Native ultrametricity of sparse random ensembles

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We investigate the eigenvalue density in ensembles of large sparse Bernoulli random matrices. We demonstrate that the fraction of linear subgraphs just below the percolation threshold is about 95% of all finite subgraphs, and the distribution of linear chains is purely exponential. We analyze in detail the spectral density of ensembles of linear subgraphs, discuss its ultrametric nature and show that near the spectrum boundary, the tail of the spectral density exhibits a Lifshitz singularity typical for Anderson localization. We also discuss an intriguing connection of the spectral density to the Dedekind \( \eta \)-function. We conjecture that ultrametricity is inherit to complex systems with extremal sparse statistics and argue that a number-theoretic ultrametricity emerges in any rare-event statistics.

I. INTRODUCTION

A. Some notions about ultrametricity

The concept of ultrametricity is related to a special class of metrics. Generally, a metric space is a set of elements equipped by pairwise distances. The metric \( d(x, y) \) meets three requirements: i) non-negativity, \( d(x, y) > 0 \) for \( x \neq y \), and \( d(x, y) = 0 \) for \( x = y \), ii) symmetry, \( d(x, y) = d(y, x) \) and iii) the triangle inequality, \( d(x, z) \leq d(x, y) + d(y, z) \). Ultrametric spaces obey the strong triangular inequality, \( d(x, z) \leq \max\{d(x, y), d(y, z)\} \), allowing only acute isosceles and equilateral triangles.

The strong triangle inequality radically changes the properties of spaces in comparison with our intuitive expectations based of Euclidean geometry. In ultrametric spaces, the sum of distances does not exceed the largest summand, i.e. the Archimedean principle does not hold. Few counterintuitive properties are related to ultrametric balls: Every point inside a ball is its center; the radius of a ball is equal to its diameter; there are no balls that overlap only partially, if two balls have a common point, then one of them is completely embedded into the other. The last property implies that any ultrametric ball can be divided into a set of smaller balls, each of them can be divided into even smaller ones, etc. In other words, an ultrametric space resembles not a line of segments, but a branching tree of hierarchically nested balls (“basins”). The terminal points of a tree, i.e. the tree boundary, also obey the strong triangle inequality: for any two such points, A and B, the distance to the root of
the minimal subtree to which these points belong, determines the distance between A and B. As a result, ultrametric geometry, in general, fixes taxonomic (i.e. hierarchical) tree-like relationships between the elements.

Ultrametricity is closely related to the $p$-adic numerical norm, which originally arose in the context of number theory \[1\]. The role of ultrametric spaces and $p$-adic numbers in number theory \[2, 3\] and in other branches of mathematics, such as $p$-adic analysis \[4\] and non-Archimedean analytic geometry \[5\], is growing. More recently, there have been numerous attempts to apply $p$-adic numbers and ultrametric spaces to various branches of physics ranging from string theory to inflationary cosmology (see e.g. \[6, 7\]).

Perhaps most naturally ultrametric spaces emerge in the realm of complex systems. Indeed, the ultrametric description is inherently multi-scale, thus it is not surprising that ultrametricity provides a fertile framework for describing complex systems characterized by a large number of order parameters. For instance, signatures of ultrametricity were discovered in the context of spin glasses (see \[8\] for an earlier review). Spin glasses are characterized by a huge number of metastable states and the equilibration in spin glasses proceeds via tree-like splitting of the phase space into hierarchically nested domains. When the temperature decreases, the most distant groups of spin-glass states get factorized first. Then, each of these groups splits into a number of subgroups, etc. The scales of hierarchically nested sets of equilibrated states satisfy the strong triangle inequality, i.e. the multi-scale splitting of spin-glass phases obey ultrametric relations.

