Universality class of site and bond percolation on multi-multifractal scale-free planar stochastic lattice

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In this article, we investigate both site and bond percolation on a weighted planar stochastic lattice (WPSL) which is a multi-multifractal and whose dual is a scale-free network. The characteristic properties of percolation is that it exhibits threshold phenomena as we find sudden or abrupt jump in spanning probability across \( p_c \), accompanied by the divergence of some other observable quantities which is reminiscent of continuous phase transition. Indeed, percolation is characterized by the critical behavior of percolation strength \( P(p) \sim (p_c - p)^\beta \), mean cluster size \( S \sim (p_c - p)^{-\gamma} \) and the system size \( L \sim (p_c - p)^{-\nu} \) which are known as the equivalent counterpart of the order parameter, susceptibility and correlation length respectively. Moreover, the cluster size distribution function \( n_s(p_c) \sim s^{-\tau} \) and the mass-length relation \( M \sim L^{d_f} \) of the spanning cluster also provide useful characterization of the percolation process. We obtain an exact value for \( p_c \) and for all the exponents such that there is the probability that the spanning cluster also provide

I. INTRODUCTION

Percolation is perhaps one of the most studied problems in statistical physics. This is not only because of the simplicity of its definition but also because of the versatility of its applications. To study percolation one needs to choose a skeleton first. It can be a lattice or a graph that has two entities namely sites (nodes) and bonds (edges). We then occupy each site or bond, depending on whether we want to study site or bond percolation, with probability \( p \) independent of the state of its neighbors. Broadbent and Hammersley in 1957 first presented the percolation model to understand the motion of gas molecules through the maze of pores in carbon granules filling a gas mask. Since then the intuitive idea of percolation has been found relevant to so many seemingly disparate systems that its concept has literally percolated across a vast area of science and social science. Examples include flow of fluid in porous media, infiltration in composite materials processing, spread of fluids, rumours, opinion, biological and computer viruses are just a few to mention.

Besides the simplicity of its definition and the versatility of its application there exists yet another reason why percolation model is so popular. In percolation we primarily observe how clusters, set of contiguous occupied sites, are formed and grown as a function of \( p \) which is the only control parameter. As \( p \) value increases from negligibly small, there appears for the first time a cluster that spans across the entire system. In the case of infinite system size, we find a unique threshold value \( p_c \) such that there is the probability that the spanning cluster \( W(p) = 0 \) for \( p \leq p_c \) and \( W(p) = 1 \) for \( p > p_c \). Interestingly, such transition, despite being geometric in nature, yet we find many of its aspects reminiscent of continuous thermal phase transition (CTPT) \[12,13\]. Thus, percolation serves as a relatively tractable model for the investigation of phase transition and critical phenomena that lie at the heart of the modern development of statistical physics. This is perhaps the most important reason why percolation is still studied extensively even after almost 60 years of its inception.

Indeed, for almost every observable quantities in percolation there exist an equivalent counterpart in CTPT. These observables like their counterpart in CTPT, exhibit power-law, at least near \( p_c \), which is typically attributed to critical phenomena. For instance, the system size \( L \) is like correlation length \( L \sim (p_c - p)^{-\nu} \), mean cluster size \( S \) is like susceptibility \( S \sim (p_c - p)^{-\gamma} \), percolation strength \( P \) is like order parameter \( P \sim (p_c - p)^\beta \) etc. Like thermal phase transition, percolation transition too can be classified in terms of \( p_c \) and by a set of critical exponents \( \beta, \gamma, \nu \). One of the extraordinary findings in percolation is that the numerical value of its critical exponents depend neither on the detailed nature of the lattice structure nor on the type of percolation, bond or site. Their values depend only on the dimension of the embedding space of the lattice. It is, therefore, said that percolation on all planar lattices belong to the same universality class.

Unique universality class has been found true for a variety of periodic and non-periodic planar lattices having fixed and mixed-valued coordination number, random planar lattices and their dual, random multifractal lattices etc. \[14,17\] (see also Ref. \[18\], which is the most
recent review article). Yet, have we exhausted all the possible lattices to conclude that percolation on all planar lattices belongs to the same universality class? The answer is no. Recently, we have reported that the site percolation on a weighted planar stochastic lattice (WPSL) belongs to separate and distinct universality class [19]. The WPSL is quite non-trivial as it has mixed properties of both lattice and network or graph [20]. On one hand, unlike networks it is embedded in the space of dimension $d = 2$, on the other unlike regular lattice, its coordination number distribution obeys a power-law. We found that the critical exponents for site percolation on the WPSL are totally different from the known values for all other planar lattices studied till today. We, therefore, claim that the random site percolation on the WPSL belong to a separate and distinct universality class.

In this article, we investigate the bond percolation on the WPSL and present detailed results of its site counterpart in order to see the contrast. One of the goals of the present article is to check if the bond and site percolation on WPSL belong to the same universality class like for all known planar lattices studied to date. First, we find the percolation threshold $p_c$, for both bond and site percolation, using the idea of spanning probability $W(p)$. Second, we attempt to find an estimate for the various critical exponents such as $\nu, \beta$ and $\gamma$ using the finite-size scaling hypothesis where precise value of $p_c$ is necessary. Then, we use the idea of data collapse for further fine tuning of the estimated values for the exponents till we get the best data-collapse. Besides critical exponents, we also find the exponent $\tau$ that characterizes the cluster size distribution function $n_c(p_c) \sim s^{-\tau}$ and the fractal dimension $d_f$ that characterizes the mass of the spanning cluster $M(p_c) \sim L^{d_f}$. Note that the values of the various critical exponents and the exponents $\tau, d_f$ etc. are not at all independent rather they are bound by some scaling relations. We use these scaling relations for self-consistency check. We find that our estimate for various exponents satisfy these relations up to quite a good extent. Our results based on extensive Monte Carlo simulation suggest that both site and bond percolation on WPSL belong to the same universality class and it is different from the one where percolation on all the planar lattices belong.

