-energy density f(L) at the estimated critical points for even system sizes up to L=18. A fit of Eq. (9) to the data was then used to determine the conformal anomaly c. The best estimates of x_c and c are included in Table IV for several values of n>2. Our estimates of c agree well with the expected value c=1/2 for the Ising universality class, at least for large n. However, for smaller values of n, the estimates of c deviate from 1/2. A likely explanation is provided by the predicted strong corrections to scaling associated with the expected marginal temperature field at n=2.

TABLE IV. Numerical results for the critical points $x_c(n)$ and the conformal anomaly c(n) for the lattice-gas-like phase transitions. Results are shown for the two cases y=x and y=0. Estimated numerical uncertainties in the last decimal place are shown in parentheses. The results for c agree well with the expected Ising conformal anomaly c=1/2, except for n=2 and for some relatively small values of n where poor convergence occurs.

	Case $y = x$		Case $y = 0$	
n	$x_c(n)$	c(n)	$x_c(n)$	c(n)
2			0.784(1)	1.49(2)
3	1.101(2)	2.0(6)	0.809(1)	1.3(2)
4	1.12(2)	1.5(2)	0.800(4)	0.9(3)
5	1.00(2)	1.3(2)	0.787(3)	0.6(3)
8	0.907(2)	0.5(1)	0.746(1)	0.50(1)
10	0.857(3)	0.51(1)	0.7202(1)	0.50(1)
15	0.7716(3)	0.501(3)	0.6696(1)	0.50(1)
20	0.7147(3)	0.500(1)	0.6322(1)	0.50(1)
30	0.64060(1)	0.500(1)	0.5795(2)	0.498(2)
40	0.59240(1)	0.500(1)	0.54318(1)	0.500(2)
50	0.55754(1)	0.500(1)	0.51593(1)	0.500(2)
75	0.4995(1)	0.500(1)	0.46888(1)	0.500(1)
100	0.4622(2)	0.500(1)	0.43760(1)	0.500(1)
200	0.3841(1)	0.500(1)	0.36958(1)	0.500(1)
400	0.32004(1)	0.5001(1)	0.311448(1)	0.5000(1)
800	0.26726(1)	0.5001(1)	0.262179(2)	0.5000(1)
1000	0.25230(1)	0.5001(1)	0.248007(1)	0.5000(1)
10000	0.14033(1)	0.5000(1)	0.139573(1)	0.50000(1)

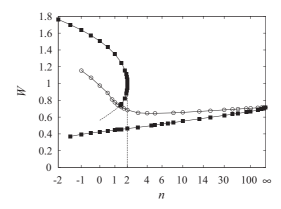


FIG. 2. The phase diagram of the O(n) loop model on the square lattice in the (n,x) parameter space. In order to map the range $-2 < n < \infty$ on a finite interval, a 1 - 8/(n + 10) scale is chosen along the horizontal axis. The vertical axis shows the temperaturelike quantity W, which is inversely proportional to the bond weight x as defined in the text. Data for the case y = x are shown as black squares, for y = 0 as open circles. The lines connecting these symbols are for visual aid only. The topmost curved line on the left is the O(n) critical line for x = y. Also shown is a part of its continuation into the low-temperature O(n) phase, which exists only for $n \le 2$. The vertical dashed line shows the boundary of this phase at n = 2. The lines of lattice-gas-like transitions extend all the way to $n = \infty$, where they meet in a point that is accurately known as described in the text.

In the limit $n \to \infty$, only loops of the smallest possible size occur, with the size of an elementary square of the lattice. The model of Eq. (1) then reduces to the lattice gas on the square lattice with nearest-neighbor exclusion and no further interactions. We make use of the existing numerical result $\mu_c = 1.334\,015\,100\,277\,4(1)$ for the critical value of the chemical potential of this model [24]. By relating the weight of an elementary loop to this chemical potential, which leads to $nx_c^4 = \exp(\mu_c)$, we obtain the large-n limiting behavior of x_c .

The phase diagram for n in the range $(-2,\infty)$ is shown in Fig. 2. In order to map the range $-2 < n < \infty$ on a finite interval, a 1 - 8/(n + 10) scale is chosen along the horizontal axis. The vertical axis shows the temperaturelike quantity $W \equiv 1/[x(n+10)^{1/4}]$, which parametrizes the bond weight, while remaining finite in the mentioned interval. The curved line on the left is the O(n) critical line and a part of its continuation into the low-temperature O(n) phase which exists only for $n \le 2$.