The physical origin behind the emergence of ultrametricity in spin-glass systems is quite general. In a complex system with an extremely large number of metastable states, the phase trajectories cannot totally explore all states: they cover only a tiny part of phase space almost without returns to already visited regions. For this reason, the factorization of spin-glass phases is similar to random branching in a high-dimensional (infinitely-dimensional in the thermodynamic limit) space. The factorization of spin-glass phases is specified only by branching points because once two branches are dispersed in a high-dimensional space, they never join. The ultrametric ansatz \[13, 14\] proposed to describe the spin-glasses by infinite (in the thermodynamic limit) number of order parameters, was proven in \[15\] to be an exact ground state of the Sherrington-Kirkpatrick model \[16\].

Shortly after the formulation of the ultrametric ansatz in the context of spin glasses, similar ideas were proposed for the description of native states of protein molecules \[17, 18\]. Proteins are highly frustrated polymer systems characterized by rugged energy landscapes of very high dimensionality. The ultrametricity of proteins implies a multi-scale representation of the protein energy landscape by means of ranging the local minima into hierarchically embedded basins of minima. Ultrametricity follows from the conjecture that the equilibration time within the basin is significantly smaller than the time to escape the basin. As a result, the transitions between local minima obey the strong triangle inequality. In such an approximation, dynamics on a complex energy landscape can be modelled via a jump-like random process, propagating in an ultrametric space of states. The approximation of the protein energy landscape by a self-similarly branching tree of basins has been shown to be very fruitful for describing a large body of experimentally justified features of protein conformational dynamics and protein fluctuation mobility at temperatures covering very wide range from room temperatures up to the deeply frozen states \[19\].

With respect to ultrametricity and the self-similarity of protein energy landscapes, it is important to keep in mind that proteins are functional systems. Proteins precisely operate
at the atomic level using sparse-event statistics, which is a bridge between non-local protein dynamics in high-dimensional conformational space and low-dimensional movements of particular atomic units related to protein functionality.

B. Ultrametricity and rare-event statistics

In the context of high dimensionality, randomness, and rare-event statistics, ultrametricity emerges in diverse subjects, e.g., in genetic trees, in DNA folding (or crumpling) in chromosomes, in the behavior of hierarchically organized ecological, social, and economic systems, etc. The fact that ultrametricity is naturally rooted in high dimensionality, randomness, and sparse statistics, has been unambiguously observed recently in [20]. It was proven that in a $D$-dimensional Euclidean space the distances between points in a highly sparse sampling tends to the ultrametric distances as $D \to \infty$.

In this work we bring another intriguing connection between rare-event statistics and ultrametricity. We demonstrate that ultrametricity is inherit for ensembles of sparse topological graphs. In particular, comparing a limiting behavior of spectral density of bi-diagonal random matrices, with the behavior of the function $f(z) = \sqrt{-\log|\eta(z)|}$, where $\eta(z)$ is the Dedekind $\eta$-function, and $z$ tends to rational points on real axis, we conjecture that the eigenvalue distribution of sparse ensembles shares some number-theoretic properties related to ultrametricity.

Our finding provides an example of a nontrivial manifestation of rare-event Bernoullian noise in some collective observables in large systems (like spectral density, for example). From this point of view, our work should warn researchers working with effects of ultra-low doses of chemicals: in the statistical analysis of, say, absorption spectra of extremely diluted solutions of chain-like polymer molecules, the valuable signal should be clearly purified from the background noise, which itself could have a very peculiar ultrametric shape.

The paper is organized as follows. In the Section II we formulate the model under consideration and describe the results of our numerical simulations. In what follows we pay attention to statistics of linear subgraphs, presented by bi-diagonal matrices. Specifically, we derive the distribution of the fraction of linear subgraphs at the percolation threshold in the Section III, and study the spectral properties of Bernoulli ensembles of bi-diagonal matrices in the Section IV. The number-theoretic properties of limiting spectral statistics in ensembles of random bi-diagonal matrices are discussed in the Section V. The results of the work and the relation with other natural systems possessing the number-theoretic properties under consideration, are summarized in the Discussion.