The rest of the article is organized as follows. In section II, we discuss the algorithm for the construction of WPSL and some of its key features. In section III, we briefly discuss the Newman-Ziff algorithm as it is the most efficient algorithm for percolation. We also discuss the finite-size scaling and underline its deep connection to the Buckingham II-theorem in section IV. In section V, we present our results about bond and site percolation on the WPSL side by side so that we can appreciate the contrast. Finally, we summarize our results in section VI.

II. WPSL AND ITS PROPERTIES

We first give a brief description of the construction process of the WPSL. It starts with an initiator which we choose to be a square of unit area. The generator is then defined as the one that divides the initiator (in step one) randomly into four smaller blocks. In step two and thereafter the generator is applied to only one of the blocks by picking it preferentially with respect to their areas. Consider the $t$th time step of the generation of the WPSL at which the system has $3t - 2$ number of blocks available whose areas are say $a_1, a_2, a_3, ..., a_{3t-2}$. To pick one from $3t - 2$ blocks we subdivide an interval of unit length $[0, 1]$ into $(3t - 2)$ sub-intervals of size $[0, a_1]$, $[a_1, a_1 + a_2]$, ..., $[\sum_{i=1}^{3j-3} a_i, 1]$ so that the higher the area the greater the size of the sub-intervals. We then generate a random number, say $R$, from the interval $[0, 1]$ and find which of the $(3t - 2)$ sub-intervals contain this $R$ and pick that block. This process ensures that the blocks are being picked preferentially according to their size. In Fig. 1 we give a snapshot of the lattice to give a visual impression of how it actually looks at any given time. It is a space-filling planar cellular structure where the size or the area of the cells in the lattice are not equal rather their distribution is random. This is in sharp contrast to many of the cellular structures that we are familiar with. One advantage of creating WPSL by random sequential partitioning of the square into ever smaller mutually exclusive rectangular blocks helps defining each step of the division process as one time unit. The number of blocks $N$ at time $t$ therefore is $N = 1 + 3t$ and hence it grows albeit the sum of the areas of all the blocks is always equal to the size of the initiator. Thus, the number of blocks $N$ increases with time at the expense of the size
of the blocks.

Recently, we have shown that the area size distribution of the blocks of WPSL obey dynamic scaling

\[ c(a, t) \sim t^\theta \phi(a/t^z), \tag{1} \]

where we found \( \theta = 2 \) and \( z = 1 \) [21]. It implies that the snapshots of the lattice at different times are similar. Yet another interesting properties of this lattice is that the dynamics of the system is governed by infinitely many conservation laws one of which is the conservation of total area. To be more precise, if we denote \( x_i \) and \( y_i \) as the length and width of the \( i \)th block then we can show analytically that

\[ M_n = \sum_{i=1}^{N} x_i^n y_i^{4/n-1} \]

assumes statistically a constant value regardless of the time \( t \) when the snapshot is taken [20]. We have also shown that, except the conservation of total area, each of the infinitely conserved quantity is a multifractal measure. That is, we can assume that the \( i \)th block of the lattice is populated with probability

\[ p_i \sim x_i^{n-1} y_i^{4/n-1}. \]

We have shown that within the multifractal formalism we can construct the partition function which is the \( q \)th moment of \( p_i \), i.e.,

\[ Z_q = \sum_i p_i^q. \tag{2} \]

Measuring \( Z_q \) as a function of the square root of the mean block area

\[ \delta = \sqrt{\frac{\text{area of the initiator}}{\text{total number of blocks}}} = \sqrt{\frac{1}{1 + 3t}} \sim t^{-1/2}, \tag{3} \]

one can show that \( Z_q \) exhibits power-law

\[ Z_q(\delta) \sim \delta^{-\tau(q, n)}, \tag{4} \]

with exponent

\[ \tau(q, n) = \frac{4/3 - n^2q^2 + 16 - ((4/n + n - 2)q + 2)}{2}. \tag{5} \]

One of the characteristic features of this exponent is that it is non-linear for \( n = 2 \).

Note that the exponent \( \tau(q, n) \) has two interesting properties. First, \( \tau(q, n) = 2 \) \( \forall n \) at \( q = 0 \) which is the dimension of the embedding space of the WPSL. Second, \( \tau(q, n) = 0 \) \( \forall n \) at \( q = 1 \) as it is required by the normalization condition [22]. The Legendre transform of \( \tau(q, n) \) is a method whereby its derivative

\[ \alpha = -\frac{d\tau(q, n)}{dq}, \tag{6} \]

can be considered as an independent variable instead of \( q \) itself. In general, if we denote \( \alpha \) as the slope and \( f \) as the intercept then the equation for the straight line is

\[ \tau(q) = -\alpha q + f(\alpha). \tag{7} \]

The function \( f(\alpha) \) is the Legendre transform of the function \( \tau(q) \) which is always concave in character. It implies that for every \( n \) value there exist a spectrum of spatially intertwined fractal dimensions

\[ f(\alpha(q, n)) = \frac{16}{\sqrt{(4/n - n)^2q^2 + 16}} - 2, \tag{8} \]

which are needed to characterize the WPSL except for \( n = 2 \). Note that the maximum of \( f(\alpha, n) \) occurs at \( q = 0 \) which corresponds to the dimension of the embedding space of the WPSL when blocks are assumed empty. We thus find that the WPSL is a multi-multifractal planar lattice.

Besides, WPSL is a planar cellular structure whose cells or blocks has coordination number disorder in the sense that unlike regular lattice it has great many different number of neighbors. In fact, its coordination number distribution exhibits a power-law [20]. This is in sharp contrast to the coordination number distribution in the Voronoi diagram where it is also random but its distribution is peaked around the mean [23]. In the Voronoi diagram it is almost impossible to find cells or blocks which have significantly higher or fewer neighbours than the mean coordination number. That is, here the mean describes the characteristic scale. Such characteristic scale is absent in the WPSL since the distribution function follows a power-law. The power-law coordination number distribution also means that the majority of the blocks in the WPSL are very poor in coordination number and there are few cells or blocks which have significantly high number of nearest neighbours. A lattice, so rich in properties can be of great interest as it can mimic disordered medium on which one can study problems like percolation or random walk. In brief, the WPSL has the following properties:

i) Its area size distribution function obeys dynamic scaling.

ii) It obeys infinitely many conservation laws.

iii) It is a multi-multifractal.

iv) Its coordination number distribution function obeys power-law.

