Duality and the universality class of the three-state Potts antiferromagnet on plane quadrangulations

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1. INTRODUCTION

Ever since Kramers and Wannier’s [1] pioneering work on the two-dimensional (2D) Ising model, the concept of duality has led to important insights in statistical mechanics and quantum field theory [2] and more recently also in string theory [3]. The purpose of this Rapid Communication is to show an unusual application of duality to the study of the 3-state Potts antiferromagnet (AF) on a class of 2D lattices.

The q-state Potts model [4, 5] plays a key role in the theory of critical phenomena, especially in 2D [6–8], and has applications to various condensed-matter systems [5]. Ferromagnetic Potts models are by now fairly well understood, owing to universality; but the behavior of AF Potts models depends strongly on the microscopic lattice structure, so that many basic questions about the phase diagram and critical exponents must be investigated case-by-case. One expects that for each lattice \( \mathcal{L} \) there exists a value \( q_c(\mathcal{L}) \) [possibly noninteger] such that for \( q > q_c(\mathcal{L}) \) the model has an exponential decay of correlations at all temperatures \( T \) including zero, while for \( q = q_c(\mathcal{L}) \) the model has a zero-temperature critical point. The first task, for any lattice, is thus to determine \( q_c \).

Some 2D AF models at \( T = 0 \) have the remarkable property that they can be mapped exactly onto a “height” model (in general vector-valued) \([9–13]\). Since the height model must either be in a “smooth” (ordered) or “rough” (massless) phase, the corresponding zero-temperature spin model must either be ordered or critical, never disordered. When the height model is critical, the long-distance behavior is that of a massless Gaussian with some \((a \text{ priori unknown})\) “stiffness matrix” \( K > 0 \). The critical operators can be identified via the height mapping, and the corresponding critical exponents can be predicted in terms of \( K \). Height representations thus provide a means for recovering a sort of universality for some (but not all) AF models and for understanding their critical behavior in terms of conformal field theory (CFT).

In particular, on any plane quadrangulation (i.e., any
implies a one-to-one correspondence between quadrangulations $\Gamma$, $Q$. Conversely, given a dual pair $(\Gamma, Q)$ of plane graphs, we can construct a quadrangulation $Q(Q_0) = \mathcal{Q}(G_1)$ with the vertex set $V = V_0 \cup V_1$ by connecting each vertex in $G_0$ to the neighboring vertices in $G_1$. There is thus a one-to-one correspondence between quadrangulations $\Gamma$ and dual pairs of plane graphs $(G_0, G_1)$. We shall say that the quadrangulation $Q(Q_0)$ is of self-dual type if $G_0$ is self-dual, and of non-self-dual type otherwise. For instance, the square lattice is a quadrangulation of self-dual type (both $G_0$ and $G_1$ are themselves square lattices), while the diced lattice is a quadrangulation of non-self-dual type (the sublattices are triangular and hexagonal).

Let us henceforth restrict attention to periodic planar lattices. It is well known (and obvious) that the square lattice is self-dual; what seems to be less well known is that there exist infinitely many examples of self-dual periodic planar lattices [20–26], including the “hextri” lattice [21, Figs. 1 and 10], [23, Fig. 16] [24, Fig. 1b], the “house” lattice [21, Fig. 2], the martini-B lattice [25, Fig. 8], and the cmm-pmm lattice [23, Fig. 29]. In particular, from each of these lattices we can construct the corresponding quadrangulation of self-dual type.

In this Rapid Communication we present the results of our study — using Monte Carlo (MC), transfer matrices (TM), and critical polynomials (CP) [27] — of the 3-state Potts AF on a variety of quadrangulations of both types. We find empirically, without exception, the following behavior:

**Conjecture 1** For the 3-state Potts AF on a (periodic) plane quadrangulation $\Gamma$:

1. If $\Gamma$ is of self-dual type, the model has a zero-temperature critical point, so that $q_c = 3$. This critical point has central charge $c = 1$.
2. If $\Gamma$ is of non-self-dual type, the model has a finite-temperature phase transition, so that $q_c > 3$. This transition is second-order and lies in the universality class of the 3-state Potts ferromagnet.

To our knowledge, prior to this Rapid Communication, only four AF Potts models on planar lattices with a critical point at $T = 0$ were known [9, 12]: the square and kagome lattices with $q = 3$, and the triangular lattice with $q = 2$ and $q = 4$. By contrast, Conjecture 1 implies that this phenomenon is not so exceptional: There are infinitely many $q = 3$ models displaying it!