II. THE MODEL

The Bernoulli matrix model is defined as follows. Take a large $N \times N$ symmetric matrix $A$. The matrix elements, $a_{ij} = a_{ji}$, are assumed to be independent identically distributed binary random variables such that non-diagonal elements are equal to one with probability
$q$, while diagonal elements vanish. Thus, $a_{ii} = 0$, and for $i \neq j$ one has

$$a_{ij} = \begin{cases} 1 & \text{with probability } q \\ 0 & \text{with probability } 1 - q \end{cases}$$

The matrix $A$ can be regarded as an adjacency matrix of a random Erdős-Rényi graph, $\mathcal{G}$, without self-connections. The eigenvalues, $\lambda_n$ ($n = 1, ..., N$), of the symmetric matrix $A$ are real. Let $\rho(\lambda)$ be the eigenvalue density of the ensemble of such matrices. For $N \gg 1$ the limiting shape of $\rho(\lambda)$ is known in various cases. If $q = O(1)$, the function $\rho(\lambda)$ tends to the Wigner semicircle law, $\sqrt{4N - \lambda^2}$, typical for the Gaussian matrix ensembles. For $q = c/N$ ($c > 1$), the matrix $A$ is sparse and the density $\rho(\lambda)$ in the ensemble of sparse matrices has singularities at finite values of $c$ [21–23]. In Refs. [24–27], the behavior of the spectral density, $\rho(\lambda)$, has been analyzed near $c = 1$ ($N \gg 1$). It has been pointed out that the function $\rho(\lambda)$ becomes more and more singular as $c$ approaches 1 from above.

For ensemble of matrices $A$ the percolation transition occurs at the value $q_c = 1/N$. (The true transition occurs, certainly, in the thermodynamic limit, $N \to \infty$.) For $q > q_c$, the graph $\mathcal{G}$ has a giant component whose size is proportional to $N$, and numerous small components (almost all of them are trees); for $q < q_c$, the graph $\mathcal{G}$ is the collection of small components (almost all of them are again trees). For $q \ll q_c$ the isolated vertices dominate, contributing to the trivial spectral density, $\rho(\lambda) = \delta(\lambda)$. The most interesting region is close to $q_c$. As long as $|q - q_c| = O(N^{-\alpha/2})$, the giant component is still absent. In what follows we discuss the behavior of the spectral density $\rho(\lambda)$ at $q$ in this region; more precisely, we usually chose $q$ slightly below $q_c$.

An example of a random graph with a randomly generated $N \times N = 500 \times 500$ adjacency matrix at $q = 2.0028 \times 10^{-3}$ is shown in Fig. 1. Note that the components are predominantly linear subgraphs. In other realizations we observed components with a single loop, but they are rare, usually a few per realization; more complicated components, namely those with more than one loop, appear very rarely.

The whole spectrum of a symmetric matrix $A$ in a sparse regime below the percolation point consists of singular peaks coming from the graph components (maximal disjoint subgraphs constituting the random graph). For a subgraph of size $k$, the corresponding adjacency matrix $A_k$ of size $k \times k$ and the spectrum is determined by the equation $\det(A_k - \lambda I_k) = 0$, where $I_k$ is the $k \times k$ identity matrix. There are $k$ eigenvalues; apart from $k$, the eigenvalues depend on the topology of the subgraph. The heights of peaks are the eigenvalue multiplicities, the eigenvalue $\lambda = 0$ appears with the highest multiplicity. For instance, the graph shown in Fig. 1 has 222 subgraphs of size one (isolated vertices) each of them contributing to $\lambda = 0$. There are also 10 linear chains of length three, 3 chains of size five, and 1 chain of size seven contributing to $\lambda = 0$; generally any chain of odd length has eigenvalue 0. Trees which are not linear chains also can have eigenvalue 0. One can bipartite any tree into two disjoint parts $X_1$ and $X_2$ with edges only between $X_1$ and $X_2$. If the sizes $n_1 = |X_1|$ and $n_2 = |X_2|$ of the parts are different, say $n_1 > n_2$, the adjacency matrix has eigenvalue 0 with multiplicity at least $n_1 - n_2$.