III. NEWMAN-ZIFF ALGORITHM

In the standard algorithms, such as the Hoshen-Kopelman (HK), one must create an entire new state for every given value of occupation probability \( p \) in every independent realization. Investigation of the various observable using such traditional algorithms are highly expensive in terms of computational time and accuracy of finding various observable quantities. In 2000, Newman and Ziff (NZ) proposed an algorithm which is highly efficient in both account [24]. The efficiency in the NZ algorithm lies in the fact that one creates a new state with \( n + 1 \) occupied sites or bonds from the immediate
previous state with $n$ occupied sites or bonds simply by occupying one extra randomly chosen site or bond. It is based on the intuitive idea of random sequential adsorption of sites or bonds on a given lattice or graph. The algorithm is trivially simple. One starts with an empty lattice. Then at each step an empty site or bond is chosen at random and then is occupied if empty; else the attempt is discarded. However, in order to further reduce the computation time we first decide an order in which the sites or bonds will be occupied. That is, we wish to choose a random permutation of the bonds or sites. This is done by creating a list of all the bonds in any convenient order. Positions in this list are numbered from $1, 23, ..., M$. Choose a number $j$ at random with uniform probability in the range $i \leq j \leq M$. Then use any standard textbook algorithm to randomize the number $i = 1$ to $M$ and put them in a new order in which they will be occupied. Having chosen an order of all the sites, we start occupying them in that order. The first site or bond to be occupied will definitely form a cluster of size one. The second, third, fourth etc too are highly likely to form clusters of size one. However, the likelihood of forming clusters of size one will decrease with the number of occupied sites since some sites when occupied, will become contiguous occupied sites thus making clusters of size more than one.

The formation of clusters and the statistics of their sizes are the key to the study of percolation theory. In the case of NZ algorithm we measure an observable, say $O$, for fixed numbers of occupied sites (or bonds), and obtain a data for $H$ as a function of occupation number $n$. This is in sharp contrast with the HK algorithm where the number of sites being occupied at a given $p$ is random and different at every independent realization. However, if the system size is large enough then the mean occupation number will almost equal to $pN$ where $N$ represents the system size. The weight factor of obtaining different $n$ for a given $p$ are not the same. The exact weighting factor of there being exactly $n$ occupied sites on the lattice for a given $p$ is given by binomial distribution

$$C(n, N, p) = \sum_{n=1}^{N} \binom{N}{n} p^n (1-p)^{N-n}. \quad (9)$$

The binomial coefficient $\binom{N}{n}$ represents the number of possible configurations of $n$ occupied sites and $N-n$ empty sites. Using this and the data for the observable $O$ for all values of $n$ we can find $O$ for any value of $p$ by the following relation

$$O(p) = \sum_{n=1}^{N} \binom{N}{n} p^n (1-p)^{N-n}O_n. \quad (10)$$

It is interesting to note that the ensemble of states with exactly $n$ occupied sites or bonds obtained according to NZ algorithm can referred to as a microcanonical percolation ensemble, where the number $n$ is the equivalent counterpart of the energy $E$ in thermal statistical mechanics. On the other hand, if we keep $p$ fixed instead of $n$ we can regard it as the canonical ensemble.

## IV. Finite-Size Scaling and II-Theorem

We offer here a brief introduction to the spirit and scope of the scaling approach to phase transitions and critical phenomena in general. It is well-known as finite-size scaling (FSS) hypothesis. It has been extensively used as a very powerful tool for estimating finite size effects near the threshold value of the controlling parameter. In the continuous phase transition, the various response functions, typically the second derivative of the free-energy, diverges. Such transitions are classified by a set of critical exponents. The best known example of continuous phase transition is the paramagnetic to ferromagnetic transition where it has been found that

- magnetization $M \sim (T - T_c)^\beta$,
- susceptibility $\chi_M \sim (T - T_c)^{-\gamma}$,
- and correlation length $\xi \sim (T - T_c)^{-\nu}$.

In percolation, their equivalent counterparts are

- percolation probability $P \sim (p - p_c)^\beta$,
- Mean cluster size $S \sim (p - p_c)^{-\gamma}$,
- and system length $\xi \sim (p - p_c)^{-\nu}$. \quad (12)

These relations are only true in the thermodynamic limit in the sense that the system size is infinite. It is important to appreciate the fact that we can neither do experiment nor simulation on infinite systems where the correlation length $\xi \sim L$. To overcome this impediment, physicists have come up with a smart solution which is known as finite-size scaling. In general, an observable quantity, say $X$, of the threshold phenomena that exhibit continuous phase transition is said to obey finite-size scaling if it satisfies

$$X(p, L) \sim L^{a/\nu} \phi((p - p_c)L^{1/\nu}), \quad (13)$$

where $a$ and $\nu$ are said to be critical exponents. It provides an elegant way of extrapolating critical exponents for infinite system from a set of data for finite systems using the idea of data collapse.

We shall here show that the origin of the FSS theory is actually deeply rooted to the Buckingham II-theorem as it can be systematically obtained following the prescription of that theorem [22]. Consider that a quantity $X$ is the primary quantity of interest which depends on the control parameter $x$ and the system size $L$ so that we can write

$$X = X(x, L). \quad (14)$$

Note that in the case of threshold phenomena, where there is a critical or threshold value $x_c$ across which the
system under goes a sudden or abrupt change, we find that the distance \( x - x_c \) is a better variable than \( x \) itself. Indeed, the observable quantity \( X \) is found to depend on \( x - x_c \) and hence we write

\[
X \sim X(x - x_c, L).
\]

We almost always find that the quantity \( x - x_c \) diminishes with \( L \) following a power-law \( (x - x_c) \sim L^{-a} \). It implies that we can choose one of the parameters, say \( L \), to have an independent dimension. Thus the dimension of \( X \) too can be expressed in terms of \( L \) alone

\[
X \sim L^b.
\]

Following the argument of the \( \Pi \)-theorem we can now define two dimensionless quantities

\[
\xi = \frac{x - x_c}{L^{-a}},
\]

and

\[
\Pi = \frac{X}{L^b} \equiv \phi(\xi, L).
\]

Note that \( \phi \) being a dimensionless quantity its numerical value must remain invariant, for a given value of \( \xi \), even if we change \( L \) by an arbitrary factor and hence \( \phi(\xi, L) = \phi(\xi) \). We can thus immediately write that

\[
X(x, L) \sim L^b \phi((x - x_c)/L^{-a}).
\]

The reduction of initially two variable problem into one variable problem constitutes the basic statement of the Buckingham \( \Pi \)-theorem. This is traditionally known as an hypothesis in the literature namely as the finite-size scaling hypothesis.

