A Brief Introduction to Percolation Theory

Joshua Mankelow 1902187

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Abstract

Consider a cube of water-permeable material. What is the probability that that if water is poured on top of the cube it may drain all the way through the cube and out the opposite face? Initially developed by Paul Flory and Walter Stockmayer in 1944, percolation theory attempts to answer such questions by rephrasing them in terms of vertices (sites) and edges (bonds) of graphs and examining the connectedness of such graphs. The connectedness of these graphs—in the infinite case—is determined by a threshold probability, p_c , describing whether the water may pass through each site or bond. This essay will introduce the ideas of site and bond percolation as well as the notion of clusters and critical (threshold) probabilities. We will also analyse the one dimensional case to garner a basic understanding before exploring higher dimensional cases. After discussing the concepts of percolation theory, we will move on and look at the many applications of the theory discussed in the earlier parts of the essay.

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1 Introduction

1.1 The canonical example

Let us consider the example from the abstract of water filtering through a porus medium, but this time in two dimensions. How do we model this? One might imagine that the medium consists of many particles arranged (for simplicity) in an $n \times n$ square lattice and linked to each of their nearest neighbours. Clearly, this is the lattice on \mathbb{Z}^2 . To set up the problem, each of the particles will be expressed as a vertex in a graph and each of the links will be an edge. In the context of percolation, a vertex is called a site and an edge is called a bond; these sites and bonds form a network.

Definition 1.1. A vertex in a graph is referred to as a **site**.

Definition 1.2. A edge in a graph is referred to as a **bond**.

Definition 1.3. A graph is referred to as a **network**.

So what does percolation actually mean? First we shall introduce the notion of open and closed sites and bonds and then we can discuss percolation.

Definition 1.4. A site or bond in a network is labeled **open** if it allows whatever we're considering to pass through.

Definition 1.5. A site or bond in a network is labeled **closed** if it doesn't allow whatever we're considering to pass through.

And now for the percolation definitions.

Definition 1.6. We say that we are considering **site percolation** if we let all of the sites in the network be open with probability $p \in [0,1]$ and closed with probability $1-p \in [0,1]$. We refer to p here as the percolation probability.

Definition 1.7. We say that we are considering **bond percolation** if we let all of the bonds in the network be open with probability $p \in [0,1]$ and closed with probability $1-p \in [0,1]$. We refer to p here as the percolation probability.

Now that we've defined site and bond percolation, what's the problem that we're trying to solve? In the case of water being poured on a porus medium, we would like to know whether there is an open path from the top of the network to the bottom. We shall model this using site percolation (in fact, all examples in this essay will be using site percolation unless explicitly stated otherwise).

Definition 1.8. We say that a path in a network is **open** if:

- when considering site perolation, every site in the path is open.
- when considering bond percolation, every bond in the path is open.

Definition 1.9. Let N = (V, E) be a network and let $A, B \in V$. The sites A, B are **openly connected** if there exists an open path connecting A and B.

Definition 1.10. Let N = (V, E) be a network and let $A, B \in V$. The sites A, B are **openly disconnected** if there does not exist an open path connecting A and B.¹

The probability that an open path from the top of the network to the bottom exists depends on both our choices of both p and n from before. As a result of our context, our value for n should be large—this is the case with most percolation models—but we shall use small n for the sake of example and simplicity. Let us now fix n and see what happens as we vary p. Obviously we have two trivial cases, p=0 and p=1, where the network is completely openly disconnected and completely openly connected respectively. What about when $p \in (0,1)$? Let's inspect three different values of p on our network: p=0.25, p=0.5 and p=0.75 as shown in figures 1a, 1b and 1c on page 5. As one might expect, as p increases, so does the "connectedness" of the network; i.e. the probability of having an open path from the top of the network to the bottom increases with p. Also, observe that as p increases we also get larger "clusters" of open connected sites or bonds.

Definition 1.11. Let N = (V, E) be a network. A **cluster** is a set of vertices, $C \subset V$, such that if $v_1, v_2 \in C$ then v_1 and v_2 are openly connected.

Definition 1.12. Let N = (V, E) be a network and let $C \subset V$ be a cluster. The **size** of C, denoted by |C|, is the number of sites in the cluster. A cluster of size s may be referred to as an s-cluster.

We will explore clusters in more detail in the next section, but for now let us introduce the idea of a critical probability. Although they don't exist in the finite cases, they do exist in the infinite cases.

Definition 1.13. Consider a network, N = (V, E). A cluster, $C \subseteq V$, is considered **infinite** if and only if $|C| = \infty$.

Definition 1.14. Consider an infinite network N=(V,E). The **critical probability**, denoted p_c , is the percolation probability such that the probability that there exists a cluster, $C \subseteq V$ with $|C| = \infty$ is 1.

Add computational examples of the connectedness as p changes

¹Definitions 1.8, 1.9 and 1.10 are equivalent for both site and bond percolation.