The singular spectral density, $\rho(\lambda)$, for an ensemble of 1000 random symmetric matrices $A$, each of size $500 \times 500$ and generated with probability $q = 2.0028 \times 10^{-3}$, is shown in Fig. 2a. The spectrum possesses the regular ultrametric hierarchical structure which becomes even more profound while plotting $\ln \rho(\lambda)$, as shown in Fig. 2b. Near the value
Figure 1: Typical sample of components constituting a random graph associated with an adjacency matrix $A$ generated by the rule (1); the parameters are $N = 500$ and $q = 2.0028 \times 10^{-3}$.

$\lambda = \pm 2$ the enveloping curve of the function $\ln \rho(\lambda)$ exhibits a fracture, clearly visible in the insert of Fig. 2a, where the spectrum is replotted as $\rho^{1/3}(\lambda)$ and the slopes are depicted by the dashed lines $s_1$ and $s_2$. (The choice of the exponent 1/3 for the curve $\rho^{1/3}(\lambda)$ is an experimental selection: we have just noted that such a scaling gives the best linear shape for the enveloping curve and clearly demonstrates the fracture of the shape near $\lambda = \pm 2$.)

The largest eigenvalue of a tree can be roughly estimated as $|\lambda_{\text{tree}}^{\text{max}}| = 2\sqrt{p - 1}$, where $p$ is the maximal vertex degree of a tree (see e.g. [28,31]). For linear chains, the maximal degree is $p = 2$ (when the chain length is at least three) and the maximal eigenvalue is $|\lambda_{\text{lin}}^{\text{max}}| = 2$.

In the bulk of the spectrum of sparse ensemble, the randomly branching trees (with $p = 3$) and linear chains (with $p = 2$) dominate (some estimates are given in the following sections). For $|\lambda| > |\lambda_{\text{lin}}^{\text{max}}| = 2$, the linear chains do not contribute to the spectral density, and only graphs with $p \geq 3$ remain. For Erdős-Rényi random graphs, trees with $p \geq 4$ are rather rare,
so the dominant contribution comes from graphs with $p = 3$, giving $|\lambda_{\text{tree}}^{\text{max}}| = 2\sqrt{2} \approx 2.83$, consistent with simulation results (Fig. 2b).

III. DISTRIBUTION OF LINEAR CHAINS AT THE PERCOLATION THRESHOLD

The spectral analysis of ensemble of random adjacency matrices $A$ at the percolation threshold is cumbersome since the spectra of the components depend on their topology. For random Erdős-Rényi graphs, however, almost all components are trees (see e.g. [32, 33]). Furthermore, at the percolation point (and certainly below it) the majority of subgraphs (about 95% of them) are linear chains as we show below. The spectrum of each linear chain is easy to compute, so to get the spectral density, $\rho_{\text{lin}}(\lambda)$, of an ensemble of chain-like subgraphs we should know the distribution, $Q_n$, of linear chains of length $n$ at the percolation point. Below we derive an explicit expression for $Q_n$. We first recall a kinetic theory approach which allows one to determine the size distribution of generic clusters (irrespective of their topology) and then adopt this framework to the computation of the size distribution of linear chains.

A. General formalism: Cluster size distribution

By definition, each cluster is a maximal connected component (see Fig. 3). To determine the cluster size distribution we employ a dynamical framework (see e.g. [33]), namely we
begin with a disjoint graph represented by a collection of \( N \) separated vertices and start linking them with a constant rate. When two vertices from distinct clusters become linked, the clusters join. For example, the latest link in Fig. 3 joins two clusters of size \( i = 2 \) and \( j = 4 \) into a cluster of size \( k = i + j = 6 \).