A quantitative way of interpreting how the experimental data exhibit finite-size scaling is done by invoking the idea of the data-collapse method - an idea that goes back to the original observation of Rushbrooke [12]. The plots of \( X(x, L) \) vs \( x \) for different \( L \) always result in distinct curves. However, the same data can be made to collapse on a single universal curve if one plot \( XL^{-b} \) vs \( (x - x_c)L^a \) instead of \( X(x, L) \) vs \( x \) regardless of the size of \( L \). The quality of data collapse depends on how exact the value of \( x_c \) and the exponents \( a \) and \( b \). Data-collapse means that the characteristic properties of the system represented by \( X \) are similar on different system size \( L \). Note that two systems of different sizes are said to be similar if they differ in the numerical value of their dimensional quantities \( X \) and \( x \), however, the numerical value of the corresponding dimensionless quantities \( XL^{-b} \) and \( (x - x_c)L^a \) coincide and that is why we obtain data-collapse. Obtaining data-collapse guarantees that the system exhibits scaling or similarity with respect to different independent system size. It is an extension of the idea of similarity of two triangles. For instance, two right triangles (characterized by their area \( S \) and the sides \( a, b \) and the hypotenuse \( c \)) may differ in the numerical value of their dimensional quantities. Now, one can vary \( b \) keeping \( a \) fixed and measure \( S \) for both the triangles. Plotting \( S \) as a function of \( b \) will definitely give two distinct curves one for each. However, the plots of the corresponding dimensionless quantities \( S/c^2 \) vs \( b/c \) will give rise to single universal curve since the numerical value of \( S/c^2 \) will always coincide for a given value of the acute angle \( \theta \) regardless of the size of the triangle. This happens because triangles are similar.

V. SITE/BOND PERCOLATION ON WPSL

What is site and bond in WPSL? Before answering this question we find it worth discussing first what they are in the context of conventional lattices. For instance, we can regard a square lattice as a grid or mesh. Each cell of the grid has four sides and each side is a common border of two cells only. In the case of square grid, we can thus regard each cell as a site since it contains exactly one lattice point. Equivalently, we could also regard the vertices of each cell as sites. However, in the present context we stick to the former definition. The dual of the square grid, obtained by replacing the center of each cell by a node and the common border between neighbouring cells by a link connecting the two nodes. We can thus regard the links of the dual as the bond of the square lattice.

Following the same argument we regard the blocks of the WPSL as its sites not the vertices of the lines that tessellated the initiator. To define bond, we first find its dual. It is obtained by replacing the center of each block by a node and the common border between two neighbouring blocks by a link connecting the corresponding nodes. We regard these links as the bonds of the WPSL. Using these ideas we first performed site and bond percolation on the square lattice and reproduced all the known results and then we applied them to the WPSL.

Recently, we have studied site percolation on WPSL, and found non-trivial results. That is, it belongs to a separate universality class than the universality class where percolation on all planar lattices are believed to belong. However, we are yet to check whether the site and bond percolation on WPSL belong to the same class or not. The dual of the WPSL can be well described as complex network and we have shown in Ref. [20] that the corresponding degree distribution follow a power-law \( P(k) \sim k^{-\gamma} \) with exponent \( \gamma = 5.58 \). Interestingly, the degree distribution \( P(k) \) in the context of network is the same as the coordination number distribution in the context of lattice. However, there is a sharp difference between networks based on graph theory and the network obtained from the dual of a lattice which is embedded in a space. The difference lies in the fact that networks based on graph theory have no edge or surface but networks based on the dual of a lattice have edge or surface which is crucial in the case of percolation as it is useful in defining the spanning cluster.

In the case of bond percolation, the lattice consists
initially of \( N \) blocks and hence the system has exactly \( N \) number of cluster of size one since the center of each block represents a site. Thereafter, each time we occupy a bond, a cluster at least of size two or more is formed. In the case of site percolation, each time we occupy a block, the size of the cluster may vary as we measure it by the area of contiguous occupied blocks. Initially all the blocks are empty and we won’t know the size of the cluster even after the first block is occupied. For regular lattice like square lattice of \( L^2 \) sites have \( 2L(L-1) \) and \( 2L^2 \) bonds with open and periodic boundary condition respectively. Now in the case of WPSL, being a disordered lattice, we cannot have such exact relation. We still find that the number of bonds or sites when we take average over ensemble of independent realizations follow a relation valid for all size of the lattice. For instance, for the lattice at time \( t \) there are exactly \( 3t + 1 \) sites and on the average there are \( 3t \) bonds with periodic boundary condition. Thus the mean coordination number is equal to \( 16t/3t \approx 5.33 \) which is higher than the square lattice. We know that the percolation threshold \( p_c \) depends on coordination number of the lattice and the higher the mean coordination number of a lattice the lesser is the value of \( p_c \). In the case of for square lattice, for instance, each site has exactly four nearest neighbours and each bond has six and hence \( p_c \) of site percolation is higher than that of the bond. In the case of WPSL, we find that the mean number of nearest neighbors of a bond is 10.01 which is almost double the mean nearest neighbour of a site. So, it is expected that the \( p_c \) value for bond percolation in WPSL will be quite less than \( p_c = 0.5265 \) for the site percolation [10].

Percolation is all about formation of clusters and the statistics of their various properties as a function of control parameter \( p \) and \( L \). The typical observable quantities in percolation are (i) Spanning probability \( W(p) \), (ii) percolation probability or percolation strength \( P \), (iii) The mean cluster size \( S \), (iv) cluster size distribution function \( n_s(p) \) etc and their variation with \( p \) or \( L \).

