Exact bond percolation thresholds in two dimensions

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Abstract
Recent work in percolation has led to exact solutions for the site and bond critical thresholds of many new lattices. Here we show how these results can be extended to other classes of graphs, significantly increasing the number and variety of solved problems. Any graph that can be decomposed into a certain arrangement of triangles, which we call self-dual, gives a class of lattices whose percolation thresholds can be found exactly by a recently introduced triangle–triangle transformation. We use this method to generalize Wierman’s solution of the bow-tie lattice to yield several new solutions. We also give another example of a self-dual arrangement of triangles that leads to a further class of solvable problems. There are certainly many more such classes.

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1. Introduction

The simple geometrical model of percolation on regular lattices has been studied extensively since it was first proposed nearly 50 years ago [1]. It continues to find broad applications to such diverse problems as understanding conductive materials [2, 3], the fractality of coastlines [4], networks [5–7], turbulence [8, 9], colloids [10] and the spin quantum Hall transition [11]. Along with the Ising model of a ferromagnet and the related Potts model, percolation serves as a paradigm of statistical mechanics.

In the usual bond percolation model each edge (bond) of an infinite graph or lattice is defined to be open with probability \( p \) (and closed with probability \( 1 - p \)) independently of all other bonds. If \( p \) is near 0 most bonds are closed, but there are small and sparsely distributed open clusters. With \( p \) close to 1, the landscape is dominated by an infinite open cluster. Between these two regimes, there lies a sharply defined critical threshold, \( p_c \), which is a characteristic of each lattice, at which the infinite cluster makes its first appearance. Finding this critical point is central to many theoretical and computational studies of percolation.
Figure 1. The triangle–triangle transformation is shown in (a). The lattices in (b) and (d) are invariant under this transformation, as shown in (c) and (e).

for many years, exact thresholds were known for only a relatively small number of two-dimensional lattices (square, triangular, honeycomb, bow-tie . . . ) found through duality and a technique called the star–triangle transformation [12, 13].

Recently, it has been recognized that the star–triangle transformation can be generalized for systems with correlated bonds [14, 15] or for systems composed of triangular cells more complicated than a simple star or triangle [16]. Using this generalization, site (in the site problem we consider vertices occupied with probability \( p \)) and bond thresholds have been found for many new lattices, including the ‘martini’ (figure 3), ‘martini-A’ and ‘martini-B’ lattices [14, 16]. Recently, Wu [17] has shown that the transition point of the Potts model (which includes both the Ising model and percolation as special cases) can also be found explicitly for these three lattices—the first new exact transition points for the Potts model found in many years.

In this paper we briefly discuss these new results, and show how they can be extended further to systems where the triangular cells do not necessarily lie in a triangular lattice, but in any self-dual arrangement (in a sense defined below). In this way we provide a derivation of the threshold for Wierman’s bow-tie lattice [13] and generalizations of it, as well as other new lattices.

2. Triangle–triangle transformation

We consider any system that can be decomposed into distinct triangles, like those shown in figure 1. For lattices of this type that are also invariant under a certain transformation, which we call the triangle–triangle transformation, it is possible to determine exactly the bond threshold. The triangle–triangle transformation is illustrated in figure 1(a) where we superpose a reversed (hatched) triangle over each (shaded) triangle of the original lattice (figure 1(c)). If the lattice formed by the hatched triangles is the same as the original, we say that the lattice is self-dual under the triangle–triangle transformation. The shaded triangle (the generator) can represent any network of bonds, correlated bonds, and sites, as long as they are contained between three vertices. This means that every self-dual arrangement of triangles gives an infinite family of solvable lattices. Figure 2 shows an example, the kagomé arrangement, that can be decomposed into triangles but is not self-dual under the triangle–triangle transformation.
Figure 2. (a) The kagomé lattice. As clear from (b) this system is not self-dual under the triangle–triangle transformation.

Although some special lattices fall into this arrangement, such as the double bond honeycomb lattice, and can be solved exactly, this is not true for every choice of generator. For example, choosing a simple triangle with uncorrelated bonds, in figure 2(a), gives the kagomé lattice. This lattice’s bond threshold cannot be found by the following method and, in fact, the problem remains unsolved [18].

To find the general condition for criticality, consider any self-dual arrangement with vertices $A, B, C$ on the original triangles, and $A', B', C'$ on the dual triangles as in figure 1. To be at the critical threshold, we require two things (in exact analogy with the usual star–triangle transformation): (1) that the connection probabilities between the vertices of $A, B, C$ and those of the vertices of $A', B', C'$ are the same, and (2) that $A, B, C$ and $A', B', C'$ are dual to each other. The former implies that the overall connection probabilities between points on the lattice and corresponding ones on the dual lattice are the same, and the latter implies that this equi-probability point must be the critical point (because only at the critical point can a lattice and its dual have identical correlations).