Generally, there are \( i \times j \) ways to join distinct clusters of size \( i \) and \( j \). Hence, the clusters undergo an aggregation process

\[
(i, j) \xrightarrow{ij/(2N)} i + j
\]

which proceeds with a rate proportional to the product of their sizes. We want to determine the cluster size distribution, \( c_k(t) \). By definition, \( c_k(t) = N_k(t)/N \), where \( N_k(t) \) is the total number of components with \( k \) nodes. In (2) the linking rate is chosen to be \( (2N)^{-1} \). With this choice, the cluster size distribution \( c_k(t) \) satisfies

\[
\frac{dc_k}{dt} = \frac{1}{2} \sum_{i+j=k} (ic_i)(jc_j) - kc_k
\]

The initial condition is \( c_k(0) = \delta_{k,1} \). The gain term in (3) accounts for clusters generated by joining two smaller clusters whose sizes sum up to \( k \). The second term on the right-hand side of Eq. (3) represents loss due to linking of clusters of size \( k \) to other clusters. The corresponding gain and loss rates follow from the aggregation rule (2). Stopping the aggregation process at time \( t \) leads to a random graph with adjacency matrix characterized by the connectivity probability \( q = t/N \).

![Figure 3: An evolving random graph with \( N = 9 \). Links are indicated by solid lines. The graph is composed of three clusters: Two clusters are chains (of length 2 and 3), the third cluster is a star of size 4. When the link shown by a dashed line is added, the chain of length 2 and the star merge.](image)

Equations (3) are well-known and they can be solved via various techniques [33]. Here we recall one approach as we shall use it in the next sub-section to determine the distribution of linear chains. The idea is to eliminate the time dependence. Solving (3) recursively one gets

\[
c_1(t) = e^{-t}; \quad c_2(t) = \frac{1}{2} t e^{-2t}; \quad c_3(t) = \frac{1}{2} t^2 e^{-3t}; \quad c_3(t) = \frac{2}{3} t^3 e^{-4t}; \quad \text{etc}
\]

This suggests to seek the solution in the form

\[
c_k(t) = C_k t^{k-1} e^{-kt}
\]
Substituting this form into (3), we find that the coefficients $C_k$ can be obtained recursively starting from $C_1 = 1$ and using

$$(k - 1) C_k = \frac{1}{2} \sum_{i+j=k} (iC_i) (jC_j)$$  \hspace{1cm} (6)$$

for $k > 1$. Using a generating function technique one gets $C_k = \frac{k^{k-2}}{k!}$. Thus

$$c_k(t) = \frac{k^{k-2}}{k!} t^{k-1} e^{-kt}$$  \hspace{1cm} (7)$$

The large $k$ tail is always exponential apart from the percolation point $t = t_c = 1$ where

$$c_k(1) \simeq (2\pi)^{-1/2} k^{-5/2} \quad \text{when} \quad k \gg 1$$  \hspace{1cm} (8)$$

**B. Distribution of linear chains**

Consider now linear clusters (open chains). Let $Q_k(t)$ be the density of linear clusters of length $k \geq 2$. We have $Q_2 = c_2$, and for $k \geq 3$ the chain size distribution evolves according to

$$\frac{dQ_k}{dt} = 2c_1 Q_{k-1} + 2 \sum_{i+j=k} Q_i Q_j - k Q_k$$  \hspace{1cm} (9)$$

where the summation is taken over $i \geq 2, j \geq 2$. It is convenient to define $Q_1 = \frac{1}{2} c_1$. This allows us to rewrite (9) as

$$\frac{dQ_k}{dt} = 2 \sum_{i+j=k} Q_i Q_j - k Q_k$$  \hspace{1cm} (10)$$

here the summation runs over all $i, j$ satisfying $i + j = k$. Another virtue of the agreement $Q_1 = \frac{1}{2} c_1$ is that (10) is valid for all $k \geq 2$. Computing the first few densities we get