### A. Spanning Probability \( W(p) \)

The spanning probability \( W(p) \) for both bond and site describes the likelihood of finding a cluster that spans across the system either horizontally or vertically at the occupation probability \( p \). To find how \( W(p) \) behaves with the control parameter \( p \) we perform many, say \( M \), independent realizations under the same identical conditions. In each realization for a given finite system size we take record of the \( p_c \) value at which the spanning cluster appears for the first time. To find a regularity or a pattern among all the \( M \) numbers of \( p_c \) values recorded, one usually looks at the relative frequency of occurrence within a class or width \( \Delta p \). To find \( W(p) \), we can process the data containing \( M \) number of \( p_c \) values to plot histogram displaying normalized relative frequency as a function of class of width \( \Delta p \) chosen as per convenience. In Figs. [23] and [24] we show a set of plots of \( W(p) \) for bond and site percolation respectively as a function of \( p \) where distinct curves are for different system size \( L = \sqrt{N} \). One of the significant features of such plots is that they all meet at one particular \( p \) value regardless of the value of \( L \). It means that even if we had data for infinite system the resulting plot would still meet at the same point revealing that it must have a special significance and the significance is that it is the threshold probability \( p_c \). Note that finding the \( p_c \) value for different lattice is one of the central problems in percolation theory. In the case of bond we find \( p_c = 0.3457 \) which is exceedingly less than its site counterpart since on the average nearest neighbor that each bond has in the WPSL is much higher than for its site counterpart.

The second most significant feature of the \( W(p) \) vs \( p \) plot is the direction of shift of the curve on either side of \( p_c \). As the system size \( L \) increases. This shift with \( L \) clearly reveals that all the data points, i.e. the \( p \) values, are marching towards \( p_c \). We can quantify the extent at which they are marching by measuring the magnitude of the difference \((p_c - p)\) for different \( L \). That is, we can draw a horizontal line at a given value of \( W \), preferably at the position where this difference is the most, and take records of the difference \( p_c - p \) as a function of system size \( L \). Plotting the resulting data after taking log of both the variables or in the logarithmic scale we find a straight line whose slope gives an estimate of the inverse of \( 1/\nu = 0.613552 \pm 0.003861 \) since Fig. [24] suggests

\[
p_c - p \sim L^{-\nu}.
\]

\( (20) \)
It implies that in the limit $L \to \infty$ all the $p$ takes the value $p_c$ revealing that $W(p)$ will ultimately become a step function so that $W(p) = 0$ for $p \leq p_c$ and $W(p) = 1$ for $p > p_c$. We can use Eq. (20) to define a dimensionless quantity $(p_c - p)L^{\frac{\beta}{\nu}}$. Now, we plot $W(p)$ vs $(p_c - p)L^{\frac{\beta}{\nu}}$ in Fig. 24 and we see that all the distinct plots $W(p)$ vs $p$ for bond percolation collapse onto a one universal curve and for site onto another curve albeit they share the same $\nu$ value. By tuning the $1/\nu$ value further we can get an excellent data-collapse for $1/\nu = 0.6115$ and hence a better $\nu \sim 1.635$ value that corresponds to infinite lattice size.

**B. Percolation probability $P$**

Consider that we pick a site at random and ask: How likely is that site belong to the spanning cluster? For finite system size, it may not belong to the spanning cluster even if $p$ is larger than the percolation threshold $p_c$. Therefore, we can quantify the strength of the spanning cluster by percolation probability $P$ which describes how likely a site picked at random is to belong to the spanning cluster. The quantity $P$ is defined as the ratio of the size of the spanning cluster $s_\infty$ to the size of the lattice $N$ i.e.,

$$P = \frac{\text{Number of sites in the spanning cluster}}{\text{Total number of sites in the lattice}}.$$  

(21)

Sometimes, percolation probability is also defined as the probability that an occupied site belongs to the spanning cluster. It can be obtained if we replace the denominator $N$ of Eq. (21) by total occupied sites. We, however, will consider the former definition. There exists yet another definition where we can use the size of the largest cluster instead of the spanning cluster. Note that all of these definitions behaves in the same fashion like order parameter. That is, in the limit $L \to \infty$, $P = 0$ for $p \leq p_c$ and it rises from $P = 0$ at $p_c$ to $P = 1$ continuously and monotonically like $P \sim (p - p_c)^\beta$. Such behavior is reminiscent of order parameter like magnetization $m$ in the case of paramagnetic to ferromagnetic transition and hence $P$ is regarded as the order parameter in percolation theory. The critical exponent $\beta$ value is known to depend only on the dimension of the lattice and independent of the type of percolation. Through the site percolation on WPSL we already reported that $\beta$ value for WPSL, which is a planar lattice, is different from the value for all the known lattices whose dimension of the embedding space $d = 2$. We shall now check if the $\beta$ value for the bond percolation is the same as for the site percolation.

It is important to note that in the case of site percolation we occupy its blocks or cells which are of different size. We therefore measure the area of the spanning cluster, not the number of blocks in the spanning cluster. This is in sharp contrast to the regular lattice where all the blocks or cells are of the same size and hence the size of the spanning clusters can be described by the number of blocks or sites in the spanning cluster. In the case of bond percolation on WPSL we, however, use the traditional definition of cluster size. This is one significant difference between bond and site percolation on WPSL. Note that for bond percolation on WPSL we use the dual of the WPSL not the lattice itself. The dual of the WPSL is obtained by replacing each block of the WPSL by a node or vertex at its center and each common border between blocks by a bond connecting the nodes at the center of corresponding blocks. In the case of bond percolation we occupy these links and measure the size of the cluster by the number of nodes or vertices that the cluster contains. Below we shall see the impact of this difference in their behavior, if at all. In Figs. 3a and 3b we plot percolation probability $P$ as a function of $p$ for bond and site respectively. Looking at the plots, one may think that all the plots for different $L$ meet at a single unique point like it does for $W(p)$ vs $p$ plot. However, if one zoom in it becomes apparent it is not so and hence the $p_c$ value from this plot will not be as satisfactory as it is from $W(p)$ vs $p$ plot. We also find that $P(p)$ is not strictly equal to zero at $p < p_c$ rather there is always a non-zero chance of finding a spanning cluster even at $p < p_c$ as long as the system size $L$ is finite. However, the plots of $P$ vs $p$ for different system size $L$ reveals that the chances of getting spanning cluster at $p < p_c$ diminishes with increasing $L$. There is also a lateral shift of $P$ value to the left for $p > p_c$ but the extent of this shift $p - p_c$ decreases to such an extent that it never diminishes. On the other hand, the extent of shift $p - p_c$ to the right for $p < p_c$ diminishes to zero following Eq (20). We shall now check if $P$ above $p_c$ grows like $P \sim (p - p_c)^\beta$. If it does so then we shall find the value of the critical exponent $\beta$ and compare it with that of its site counterpart.