B. Subspace y = 0, $z = x^2$

As mentioned in Sec. I, for y=0 there exists an exactly solvable case $x_c=z_c=1/2$, which is called branch 0 [16]. For n=0 it describes the θ point of a collapsing polymer [25]. For other values of n, it describes a higher critical point (but not the tricritical point analyzed in Ref. [15]). Since the present value $z=x^2$ is smaller than that for branch 0, we do not expect that the y=0, $z=x^2$ subspace contains the n=0 collapse transition. However, the fact that an Ising degree of freedom is associated with each separate loop implies that a degree of Ising ordering is introduced at the critical points for n<2, where the largest loops are expected to diverge. Thus, we may expect that, at least for some values of n, transitions occur in a different universality class than that of branch 1. Furthermore

it remains to be investigated if the phase diagram displays the same topology as that for the $y \neq 0$ case.

First, we investigate whether the lattice-gas-like transition persists in the present subspace with y = 0. For $n \ge 8$, we solved for x_c in the scaling equation Eq. (23), after substituting the Ising magnetic scaling dimension $X_s(L) = 1/8$. For n =2,3,4,5, we solved instead the scaling equation Eq. (20) for $X_s(L)$, for the same reason as in the preceding subsection. Only even L were used in these calculations. We found that the solutions converge in the way described by Eq. (21), confirming that Eq. (13) applies to $X_s(L)$, thus indicating that algebraic decay of the staggered correlation function occurs in the thermodynamic limit. After extrapolation of the critical points x_c , we calculated the free-energy density f(L) at x_c for even system sizes up to L = 18. A fit of the data according to Eq. (9) then yields the conformal anomaly c. The results behave in a way similar to the y = x case: in accurate agreement with Ising universality (c = 1/2), except for a few relatively small n values. The numerical results for x_c and c are included in Table IV.

Next we address the question of whether the critical manifold continues into the $n \leq 2$ range and connects to an Ising-dense O(n) transition. We handle this problem by solving $x_c(L)$ in the scaling equation (20), for even system sizes up to L = 18. For n < 1, the magnetic scaled gap $X_h(L)$ is used in Eq. (20). We find that the solution $x_c(L)$ converges with L in the way described by Eq. (21). The estimated critical points are included in the phase diagram in the (x,n) plane shown in Fig. 2. We then calculated the magnetic scaled gaps $X_h(L)$ for a sequence of systems with even L up to 18 at the solutions $x_c(L)$. Extrapolation of the gaps according to Eq. (22) yields the scaling dimension X_h , which is listed in Table V. These results are, in a limited range, compatible with the known scaling dimension X_h of the critical O(n) transition, but the accuracy is low because of strong corrections to scaling, with the exception of the result at n = 0.

We also calculated the free-energy density at the estimated x_c . A fit of these data by Eq. (9) then yields estimates of the conformal anomaly c for this transition, which are also listed in Table V. For most values of n these results do not agree with the known theory for the O(n) critical line, or with a superposition of O(n) criticality and Ising behavior, although the numerical results still seem consistent with the latter situation in the range $1 < n \le 2$.

For small finite systems with n = -1, the leading eigenvalue Λ_0 becomes twofold degenerate at x = 1/2. The same applies to the leading eigenvalue Λ_1 in the odd (magnetic) sector. Moreover, these two pairs of eigenvalues are also equal. On this basis we conjecture that $x_c(-1) = 1/2$ and $X_h(-1) = 0$.

For $n \ge 1$, we found no solutions of the scaling equation Eq. (20) for $X_h(L)$ with even L. The staggered scaled gap $X_s(L)$ was used instead to study the possible transition. The scaling equation was solved for a sequence of even systems up to L = 18. We find that the solutions $x_c(L)$ behave in a way consistent with convergence to a critical point x_c as described in Eq. (21). The estimated critical points are included in the phase diagram in the (x,n) plane shown in Fig. 2.

Next we calculated the scaled staggered gaps $X_s(L)$ for even system sizes up to L=18 at the solutions $x_c(L)$. The gaps converge to the scaling dimension X_s according to Eq. (22).