We have studied four quadrangulations of self-dual type: $Q(\text{hextri})$ [see Fig. 1], $Q(\text{house})$, $Q(\text{martini-B})$ and $Q(\text{cmm-pmm})$. We have also considered four quadrangulations of non-self-dual type: $Q(\text{diced})$ (see Fig. 2), $Q(\text{martini})$, $Q(\text{ruby})$, and $G_0^F$ [19, Fig. 2(b)]. As the qualitative behavior of the four lattices within each class turns out to be the same, we refrain from giving here all the details [28], and shall focus on one lattice of each type: $Q(\text{hextri})$ and $Q(\text{diced})$.

## II. QUADRANGULATIONS OF SELF-DUAL TYPE

If the quadrangulation is of self-dual type, then we expect the number of ideal states [9, 12] to be six: The system must choose which of the two sublattices to order, and in which of the three possible spin directions.

![Figure 1. The quadrangulation $Q(\text{hextri})$.](image1)

![Figure 2. The quadrangulation $Q(\text{diced})$, which is also the Laves lattice $D(3, 4, 6, 4)$ and is the dual of the ruby lattice. The black (resp. gray) vertices form a diced (resp. kagome) sublattice.](image2)
It is therefore natural to expect (by using universality arguments) that, as for the square lattice, there will be a critical point at \( T = 0 \) characterized by a CFT with central charge \( c = 1 \) [28].

We first investigated the 3-state Potts AF by extensive MC simulations on lattices of size \( L \times L \) unit cells with periodic boundary conditions (BC), using the Wang–Swendsen–Kotecký (WSK) cluster algorithm [29]. As our lattices are bipartite, this algorithm is known to be ergodic even at \( T = 0 \) [11, 30, 31]. For each lattice, we measured the staggered and uniform susceptibilities, which are expected to diverge at the critical point as \( \chi_{\text{stagg}} \sim L^{(\gamma/\nu)_{\text{stagg}}} \) and \( \chi_{u} \sim L^{(\gamma/\nu)_{u}} \). The qualitative behavior of these susceptibilities is the same for all four lattices considered here, but the critical exponents \( (\gamma/\nu)_{\text{stagg}} \) and \( (\gamma/\nu)_{u} \) do depend on the lattice. In Fig. 3 we show, as an example, the scaled staggered susceptibility on the \( Q(\text{hextri}) \) lattice (we use, instead of the standard Potts-model coupling constant \( J \), the variable \( v = e^{J} - 1 \)). All the finite-\( L \) curves meet at \( v = -1 \), implying that this point is indeed critical.

The height representation [9–12] relates these susceptibility exponents to the stiffness \( K \) (which is a scalar in this case):

\[
(\gamma/\nu)_{\text{stagg}} = 2 - \frac{\pi}{18K}, \quad (\gamma/\nu)_{u} = 2 - \frac{2\pi}{9K}.
\]

The results for the four lattices studied here, along with the known exact values for the square lattice [11, 12], are displayed in Table I. The value of the stiffness is in all cases much smaller than the critical value \( K_{c} = \pi/2 \approx 1.570796 \) where the locking potential becomes marginal, which separates the rough and smooth phases [9, 10, 12].

We also found that, for all these lattices, the WSK algorithm does not suffer from critical slowing-down (CSD). By measuring the integrated autocorrelation times \( \tau_{\text{int}} \) for the staggered and uniform susceptibilities at \( T = 0 \), we find that \( \tau_{\text{int}} \lesssim 8 \) uniformly in \( L \). This phenomenon also occurs for the square-lattice model [12, 30]. We conjecture that the WSK algorithm for the 3-state Potts AF on any quadrangulation of self-dual type has no CSD.