To show that the percolation probability behaves like $P \sim (p - p_c)^\beta$ and to find the exponent $\beta$ for infinite system size $L$ we use the idea of finite-size scaling. We first plot $P(p)$ vs $(p_c - p_c(L))L^{\frac{\beta}{\nu}}$ and find that unlike $W(p)$ vs $(p_c - p_c(L))L^{\frac{\beta}{\nu}}$ it does not collapse. Instead, we find that for a given value of $(p - p_c)L^{1/\nu}$ the $P$ value decreases with lattice size $L$. It means percolation probability is not a dimensionless quantity and hence assume that

$$P \sim L^{-a},$$

(22)

and we choose $a = \beta/\nu$ for later convenience. To find the value of $\beta/\nu$ we measure the heights at a given value of $(p - p_c)L^{1/\nu}$ for different $L$ and plot them in the log-log scale. We find straight lines for both bond and site (see Fig. 3b) with slopes $\beta/\nu = 0.135699 \pm 0.0005905$ for bond and $0.135701 \pm 0.0002768$ for site revealing that they are almost parallel. It implies that if we now plot $PL^{\beta/\nu}$ vs $(p - p_c)L^{1/\nu}$ all the distinct plots of $P$ vs $p$ should collapse into a single universal curve. In Fig. 3b we plot just that and find an excellent data-collapse using $\beta/\nu = 0.1357$ for both bond and site. We checked it for square lattice anyway. This again implies that perco-
dimensional system, we know an exact form for the cluster
the lattice. Unfortunately, only in the case of one di-

![Figure 3](image1.png)

**FIG. 3:** Percolation strength or percolation probability
$P(p, L)$ in WPSL for (a) bond and (b) site percolation. In (c) we plot log $P$ vs log $L$ using data for fixed value
of $(p - p_c)L^{1/\nu}$ and find almost parallel lines with slopes
$eta/\nu = 0.135699\pm0.0005905$ for bond and $0.135701\pm0.0002768$
for site respectively which clearly implies that the critical exponent
$\beta$ is independent of the type of percolation. For further
fine tuning of the $\beta$ value we also plot the same data of
(a) and (b) in the self-similar coordinates namely $PL^{\beta/\nu}$ and
$(p - p_c)L^{1/\nu}$ and find excellent data-collapse of the plots (a)
and (b) both using $\beta/\nu = 0.1357$ which gives $\beta \sim 0.222$

![Figure 4](image2.png)

**FIG. 4:** The mean cluster size $S(p, L)$ for (a) bond and (b) site
percolation as a function of $p$ for different size of the WPSL.
In the case of bond the cluster size is measured by the number
of sites each cluster contain and in the case of sites it is the
area of the contiguous blocks that belong to the same cluster.
In (c) we plot log $S$ vs log $L$ using the size of $S$ for fixed
value of $(p - p_c)L^{1/\nu}$ and find almost parallel lines with slope
$\gamma/\nu = 1.73153 \pm 0.001979$ and $1.72806 \pm 0.001993$ for bond
and site respectively. In order to obtain a better estimate
for the $\gamma$ value we also plot the same data of (a) and (b) in
the self-similar coordinates namely $PS^{-\gamma/\nu}$ and $(p - p_c)L^{1/\nu}$.
By tuning the $\gamma/\nu = 1.728$ value we find a set of excellent
data-collapse for both (a) and (b) that gives $\gamma = 2.825$.

infinite system which is actually the Bethe lattice. For
$1 < d < \infty$ we do not yet know an exact expression for $n_s$.
This is because in such cases there exists a large number of
different ways in which clusters of same size can ar-
range themselves, which are called lattice animals. Even
for relatively small cluster size in the square lattice we
run into difficulties in enumerating them. Nevertheless,
theoretically we can still write down the general expres-
sion

$$n_s(p) = \sum_{s,t} g_{s,t} p^s (1-p)^t,$$

where $g_{s,t}$ is the number of possible lattice configurations
of size $s$ and perimeter of size $t$. Note that the quantity
$n_s(p)$ is the probability that an arbitrary site belongs
to a cluster of size $s$. On the other hand, the quantity
$\sum_{s=1} \sum sn_s$ is the probability that an arbitrary site belongs
to a cluster of any size which is in fact equal to $p$. There-
fore, the ratio of the two

$$f_s = \frac{sn_s(p)}{\sum_{s=1} \sum sn_s},$$

is the probability that an occupied site chosen at random
belong to a cluster of size exactly equal to $s$. The mean
cluster size $S(p)$ therefore is given by

$$S(p) = \sum_s s f_s = \sum_s s^2 \sum sn_s,$$
where the sum is over the finite clusters only i.e., the
spanning cluster is excluded from the enumeration of $S$.
The definition of mean cluster size $S$, however, does not
have information about the geometric structure of the
clusters like their compactness and spatial extent. It is
important to mention that the mean area of the blocks in
the WPSL decreases as $(1 + 3t)^{-1}$ and hence increasing
the size of the lattice we need to blow up the lattice
by a factor of $3t$. It compensates the decreasing block
size with increasing block number $N$. That is, the mean
cluster size

$$ S = \frac{1}{p} \sum_s s^2 n_s \times 3t, \quad (28) $$

in the case of WPSL. In the case of bond percolation,
however, we do not need to multiply by the factor $3t$ as
the cluster size here is measured by the number of nodes
or vertices it contains not by the area.