TABLE V. Numerical results for the critical points $x_c(n)$, the conformal anomaly c(n), and scaling dimensions $X_h(n)$ and $X_s(n)$ for several values of n in the subspace y = 0. Since solutions of Eq. (20) for X_h are absent for $n \ge 1$, the result $X_h(1)$ was obtained from a fit of the $X_{h,L}$ data at the self-dual value $x = 1/\sqrt{2}$ of the bond weight. The latter value was also used to estimate $X_s(1)$ and c(1). It is in accurate agreement with the numerical result obtained by fits to the $X_s(L)$ data for n = 1. Satisfactory finite-size convergence is found only for n = -1, 0, and 1. The error margins, shown between parentheses, are difficult to estimate in some cases indicated with a question mark.

n	$x_c(n)$	$X_h(n)$	c(n)
-1.0	1/2	0	-2.00(1)
-0.8	0.51229(1)	0.06(2)?	-1.115(2)
-0.7	0.51889(1)	0.07(2)?	-0.92(1)
-0.5	0.53317(2)	0.07(1)	-0.60(1)
-0.3	0.54910(1)	0.09(2)	-0.339(1)
0.0	0.57686(2)	0.10417(5)	0
0.3	0.6106(3)	0.10(3)	0.306(1)
0.5	0.637(2)	0.16(4)?	0.506(1)
0.8	0.68(1)	0.21(3)?	0.81(1)
1.0	$1/\sqrt{2}$	0.25000000(5)	1.0000(1)
n	$x_c(n)$	$X_s(n)$	c(n)
1.0	$1/\sqrt{2}$	1.00000000(1)	1.0000 (1)
1.2	0.731(2)	0.81(4)	1.13(2)
1.5	0.755(4)	0.5(2)	1.32(1)
1.7	0.768(2)	0.3(3)	1.41(1)
2.0	0.784(1)	0.1(3)	1.49(2)

Unfortunately, the convergence of the data is not good except for n = 1. The results for X_s are listed in Table V. For n = 1 numerical result for x_c agrees with the self-dual value $x = 1/\sqrt{2}$ (see Sec. IV for further details), and the latter value was used to estimate the universal quantities for n = 1.

We also calculated the free-energy density for $1 \le n \le 2$ at the estimated critical bond weight x_c . From a fit of Eq. (9) to these data, we estimate the conformal anomaly c for this transition, as also listed in Table V.

IV. DISCUSSION

Using a finite-size-scaling analysis of results from transfermatrix calculations, we have determined the phase diagram of the O(n) loop model on the square lattice in the (x,n) plane, where x is the weight of a lattice edge covered by a loop. Two subspaces, y = x and y = 0, were investigated.

For $n \gg 2$ we find an Ising-like phase transition associated with the onset of a checkerboardlike ordering of the elementary loops. In this respect, the ordered state resembles that of the hard-square lattice gas. For large values of n the critical points shown in Fig. 2 approach the accurately known lattice-gas limit. For the case y = x, the data in this figure suggest that this approach happens with a weak cusplike singularity. This behavior can be explained by the residual presence of loops exceeding the size of an elementary face. The next-smallest loops cover a rectangle with the size of two faces, and contain two y-type vertices, at the expense of an extra weight factor x^2 . The presence of these larger loops thus corresponds with a repulsive potential of order $x^2 \propto 1/\sqrt{n}$ between

next-nearest-neighboring hard squares. In lowest order, one then expects a linear dependence of the critical chemical potential of the hard-square model on such a repulsion. Noting that the quantity W shown in Fig. 2 plays the role of this chemical potential, one expects that the critical value of W depends linearly on $1/\sqrt{n}$ for large n. This corresponds with a square-root-like singularity on the scale used in Fig. 2, which behaves as 1/n for large n. This explains the weak cusplike singularity in the y=x line in Fig. 2.

The appearance of loops exceeding the size of an elementary square, in particular loops covering two squares, can be interpreted as a softening of the nearest-neighbor repulsion. In this respect, the loop model with $n < \infty$ approaches the experimental situation of monatomic gases adsorbed on the (1,0,0) surface of a cubic crystal better than the hard-square model.

These results are in part similar to those obtained for the large-n loop model on the honeycomb lattice [26], which behaves as a hard-hexagon model [27], with a phase transition in the three-state Potts universality class. It appears that large-n loop models generically approach the behavior of systems of hard particles, with universal properties that are dependent on the microscopic lattice structure. The universal properties may also depend on the allowed set of vertices. For instance, if we put the vertex weights y = z = 0 in the loop model on the square lattice, then the corresponding hard-square model is subject not only to nearest-neighbor exclusion, but also to next-nearest-neighbor exclusion. The universal behavior of this system is not Ising-like [28,29], while the symmetry of the Hamiltonian remains the same. Crossover to a different universality class is thus possible when the parameters in the Hamiltonian are changed. An example thereof is already due to Das et al. [30], who added a staggered edge potential into the loop model on the honeycomb lattice, and observed crossover from three-state Potts to Ising behavior.