We also studied the \( Q(\text{hextri}) \) lattice by means of a TM approach. We considered strip graphs of this lattice with cylindrical BC and widths \( 2 \leq L \leq 14 \). In Fig. 1, our TM propagates from left to right. We measured the free energy (per unit area) \( f_{\text{L}}(q) \) at \( T = 0 \) in the AF regime for \( 2 \leq q \leq 4 \). The central charge \( c(q) \) can be extracted using the standard CFT Ansatz [32, 33]

\[
f_{\text{L}}(q) = f_{\text{bulk}}(q) - \frac{c(q)\pi}{6L^{2}} + o(L^{-2}).
\]

We first observed that there are parity effects depending on the value of \( L \mod 4 = 0 \). We then ignored the \( o(L^{-2}) \) terms, fitted the values corresponding to \( L, L + 4 \) to (2), and extracted the estimates \( c_{L}(q) \). This curve exhibits, for each value of \( L \), a maximum value \( c_{\text{max}}(L) \) at \( q = q_{\text{max}}(L) \). These values are displayed in Table II, together with our extrapolations to \( L = \infty \) (see details of the fits in Ref. [28]). These results agree well with our conjecture that the \( q = 3 \) Potts AF on \( Q(\text{hextri}) \) is critical at \( T = 0 \), with behavior described by a CFT with \( c = 1 \).

| \( \Gamma \) | \( (\gamma/\nu)_{\text{stagg}} \) | \( (\gamma/\nu)_{u} \) | \( K \) |
|---|---|---|---|
| \( Q(\text{cmm-pmm}) \) | 1.71762(9) | 0.8691(5) | 0.6177(6) |
| \( Q(\text{hextri}) \) | 1.7024(3) | 0.8096(9) | 0.5865(6) |
| \( Q(\text{house}) \) | 1.6978(3) | 0.7922(4) | 0.5778(8) |
| \( Q(\text{martini-B}) \) | 1.6882(3) | 0.7557(9) | 0.5609(6) |
| square | 5/3 | 2/3 | \( \pi/6 \) |

Table I. Critical exponents \( (\gamma/\nu)_{\text{stagg}} \) and \( (\gamma/\nu)_{u} \), and the estimated stiffness \( K \), for the zero-temperature 3-state Potts AF on the quadrangulations \( \Gamma \) of self-dual type studied in this Rapid Communication. We include for comparison the exact values for the square lattice [12].

| \( L \) | \( q_{\text{max}}(L) \) | \( c_{\text{max}}(L) \) |
|---|---|---|
| 2 | 3.8544146155 | 0.8508786050 |
| 4 | 3.2788982545 | 1.013354086 |
| 6 | 3.1443621430 | 1.055838007 |
| 8 | 3.0975518402 | 1.075432576 |
| 10 | 3.0795627986 | 1.0383006482 |
| \( \infty \) | 3.00(2) | 0.99(2) |

Table II. \( q_{\text{max}}(L) \) and \( c_{\text{max}}(L) \) for the \( T = 0 \) \( q \)-state Potts AF on the \( Q(\text{hextri}) \) lattice with cylindrical BC as a function of the width \( L \), and the extrapolation to \( L = \infty \).
III. QUADRANGULATIONS OF NON-SELF-DUAL TYPE

When the quadrangulation is of non-self-dual type, the asymmetry between the two sublattices suggests that at $T = 0$ one preferred sublattice will be ordered (in one of the three possible spin directions) and the other sublattice disordered (between the other two states). If this is so, then at $T = 0$ there are only three ideal states each of them with one sublattice ferromagnetically ordered. Therefore, we have the same $\mathbb{Z}_3$ symmetry and ground-state degeneracy as for the 3-state Potts ferromagnet, and hence we expect a first-order finite-temperature second-order transition in the universality class of this latter model. However, a first-order finite-temperature transition is also possible.

In particular, if the two sublattices have unequal vertex densities (as occurs most often), then we expect that the sublattice with the smaller (resp. larger) vertex density will be ordered (resp. disordered), as this maximizes the entropy. The reasoning becomes more subtle, however, if the two sublattices have equal vertex densities [as occurs, for instance, for $Q$ (diced) and $Q$ (rubby)]: Then it is not obvious how the asymmetry alone can drive the phase transition. For this reason we focus here on $Q$ (diced).