$$Q_1 = \frac{1}{2} c_1 = \frac{1}{2} e^{-t}; \quad Q_2 = c_2 = \frac{1}{2} t e^{-2t}; \quad Q_3 = c_3 = \frac{1}{2} t^2 e^{-3t}; \quad Q_3 = \frac{1}{2} t^3 e^{-4t}$$  \hspace{1cm} (11)$$

etc (we have used (10) to determine $Q_4$). One is led to the conjectural behavior

$$Q_k(t) = \frac{1}{2} t^{k-1} e^{-kt}$$  \hspace{1cm} (12)$$

This indeed provides the solution to (10) and the initial condition $Q_k(0) = \frac{1}{2} \delta_{k,1}$.

Thus, the linear chain length distribution (12) is purely exponential. Rewriting (12) as

$$Q_k(t) = \frac{1}{2t} e^{-k(t-\ln t)}$$  \hspace{1cm} (13)$$

we see that the least-stepest decay occurs at the percolation point $t = t_c = 1$ where

$$Q_k \equiv Q_k(1) = \frac{1}{2} e^{-k}$$  \hspace{1cm} (14)$$
The total chain density at the percolation point is

\[ c_1 + \sum_{j \geq 2} Q_j(1) = \frac{1}{2} \left( 2 - e^{-1} \right) e - 1 = 0.474928 \ldots \] (15)

Comparing this result with the total cluster density at the percolation point, \( \sum_{k \geq 1} c_k(1) = \frac{1}{2} \), we conclude that at the percolation point almost 95% of clusters are linear chains. This justifies the abundance of linear chains (see Fig. 1).

Note that the total mass density of linear chains at the percolation point,

\[ c_1 + \sum_{j \geq 2} jQ_j(1) = \frac{1}{2e} \left[ 1 + (1 - e^{-1})^{-2} \right] = 0.6442765 \ldots , \] (16)

is significantly smaller than unity. Large clusters are usually not linear and hence despite of their rareness they provide a notable contribution to the mass density.

C. Comments on more complicated components and different random graph models

Below the percolation point, the total number of components which are not trees remains finite in the thermodynamic limit \( N \to \infty \), so it greatly fluctuates from realization to realization. The components which are not trees are unicyclic: each such component contains one loop. At the percolation point the total number of unicyclic components diverges but very slowly, viz. as \( \frac{1}{6} \ln N \). These results were established using probabilistic techniques, see [32]; they are also easily derivable using above dynamical approach (see [34]). Overall we see that the contribution of the non-trees is negligible.

The influence of the "shape" of the tree on the spectra is more notable. We have shown, however, that the majority of the clusters are not merely trees, but linear chains. In principle, one can modify the definition [1] to ensure that all components are chains. If each row (and column) of the adjacency matrix \( A \) has no more than two non-zero elements, \( \sum_{j=1}^{N} a_{ij} \leq 2 \) for \( i = 1, \ldots, N \), the corresponding graph is obviously the collection of linear chains (including loops) of various lengths.

Equivalently, the chain model can be defined dynamically by using the same procedure as before and checking that the constraint is obeyed. In other words, we attempt to link vertices randomly, but we actually draw a link only if the degrees of both vertices are smaller than 2. The cluster size distribution characterizing the chain model evolves according to master equation

\[ \frac{dc_k}{dt} = \frac{1}{2} \sum_{i+j=k} K_{ij} c_i c_j - c_k \sum_{j \geq 1} K_{kj} c_j \] (17)

with reaction rate matrix

\[
K_{ij} = \begin{pmatrix}
1 & 2 & 2 & 2 & \cdots \\
2 & 4 & 4 & 4 & \\
2 & 4 & 4 & 4 & \\
2 & 4 & 4 & 4 & \\
\vdots & & & & \ddots
\end{pmatrix}
\] (18)
The initial condition is $c_k(0) = \delta_{k,1}$. A more general model with reaction rate matrix of the type (18) composed of three arbitrary rates has been studied in [35].