In Figs. 4 and 5, we show the plots of the mean
cluster size $S(p)$, for both bond and site percolation, as
a function of $p$ for different lattice sizes $L$. We observe
that in either cases, there are two main effects as we increase
the lattice size. First, we see that the mean cluster size
increases as we increase the occupation probability till $p$
approaches to $p_c$ and the peak height grows profoundly
with $L$ in the vicinity of $p_c$. Second, there is a slight
shift in the peak towards $p_c$ value as we increase $L$.
The extent of shift is again given by Eq. (29). To bring
the peak height to meet at the same point we first plot
$S$ as a function of dimensionless quantity $(p_c - p)L^{1/\nu}$.
We then measure the peak height for a fixed value of
$(p_c - p)L^{1/\nu}$ but for different $L$. Plotting these peak
heights as a function of $L$ in the log-log scale give straight
lines for site and bond percolation both (see the inset of
Fig. 4d). It implies that

$$ S \sim L^\theta, \quad (29) $$

where like before we again choose $\theta = \gamma/\nu$ for future
convenience and find that $\gamma/\nu = 1.73153 \pm 0.001979$ for
bond and $1.72806 \pm 0.001993$ for site. The two values
are so close that they can be well approximated to be
the same. Plotting now the same data of Figs 4a and 4b
by measuring the mean cluster size $S$ in unit of $L^\theta$ and
$(p_c - p)$ in unit of $L^{-1/\nu}$ respectively we find that
all the distinct plots of $S$ vs $p$ collapse superbly into one
universal curve (see Fig. 4d) in both cases with the
same value for the corresponding exponents $\gamma/\nu = 1.728$.
It again implies that the mean cluster size too, for both
bond and site, exhibits finite-size scaling

$$ S \sim L^{\gamma/\nu} \phi((p_c - p)L^{1/\nu}), \quad (30) $$

sharing the same critical exponents. Eliminating $L$ from
Eq. (29) in favor of $(p_c - p)$ using $(p_c - p) \sim L^{-1/\nu}$ we
find that the mean cluster diverges

$$ S \sim (p_c - p)^{-\gamma}, \quad (31) $$

where $\gamma = 2.825$ for both site and bond percolation. This
value is significantly different from the known value $\gamma =
2.389$ for all the regular planar lattices.

![Fig. 5: We plot the cluster size distribution function log($n_s(p_c)$) vs log $s$ for different size of the WPSL. Once again we find almost parallel lines since slopes are 2.07252 and 2.0728 for bond and site percolation respectively which implies that the $\tau$ value is independent of the type of percolation.](image)

The mean cluster size $S$ according to Eq. (31) thus
diverges as we approach to the threshold value $p_c$ as expected.
On the other hand, $S$ can diverge, according to
Eq. (27), if $\sum_s s^n n_s$ diverges since denominator $\sum_s s^n$
in the same limit reaches to a constant $p_c$. Generally, we know that

$$ \sum_{s=1}^{\infty} s^\alpha = \begin{cases} 
\text{convergent} & \text{for } \alpha < -1 \\
\text{divergent} & \text{if } \alpha \geq -1,
\end{cases} \quad (32) $$

and hence we can use it to find out under what condition
the numerator of Eq. (27) diverges. It is convenient to assume

$$ n_s(p) \sim s^{-\tau} \phi((p - p_c)^{1/\sigma} s), \quad (33) $$

which means $n_s(p_c) \sim s^{-\tau}$ and hence

$$ \sum_{s=1}^{\infty} s^{2n} n_s(p_c) \sim \sum_{s=1}^{\infty} s^{2-\tau}. \quad (34) $$

It implies that $S$ would diverge as $p \to p_c$ if $(2 - \tau) \geq -1$
or $\tau \leq 3$. On the other hand, we also demand $p_c \sim$
$\sum_s n_s(p_c)$ implies $\sum_{s=1}^{\infty} s^{1/\tau} n_s(p_c) \sim \sum_{s=1}^{\infty} s^{1-\tau}$. It implies that $(1 - \tau) < -1$ or $\tau > 3$. Putting the two
constraints together we find that $\tau$ must satisfy the bound
$2 < \tau \leq 3$. We can thus write that

$$ n_s(p_c) \sim s^{1-\tau} \quad (35) $$

where $\tau$ is called the Fisher exponent. We can obtain
the exponent $\tau$ by plotting the cluster size distribution
function $n_s(p_c)$ at $p_c$. In Fig. 5 we plot $n_s(p_c)$ vs $s$, for
both site and bond, in the log-log scale and find two
parallel lines except near the tail where there is a hump.
due to finite size effect. However, we also observe that as the lattice size \( L \) increases the extent up to which we get a straight line increases too. It implies that if the size \( L \) were infinitely large, we would have a perfect straight line obeying Eq. (35). The slopes of the lines are \( \tau = 2.07252 \) for bond and \( \tau = 2.0728 \) for site. It implies that the exponent \( \tau \) is almost the same \( \sim 2.072 \) for both site and bond percolation on WPSL and its value is different than the known value for all known planar lattices \( \tau = 2.0549 \). 

\[
\ln(M) = D \ln(L)
\]

with \( D = 2 \). Now, let us slightly change the situation. We do everything like before with the only difference is that at each step we throw the top right copy leaving its space empty as shown in Fig. (6b). The amount of mass of the resulting system in the \( i \)th step is \( M = 3^i \) and the linear size of the system is \( L = 2^i \). Using this two relations we can eliminate \( i \) in favor of \( L \) and we find the same mass-length relation as in Eq. (36) except that we get exponent \( D = \ln 3/\ln 2 \) \cite{22}. We could even remove any of the four copies at random and still we would get the same result. The exponent of the mass-length relation \( D = d_f \) which is now less than the dimension of the embedding space \( d = 2 \) and hence it is a fractal. The spanning cluster too is highly ramified like Fig. (6b) as it has holes of many different sizes. Now, a litmus test whether the spanning cluster is a fractal or not would be to check if it obeys the same mass-length relation with an exponent \( d_f < 2 \) since the embedding space of the spanning cluster is a plane. We plot the size of the spanning cluster \( M \) as a function of lattice size \( L \) in the log-log scale as shown in Fig. (7). Indeed, we find that \( d_f = 1.86439 \pm 0.001498 \) for site and 1.86378 \pm 0.02249 which are almost the same but significantly different from the one for regular planar lattices \( d_f = 1.895 \). It may appear that the difference between the \( d_f \) for WPSL and that for regular planar lattices is not much but it important to remember that even a small difference in fractal dimension has a huge impact in their degree of ramification.