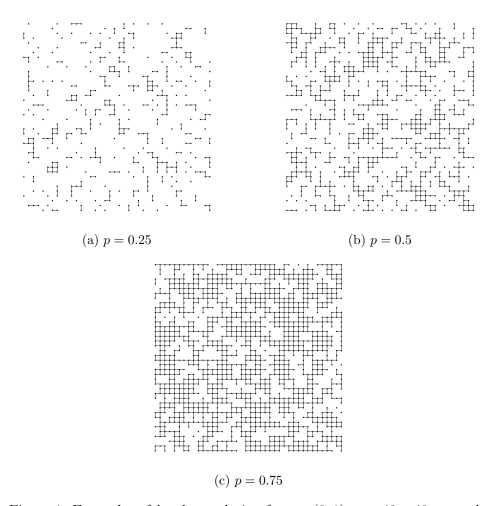


Figure 1: Examples of bond per colation for $p \in (0,1)$ on a 40×40 network.

1.2 Other network configurations

We have already seen the lattice on \mathbb{Z}^2 as an example, but there are many more. To remain within the scope of this essay, we shall only breifly mention some two and three dimensional examples and print their site and bond critical probabilities and a diagram.

Definition 1.15. A network is considered **regular** if every site in that network has the same number of bonds attached to it.

Definition 1.16. The **coordination number** of a regular network is the number of bonds attached at every site. This quantity is denoted using the letter Z. I.e. the lattice on \mathbb{Z}^2 has a coordination number Z = 4.

Two dimensional network configurations

Clearly, one two dimensional network configuration is the lattice on \mathbb{Z}^2 . In context this is referred to as the square lattice. Other regular two dimensional network configurations include, but are not limited to, the Bethe Lattice (Figure 4a), Honeycomb Lattice (Figure 4b), Kagome Lattice (Figure 4c) and the Triangular Lattice (Figure 4d). As one might imagine, each of these configurations has a different (but not necessarily distinct) critical probability. Below is a table showing the critical probabilities for each of the aforementioned network configurations. It should be noted that probabilities marked with a * (star) are exact results. [Sahimi, 1994, p. 11]

Configuration	Z	p_c for bond percolation	p_c for site percolation
Bethe	3	TO FIND	TO FIND
Honeycomb	3	$1 - 2\sin(\pi/18)^*$	0.6962
\mathbb{Z}^2 (Square)	4	1/2*	0.5927
Kagome	4	0.522	0.652
Triangular	6	$2\sin(\pi/18)^*$	1/2*

Figure 2: Critical probabilities for various configurations of two dimensional networks

Three dimensional network configurations

It shouldn't be hard to guess that the lattice on \mathbb{Z}^3 is a potential configuration for three dimensional networks. We call this configuration the Simple Cubic Lattice (Figure 5b). Similarly to the two dimensional case, there many other regular three dimensional network configurations. These include, but again are not limited to, the Diamond Lattice (Figure 5a), the

Configuration	Z	p_c for site percolation	p_c for bond percolation
Diamond	4	0.3886	0.4299
Simple Cubic	6	0.2488	0.3116
BCC	8	0.1795	0.2464
FCC	12	0.198	0.119

Figure 3: Critical probabilities for various configurations of three dimensional networks

Body Centered Cubic (BCC) Lattice (Figure 5c) and the Face Centered Cubic (FCC) Lattice (Figure 5d). Notice how none of these results are precise. [Sahimi, 1994, p. 11]

2 The one dimensional case

To develop an understanding of how to analyse these networks, we shall consider the one dimensional case where our network is the lattice on \mathbb{Z} , or a "chain". We shall use the same idea of percolation probability from definitions 1.6 and 1.7.

Theorem 2.1. The critical probability, p_c , for the lattice on \mathbb{Z}^2 when considering site percolation is $p_c = 1$.

If we recall the definition of the critical probability, the value of p_c that we're looking for is intuitively 1. In order to prove this theorem, we shall introduce some more machinery. This machinery isn't necessary for the proof, but it helps with understanding and will allow us to analyse more interesting cases later on.

Definition 2.1. If the percolation probability on the chain is p, we define the **number of s-clusters per site** by the following quantity:

$$n_s = p^s (1 - p)^2$$

This quantity represents the probability that any given site in the network is the left end of an s-cluster. So, if our network is of length L >> s, then we will have $Ln_s = Lp^s(1-p)^2$ clusters of size s on average. This quantity also allows us to explore the probability that any given site is part of an s-cluster. Such a probability is given by the quantity $n_s s$.

The idea of percolation is to understand whether a path exists from one side of the network to the other, so we have to use a slightly different definition for the one dimensional case. This new definition fixes an issue that I'll address after the proof.