Using (17)–(18) one finds that the density of isolated vertices $c_1(t)$ and the total cluster density $c(t) = \sum_{j \geq 1} c_j(t)$ satisfy a pair of coupled rate equations:

$$\frac{dc_1}{dt} = -c_1(2c - c_1), \quad \frac{dc}{dt} = -\frac{1}{2}(2c - c_1)^2 \quad (19)$$

The initial condition becomes $c(0) = c_1(0) = 1$. Treating $c$ as a function of $c_1$ we recast (19) into

$$c = c_1 \left[1 - \frac{1}{2} \ln c_1 \right] \quad (20)$$

Plugging this into the first equation (19) we obtain an implicit solution

$$\int_{c_1}^{1} \frac{du}{u^2(1 - \ln u)} = t \quad (21)$$

The following densities can be found by re-writing the rate equations (17) in a manifestly recursive form,

$$\frac{dc_2}{dt} = \frac{1}{2}c_1^2 - 2c_2(2c - c_1),$$

$$\frac{dc_3}{dt} = 2c_1c_2 - 2c_3(2c - c_1),$$

$$\frac{dc_4}{dt} = 2c_1c_3 + 2c_2^2 - 2c_4(2c - c_1),$$

$$\frac{dc_5}{dt} = 2c_1c_4 + 4c_2c_3 - 2c_5(2c - c_1),$$

etc., and solving them recursively.

We use the notation $c_k$, but we emphasize that clusters generated by the above procedure are linear chains. There is no percolation transition in this model, so when should we stop the process? Recall that for Erdős-Rényi random graphs, the percolation occurs when the average degree is equal to one. In the model (17)–(18), the fraction of vertices of degree zero ($d_0$), vertices of degree one ($d_1$), and vertices of degree two ($d_2$) are

$$d_0 = c_1$$

$$d_1 = 2 \sum_{j \geq 2} c_j = 2(c - c_1)$$

$$d_2 = 1 - d_0 - d_1 = 1 - c_1 - 2(c - c_1)$$

The average degree $\langle d \rangle = d_1 + 2d_2$ is therefore $\langle d \rangle = 2(1 - c)$. Thus, in analogy with Erdős-Rényi random graphs, we stop the process when the average degree reaches unity, the corresponding time $t_s$ is found from $c(t_s) = \frac{1}{2}$. Another choice is to run the process till the very end when the system turns into a collection of closed loops, see [36] for this and other models of ring formation.

One can generalize above model by requiring that each row (and column) of the adjacency matrix $A$ has no more than $p$ non-zero elements: $\sum_{j=1}^{N} a_{ij} \leq p$ for $i = 1, ..., N$. This model can be investigated in the dynamic framework. Unfortunately, it has proved to be much more difficult for the analysis [37] than the extreme versions corresponding to the chain model ($p = 2$) and the Erdős-Rényi variant ($p = N - 1$).
IV. SPECTRAL DENSITY OF CANONICAL ENSEMBLE OF BI-DIAGONAL RANDOM MATRICES

A. General formalism: Spectral density

We have seen in the Section III that the fraction of linear clusters of various lengths is about 95% at the percolation threshold in the ensemble of all sparse graphs. This justifies our study of ensembles of linear clusters (open chain and loops) only.

It is convenient to represent the linear cluster ensemble at the percolation point (with the distribution $Q_k$ in chain lengths) by the bi-diagonal $N \times N$ symmetric matrix $B$, where

$$B = \begin{pmatrix}
0 & x_1 & 0 & 0 & \cdots \\
x_1 & 0 & x_2 & 0 & \\
0 & x_2 & 0 & x_3 & \\
0 & 0 & x_3 & 0 & \\
   &   &   &   & \ddots
\end{pmatrix}$$

(22)

and the distribution of each $x_i$ ($i = 1, \ldots, N$) is Poissonian:

$$x_i = \begin{cases}
1 & \text{with probability } q \\
0 & \text{with probability } 1 - q
\end{cases}$$