The question about a possible physical interpretation of this large-n transition of the loop model in the language of the spin model specified by Eq. (2) is answered by substitution of the numerical results x_c for the critical point, and the length scale $1/\sqrt{n}$ of the spin vectors in that equation. This shows that the Boltzmann factors of the spin model with n>2 can become negative at the phase transition. This exposes the unphysical nature of the lattice-gas-like transition in the language of the spin model.

In Sec. III we have also investigated the critical properties of this O(n) model in the range $-2 \le n \le 2$. In the case x = y, we found that the lattice-gas-like transition line continues into the low-temperature O(n) phase. The universal behavior along this part of the transition line is interpreted as a superposition of Ising criticality and dense O(n) loop model behavior, similar to earlier findings for a related square-lattice O(n) model. For the special point n = 1, the O(n) critical point and the Isinglike transition are dual images of one another. The duality transformation includes the sum on the weights of the two z-type vertices, thus leading to a single four-leg vertex with weight 2z, and a replacement of the loop segments by empty edges and vice versa. The transformation maps the x- and y-type vertices on the same type, and interchanges the empty and the z-type vertices. The normalization of the weight of the empty vertex to 1 thus reduces x and y by a factor 2z, and changes z into 1/4z. The n = 1 results for the critical points

 x_c found in Tables I and III satisfy this dual relation with one another.

Also for the case y = 0, we find that the lattice-gas-like transition line continues into the range n < 2, but the topology of the (x,n) phase diagram is different. It does not enter into the dense O(n) phase, but continues as a line of critical points separating the disordered phase from the dense O(n) phase. Our numerical data suggest that its universal properties do not match those of the critical O(n) line (for most of the range n < 1), or those of a superposition of Ising and critical O(n)behavior. Furthermore, our results for the universal quantities differ significantly from those reported for branch 3 of the square O(n) model [8,16] and for the tricritical O(n) model [31], at least for most of the range n < 1. We remark that both branch 3 of the square O(n) model and our y = 0 critical line lie relatively close to the higher critical point of branch 0 (x =z = 1/2, y = 0), but for -1 < n < 1 they reside on different sides of branch 0. This allows for the possibility that the y = 0critical point although also twofold unstable, is attracted by a different fixed point than branch 3. Unfortunately, the limited accuracy of our numerical data for the y = 0 line case impedes the further identification of its universal nature in terms of possibly existing exact results.

As already mentioned in Sec. I, there exists an Ising-like degree of freedom that is frozen out along each loop for y = 0. Thus, for n = 0, where we allow only a single loop, it plays no role and indeed we find the O(0) or self-avoiding walk (SAW) critical behavior. For $n \neq 0$ more loops may appear, whose Ising degree of freedom may differ. But, depending on the weight z, neighboring loops will tend to meet at z-type vertices, and thus assume the same Ising variable. It may thus be expected that at the y = 0 O(n) critical point, where the largest loop diverges, there will also be some ordering of the Ising degrees of freedom, thus allowing universal behavior that is different from that of the generic O(n) critical point.

For n=1, the numerical result for x_c agrees, within an error margin of about 10^{-5} , with the self-dual location $x_c(1)=1/\sqrt{2}$. Furthermore, this self-dual point can be mapped on Baxter's eight-vertex model [32], by adding down- or left-pointing arrows on the edges covered by an O(n) loop, and up- or right-pointing arrows to the empty edges. Since two of the vertex weights are zero, the symmetry relations of the eight-vertex model allow a further mapping on the six-vertex model, with vertex weights $(a,b,c)=(1/\sqrt{2},1/\sqrt{2},1)$ in the notation used in Ref. [32].

Also the conjectured critical point $x_c(-1) = 1/2$ mentioned in Sec. IIIB can be given a more firm basis. We recall that the weight z is actually redundant for n = -1 [16]. To see this, consider an arbitrary loop configuration where four loop segments come in at a given vertex. There are two possible ways to connect these segments by a z-type vertex, and the numbers of loops closed differ by precisely 1. Taking into account that the loop weight is -1, one observes that the two contributions due to the summation on the two z-type vertices cancel. Thus, in effect, configurations with z-type vertices do not contribute to the partition sum. Therefore, the n = -1 point of branch 0 in Ref. [16], namely x = 1/2, is an exact critical point in our y = 0 subspace, in spite of the fact that the weight z is different. The results $X_h = 0$ and c = -2 found there agree with the present findings.

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