To state these conditions mathematically, we let $P(A, B, C)$ equal the probability that $A, B$ and $C$ on a triangle are connected, $P(A, B, C)$ equal the probability that $A$ and $B$ are connected but not $C$, etc, and $P(\overline{A}, \overline{B}, \overline{C})$ equal the probability that none are connected. Likewise define $P'(A', B', C')$ etc for the connection probabilities on the dual triangles. Then the condition of equality of connection probabilities is expressed by

\begin{align}
    P(A, B, C) &= P'(A', B', C') \\
    P(A, B, \overline{C}) &= P'(A', B', \overline{C'}) \quad (2)
\end{align}

Second, we require that the two triangles $A, B, C$ and $A', B', C'$ are dual to each other. This implies

\begin{align}
    P(A, B, C) &= P'(\overline{A}, \overline{B}, \overline{C}) \\
    P(A, B, \overline{C}) &= P'(A', B', \overline{C'}) \\
    P(\overline{A}, \overline{B}, \overline{C}) &= P'(A', B', C').
\end{align}

For example, equation (4) says simply that if $A, B$ and $C$ are connected to each other, then $A', B'$ and $C'$ must all be isolated. This is because the dual triangular unit $A', B', C'$ is created
by drawing bonds intersecting all the bonds of $A$, $B$, $C$, and placed down with complementary probabilities $P_i = 1 - p_i$. Thus, whenever there is an occupied bond on the original lattice, the dual-lattice bond is not occupied, and vice versa. Therefore, if $A$, $B$ and $C$ are all connected, the bonds between $A'$, $B'$ and $C'$ must necessarily be all unconnected. The same kind of argument holds for the other conditions.

Comparing the two sets of equations (1)–(3) and (4)–(6), we see that the middle (two-point) equations are identical. The first and third equations yield the requirement that the probability that all the vertices connect is the same as the probability that none of them connect:

$$P(A, B, C) = P(\overline{A}, \overline{B}, \overline{C}).$$

This simple general relation determines the critical point, and was given independently in [15] and [16]. It is only a function of the arrangement of bonds in the individual triangles and not dependent upon how the triangles are arranged to form a lattice, as long as they are arranged in an overall self-dual way as those in figure 1. In our previous work [16], (7) was applied only to systems of triangular cells that were on a triangular array (figure 1(b)). The lattices we studied, along with their generators, are shown in figure 3. As already mentioned, the site thresholds for these lattices can also be found exactly. In fact, they were initially derived by a different method [14] but were then shown to fall into the current framework by using generators that involve sites as well as bonds [16]. These bond and site thresholds are given in tables 1 and 2. Here we apply (7) to any self-dual set of triangles such as those in figures 1(d) and 6. The generalization of (7) to all self-dual arrangements is shown by the arguments above, and was suggested by the earlier work of Wierman on the bow-tie lattice [13].

3. Martini and related lattices

The result from which all our exact solutions will follow is the critical threshold of the martini lattice in which each bond of the generator is assigned a different probability, as shown in...
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Table 1. Bond percolation thresholds of the lattices in figure 3 and their associated polynomials derived from equation (8).

<table>
<thead>
<tr>
<th>Lattice</th>
<th>( p_{bc}^{\text{bond}} )</th>
<th>Equation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Martini</td>
<td>( 1/\sqrt{2} \approx 0.707107 \ldots )</td>
<td>((2p^2 - 1)(p^4 - 3p^3 + 2p^2 + 1) = 0)</td>
</tr>
<tr>
<td>Martini-A</td>
<td>0.625 457 \ldots</td>
<td>( p^5 - 4p^4 + 3p^3 + 2p^2 - 1 = 0 )</td>
</tr>
<tr>
<td>Martini-B</td>
<td>1/2</td>
<td>((2p - 1)(p^2 - p - 1) = 0)</td>
</tr>
</tbody>
</table>

Table 2. Site percolation thresholds of the lattices in figure 3 and their associated polynomials. \( \phi = (1 + \sqrt{5})/2 = 1.618034 \ldots \) is the golden ratio, which makes an interesting appearance here.