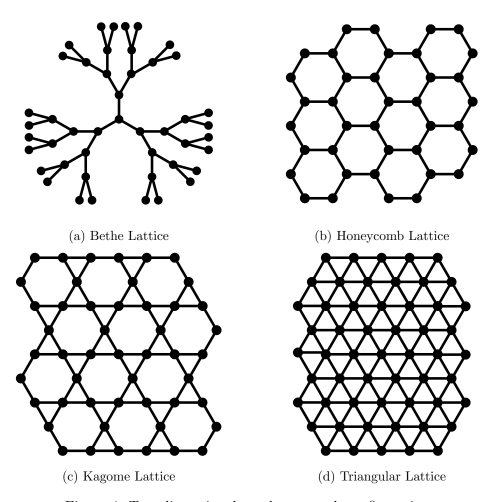


Figure 4: Two dimensional regular network configurations

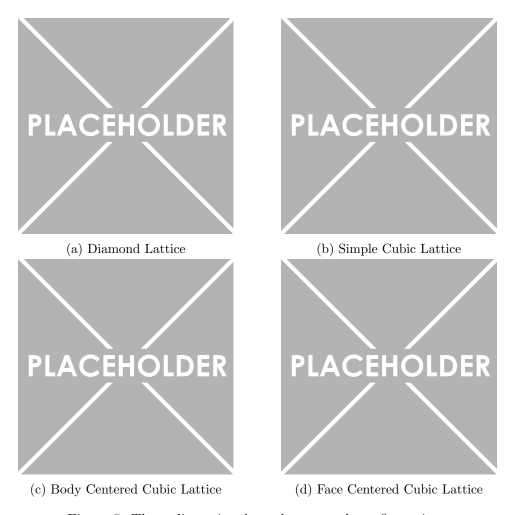


Figure 5: Three dimensional regular network configurations

Definition 2.2. Consider the lattice on \mathbb{Z} of the form N=(V,E). The critical probability, denoted p_c , is the percolation probability such that the probability that there **uniquely** exists a cluster, $C \subseteq V$, with $|C| = \infty$ is 1.

This proof and the following corollary are heavily inspired by the proof and subsequent corollory from Dietrich Stauffer's Introduction to Percolation Theory. [Dietrich Stauffer, 1991]

Proof. (Theorem 2.1) We shall prove by contradiction. Let us assume that $p_c \in [0,1)$ and is fixed. This implies that a chain of length L will have, on average, L(1-p) closed sites. As $L \to \infty$, $L(1-p) \to \infty$ showing us that there is at least one closed site in the chain and that means there is no continuous row of occupied sites. Thus $p_c = 1$ in order to have only one infinite cluster.

So why doesn't this proof work if we hadn't made that ammendment to the definition. Notice that \mathbb{Z} is a countably infinite set and all of the L(1-p)closed sites form a subset of \mathbb{Z} . This means that the set of all closed sites is also countably infinite. This situation may be rephrased in a way such that we must partition a countably infinite set into countably infinite subsets.

prove

this?

The above results allow us to get some more interesting information about the behaviour of our system. For example, we can deduce the follow-

Corollary. When considering the lattice on \mathbb{Z} with percolation probability $p \in [0,1)$, the following equality holds:

$$\sum_{s} n_s s = p$$

This result comes from the fact that every open site must belong to a cluster of some size s. So summing $n_s s$ over all s must give us p. This equality may also be derived using the definition of n_s and the formula for a geometric series.

Proof.

$$\sum_{s} n_s s = \sum_{s} p^s (1-p)^2 s$$

$$= (1-p)^2 \sum_{s} p \frac{d(p^s)}{dp}$$

$$= (1-p)^2 p \frac{d(\sum_{s} p^s)}{dp}$$

$$= (1-p)^2 p \frac{d(p/(1-p))}{dp}$$

$$= p$$

It's worth noting that this equality doesn't hold for p=1, because $n_s=1^s(1-1)^2=0$ so $\sum_s n_s s=0$. The technique of considering the sizes of clusters and the number of empty sites surrounding them is also used when analysing more compelx structures. Say, for example, you have a cluster, C, of size |C|=9 on the lattice on \mathbb{Z}^2 . This cluster could take many different shapes with different numbers of empty sites surrounding it. For example, this cluster could be a straight line of open sites which has a total of 22 closed sites surrounding it. Therefore the probability that a cluster like this exists at any site given a percolation probability, p, is $p^9(1-p)^{20}$. This cluster could also be a square of open sites with side length 3. This would have 12 closed sites surrounding it and thus the probability of a cluster of this shape existing at any given site is $p^9(1-p)^{12}$. The ability for clusters to be the same size but have different "perimeters" is what makes analysing these problems in higher dimensions so difficult.

3 Applications

References

[Dietrich Stauffer, 1991] Dietrich Stauffer, A. A. (1991). *Introduction to Percolation Theory*. Taylor & Francis, 11 New Fetter Lane, London EC4P 4EE.

[Sahimi, 1994] Sahimi, M. (1994). Applications of Percolation Theory. Taylor & Francis, University of Southern California.