Once again we studied the 3-state Potts AF using MC simulations on lattices of size $L \times L$ unit cells with periodic BC, using the WSK algorithm. In all cases, we find a finite-temperature critical point. We followed the practical methods of [14] to locate the critical point, and then fitted our numerical data to the finite-size-scaling (FSS) Ansatz

$$O_L = L^\omega \left[ O_c + a_1(v - v_c)L^{1/\nu} + a_2(v - v_c)^2 L^{2/\nu} + b_1 L^{-\omega_1} + \cdots \right].$$

For each lattice, we measured the staggered susceptibility $\chi_{\text{stagg}}$ and the Binder cumulant $R_{\text{stagg}} = \langle M_{\text{stagg}}^4 \rangle / \langle M_{\text{stagg}}^2 \rangle^2$. The qualitative behavior of these observables is the same for all four lattices considered here. As an example, we show in Fig. 4 the data for the scaled staggered susceptibility of the $Q$ (diced) lattice, together with our preferred FSS fits based on the Ansatz (3) with a varying number of terms. Since our results for the critical exponents were compatible with the predicted values $(\gamma/\nu)_{\text{stagg}} = 26/15$ and $\nu = 5/6$ [6], we then redid the fits fixing these parameters to the predicted values, in order to obtain improved estimates for $v_c$. Our results are shown in Table III, and agree well with the prediction that the model lies in the universality class of the 3-state Potts ferromagnet.

For the lattice $Q$ (diced) we have checked directly from the MC simulations that, even though the two sublattices $G_0$ and $G_1$ have the same vertex density, it is the diced sublattice $G_0$ (black vertices in Fig. 2) that becomes ordered. More precisely, the sublattice of $G_0$ consisting of degree-6 vertices is the one that is most ordered; the two degree-3 sublattices of $G_0$ are slightly more ordered than the three sublattices of the kagome sublattice $G_1$. Therefore, although both sublattices $G_0$ and $G_1$ give naively the same entropy density, it is the one having a sub-sublattice with the largest degree that becomes ordered, because fluctuations around these three ideal states maximize the system’s entropy density. A similar phenomenon occurs for the $Q$ (ruby) lattice [28].

On all these lattices (as well as on the diced lattice [14]), the WSK algorithm suffers from CSD, with dynamic critical exponents $z_{\text{int},M_2} = 0.50(1)$ and $z_{\text{int},M_4} = 0.48(1)$. If these exponents are in fact equal, then our preferred estimate (taking into account the statistical non-independence of the two estimates) would be $z_{\text{int}} = 0.49(2)$. This is compatible with the exponent $z_{\text{int},M_4} = 0.475(6)$ found in the Swendsen–Wang (SW)


| \( n \) | \( v_c(n) \) |
|---|---|
| 2 | -0.9344496491145567949 |
| 4 | -0.93889690618313817225 |
| 6 | -0.93976678350525022210 |
| 8 | -0.9401709879171472205 |
| 10 | -0.94038789257375557598 |
| 12 | -0.94051494788357303489 |

Table IV. Real roots of \( P_B(3,v) \), to 20-digit numerical precision, for \( Q(\text{diced}) \). We show the unique real root \( v_c(n) \) in the AF interval \( v \in [-1, 0] \), for \( n \times \infty \) bases, together with the extrapolation to \( n = \infty \) (where we used exponents in the range 1.2–1.5, which are much smaller than those for the ferromagnetic models investigated in Refs. [27, 37, 38]).

Finally, we have applied the CP method [27, 37, 38] to study the location of the critical point for the 3-state Potts AF on the \( Q(\text{diced}) \) lattice. We computed the CP \( P_B(3,v) \) for some bases \( B \) that admit a four-terminal representation (to be able to use the TM method of Ref. [37]). In particular, to compute the estimates of \( v_c \) shown in Table IV, we have used the more powerful eigenvalue method of Ref. [38], which allows us to use bases of size \( n \times m \) in the limit \( m \to \infty \). The last row of Table IV shows the extrapolation to \( n = \infty \) using Monroe’s implementation of the Bulirsch–Stoer [39] extrapolation scheme. This result agrees within errors with the MC estimate, but it is more precise.

\section{Conclusions}

We have studied the 3-state Potts AF on four quadrangulations of self-dual type, and on four quadrangulations of non-self-dual type (including two with equal vertex densities on the two sublattices), by extensive computations using MC simulations, TM computations, and the CP method. In all cases, we have found a perfect agreement with Conjecture 1. Our findings provide very strong empirical support for the validity of this criterion. However, we do not want to exclude the possibility that for some lattices of non-self-dual type the finite-temperature transition might be first-order.

As a side result, we have also found that the WSK algorithm has no CSD when simulating the 3-state Potts AF on any quadrangulation of self-dual type, while it has CSD (compatible with the dynamic universality class of the SW algorithm for the 3-state ferromagnet) on any quadrangulation of non-self-dual type.

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