<table>
<thead>
<tr>
<th>Lattice</th>
<th>( p_{bc}^{\text{site}} )</th>
<th>Equation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Martini</td>
<td>0.764 826 \ldots</td>
<td>( p^4 - 3p^3 + 1 = 0 )</td>
</tr>
<tr>
<td>Martini-A</td>
<td>( 1/\sqrt{2} \approx 0.707107 \ldots )</td>
<td>( 2p^2 - 1 = 0 )</td>
</tr>
<tr>
<td>Martini-B</td>
<td>1/\phi = 0.618034 \ldots</td>
<td>( p^2 + p - 1 = 0 )</td>
</tr>
</tbody>
</table>

Figure 3(a). This results in a critical surface rather than a critical point and application of (7) implies that it is given by [17, 18]

\[
1 - p_1p_2r_3 - p_2p_3r_1 - p_1p_3r_2 - p_1p_2r_1r_2 - p_1p_2r_1r_3 - p_2p_3r_2r_3 + p_1p_2p_3r_1r_2 + p_1p_2p_3r_2r_3 + p_1p_2p_3r_1r_3 + p_1p_3r_1r_2r_3 - 2p_1p_2p_3r_1r_2r_3 = 0. \tag{8}
\]

If we set all bonds equal to \( p \), we get the condition \((2p^2 - 1)(p^4 - 3p^3 + 2p^2 + 1) = 0\), which has solution on \([0, 1]) \( p_c = 1/\sqrt{2} \). Setting \( p_1 = 1 \) and \( p_2 = p_3 = r_1 = r_2 = r_3 = p \) gives the martini-A lattice, and setting \( p_2 = p_3 = 1 \) and \( p_1 = r_1 = r_2 = r_3 = p \) gives the martini-B lattice, as listed in table 1.

Taking \( r_1 = 1 \), we get the result for the critical surface of the general honeycomb lattice [12], \( 1 - p_1p_2 - p_1p_3 - p_2p_3 + p_1p_2p_3 = 0 \), and taking \( p_1 = 1 \) yields the critical surface of the general triangular lattice [12], \( 1 - r_1 - r_2 - r_3 + r_1r_2r_3 = 0 \). Setting all probabilities equal in the last two expressions gives polynomial equations with the well-known solutions \( p_c(\text{honeycomb}) = 1 - 2\sin\pi/18 \) and \( p_c(\text{triangular}) = 2\sin\pi/18 \). It is a general fact that thresholds found via (7) are solutions of polynomials with integer coefficients.

4. Bow-tie class of lattices

Although equation (8) was originally derived for the martini lattice, it gives the critical surface of any lattice in which the martini generator is arranged in a self-dual way. Here we apply the result to the arrangement given in figure 1(d). Taking \( p_1 = 1 \) in this case gives precisely the lattice of figure 1(d) where the shaded triangle is just a simple triangular cell. This lattice has a double bond where the bases of abutting triangles join. However, we can use a trick, due to Wierman [13], to combine these two bonds by setting the probability of each to be \( r_3 = 1 - \sqrt{1 - p} \). The probability of traversing the double bond is now just \( p \), which effectively combines them into one. With this replacement, we get Wierman’s bow-tie lattice (figure 4(a)). Setting the single bonds, \( r_1 \) and \( r_2 \), to \( p \) gives the condition \( 1 - p - 6p^2 + 6p^3 - p^3 = 0 \), with solution in \([0, 1]) \( p_c = 0.404 518 \ldots \) as found by Wierman. But we can just as well make the basic triangular cell the ‘A’ generator (figure 3(b)) which produces the lattice shown in
Figure 4. Some examples of generalized bow-tie lattices.

figure 4(d), with a transition point identical to that of the regular 'A' lattice (given in table 1) since no double bond arises in this case. Of course, there is an infinite variety of additional lattices that can be solved by this procedure and it becomes a somewhat subjective matter to decide which ones are most interesting; some of our choices are shown in figure 4, with their duals in figure 5 and thresholds in table 3. For (c), we use a martini generator with $p_1 = \sqrt{p}$.
and $p_2 = p_3 = r_1 = r_2 = r_3 = p$, while for (b) we use an ‘A’ generator with an extra bond on the bottom, which results in a doubled bond that must be combined.

In figure 6(a) we show another arrangement of triangles that is self-dual under the triangle–triangle transformation, defining another family of solvable lattices. For example, replacing the shaded triangles by the ‘A’ generator, one can get an interesting variation on the ‘A’ lattice with the same threshold, shown in figure 6(b). Many other self-dual arrangements can be constructed.

Thus, we have shown that a great variety of lattices with exactly determined percolation thresholds can now be found. We also see that there are many variations of a given lattice (such as the lattices with the A-lattice threshold shown in figures 3(b), 4(d) and 6(b)) that share the same threshold, even though the bonds are in different arrangements. Even after decades of study, the percolation model continues to yield new exact results.

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References