We already know that the mean cluster size diverges i.e., \( S \rightarrow \infty \) as \( p \rightarrow p_c \). According to Eq. (27), \( S \) can only diverge if its numerator diverges. Generally, we know that \( \sum_{n=1}^{\infty} s^n \) converges if \( \alpha < -1 \) and diverges if \( \alpha \geq -1 \).
Applying it into both numerator and denominator of Eq. (27) at \( p_c \) gives a bound that \( 2 < \tau < 3 \). Using Eq. (33) in Eq. (27) and taking continuum limit gives

\[
S \sim s_\xi ^{3-\tau}.
\]

(37)

We know that \( s_\xi \) diverges like \((p_c - p)^{-1/\sigma}\) where \( \sigma = 1/(\nu d_f) \) and hence comparing it with Eq. (31) we get

\[
\tau = 3 - \gamma \sigma.
\]

(38)

Besides, there is another well known scaling relation \( \tau = 1 + d/d_f \) which we can use to find \( \tau \) value. Using the \( d_f \) value for WPSL in the scaling relations, \( \tau = 3 - \gamma \sigma \) and \( \tau = 1 + d/d_f \), we find \( \tau \) equal to 2.0725 and 2.0728 respectively which is almost equal to the one we obtained straight from slope of Fig. 7. There are also a couple of other well-known scaling relations, such as \( \beta = \nu(2d_f - d) \), \( \gamma = \nu(2d_f - d) \), which we used for a consistency check of our results. To this end, we find that our estimates satisfy these relations up to a quite good extent.

| Exponents   | regular 2d lattice | WPSL bond/site |
|-------------|---------------------|----------------|
| \( \nu \)   | 1.75                | 1.635          |
| \( \beta \)  | 0.13889             | 0.222          |
| \( \gamma \) | 2.3889              | 2.825          |
| \( \tau \)  | 2.0749              | 2.0728         |
| \( d_f \)   | 1.895(8)            | 1.864          |

TABLE I: The critical and other characteristic exponents for site and bond percolation in the WPSL and in the regular planar lattice are given alongside.

VI. SUMMARY AND DISCUSSION

In this article, we have studied both bond and site percolation on WPSL using extensive Monte Carlo simulations. We thought it is important to know some key features of the WPSL so that one can understand why it is so special and unique. We therefore have first briefly discussed its construction process and then its various properties which are as follows. (i) The dynamics of its growth is governed by infinitely many conservation laws. (ii) Its area size distribution function obeys dynamic scaling. (iii) Each of the infinitely many conservation laws, except conservation of total area, gives rise to multifractal spectrum and hence WPSL is a multi-multifractal. Fourth, its coordination number distribution function follows a power-law. (iv) It has a mixture of properties of both lattice and graph. On one hand, like lattice, it is embedded in a space of dimension \( D = 2 \); On the other its coordination number distribution follow power-law like network. These unique properties have resulted in unique results too. We also briefly discussed about the finite-size scaling theory and have shown that its origin is deeply rooted to the Buckingham II-theorem. The finite-size scaling is one of the most crucial aspects in percolation as it helps extrapolating critical exponents for infinite system using data for a set of finite size systems. This is done by using the idea of data collapse. Note that an excellent data collapse is one of the clear testaments that the numerical values we obtained for various exponents are quite satisfactory. Besides, we show that these satisfy a set of scaling relations which also provide a consistency check.

In this work we first obtained percolation threshold \( p_c = 0.3457 \) and \( p_c = 0.5265 \) for bond and site percolation on WPSL. Naturally, the \( p_c \) for bond is less than that of its site counterpart as expected. We also obtained numerically the various observable quantities such as the spanning probability \( W(p) \), the percolation strength \( P(p) \), the mean cluster size \( S(p) \) etc. using NZ algorithm. The initial data obtained from the NZ algorithm correspond to microcanonical ensemble. To get the corresponding data that correspond to canonical ensemble we used the convolution equation given by Eq. (10) for each observable quantities. With the help of a comprehensive finite-size scaling theory we also obtained numerically the critical exponents \( \nu, \beta \) and \( \gamma \) for both bond and site percolation on WPSL and confirm they are equal (see table IV for detailed comparison). To check further if they are equal or not we used the idea of data collapse and found an excellent data collapse for the same critical exponents albeit different \( p_c \). Note that good estimate of \( p_c \) and of the critical exponents a must for obtaining satisfactory data collapse. These values also satisfy the scaling relations. All these provide a clear testament that the critical exponents for bond and site percolation in WPSL are the same. It happens in spite of the significant difference in the definition of clusters. Interestingly, these values are significantly different from the ones for all known planar lattices. We can thus conclude that the universality class of WPSL (bond and site) is distinct from the ones for all the known planar lattices. It happens in spite of the significant differences in the definition of site and bond in the WPSL.

Hsu and Huang also studied percolation in a class of random planar lattices and their duals yet they found the same critical exponents as the ones for regular lattices. Corso et. al. studied percolation on multifractal planar lattices and they too found the same critical exponents as the ones on regular lattices. So, it is neither the randomness nature of the lattice nor the multifractal nature of the lattice can be held responsible for making WPSL unique. The planar random lattice that Hsu and Huang studied is quite different than WPSL. The coordination number distribution of their lattice do not obey power-law. This is perhaps one of the most significant differences. Or, it may be the case that when a lattice is multifractal and at the same time it is random then depending on further detailed nature may be responsible for giving a new set of exponents. However, it is too soon to draw any conclusion. We hope to devise more variants of WPSL in our future endeavour and see what happens. Nevertheless, we still hope that our findings will have a
significant impact in the percolation theory.

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