(23)

Due to ergodicity, the spectral density, $\rho_{\text{lin}}(\lambda)$, of the ensemble of $N \times N$ matrices $B$ in the limit $N \to \infty$ coincides with the spectral density of an individual matrix $B$ and can be computed straightforwardly. Note that for any $x_k = 0$, the matrix $B$ splits into independent fragments. So, it is instructive to consider subchains consisting of sequential set of $x_k = 1$. The probability to have a subchain with $n$ consecutive 1 is $\tilde{Q}_n = q^n$, as it follows from (23). Comparing $\tilde{Q}_n$ with the distribution $Q_n = \frac{1}{2} e^{-n}$ of linear subchains in ultra-sparse matrix ensemble at the percolation point (see (14)), we conclude that $q = e^{-1}$.

In what follows we derive $\rho_{\text{lin}}(\lambda)$ for arbitrary values of $q$. This allows us to investigate the spectral density of the ensemble of bi-diagonal random operators in the limit $q \to 1$ and uncover it number-theoretic structure.

The spectrum of linear chain of length $n$ (i.e. of symmetric $n \times n$ bi-diagonal matrix $B$ with all $x_k = 1$, $k = 1, \ldots, n$) reads

$$\lambda_k = 2 \cos \frac{\pi k}{n + 1}; \quad (k = 1, \ldots, n)$$

(24)

The spectral density of the ensemble of $N \times N$ random matrices $B$ with the Poissonian distribution of matrix elements can be written as

$$\rho_{\text{lin}}(\lambda) = \lim_{N \to \infty} \frac{1}{N} \left\langle \sum_{k=1}^{N} \delta(\lambda - \lambda_k) \right\rangle$$

$$= \lim_{N \to \infty} \frac{\varepsilon}{\pi N} \text{Im} \left\langle G_N(\lambda - i\varepsilon) \right\rangle$$

$$= \lim_{N \to \infty} \frac{\varepsilon}{\pi N} \sum_{n=1}^{N} \tilde{Q}_n \text{Im} G_n(\lambda - i\varepsilon)$$

(25)
where we have used an identity
\[
\lim_{\varepsilon \to +0} \text{Im} \frac{1}{\lambda \pm i\varepsilon} = \mp i\pi \delta(\lambda)
\]
The Green function
\[
G_n(\lambda - i\varepsilon) = \sum_{k=1}^{n} \frac{1}{\lambda - \lambda_k - i\varepsilon}
\] (26)
with \(\lambda_k\) given by (24), is associated with each particular gapless matrix \(B\) of \(n\) sequential ones on the sub-diagonals, while \(\langle \ldots \rangle\) means averaging over the distribution \(\tilde{Q}_n = q^n\).

Collecting (24), (25) and (26), we find an explicit expression for the density of states, \(\rho_{\text{lin}}(\lambda)\):
\[
\rho_{\text{lin}}(\lambda) = \lim_{N \to \infty} \frac{\varepsilon}{\pi N} \sum_{n=1}^{N} q^n \sum_{k=1}^{n} \frac{1}{(\lambda - 2 \cos \frac{\pi k}{n+1})^2 + \varepsilon^2}
\] (27)

The sample plots \(\rho_{\text{lin}}(\lambda)\) for two different values of \(q\), namely for \(q = 0.7\) and \(q = e^{-1} \approx 0.37\) computed numerically via (27) with \(\varepsilon = 3 \times 10^{-3}\) are shown in the Fig. 4.

Figure 4: The spectral density \(\rho_{\text{lin}}(\lambda)\) for the ensemble of bi-diagonal matrices at \(q = 0.7\) (a) and \(q = e^{-1} \approx 0.37\) (b); the regularization parameter \(\varepsilon\) is taken \(\varepsilon = 2 \times 10^{-3}\).

**B. Analysis of enveloping curves and tails of spectral density \(\rho_{\text{lin}}(\lambda)\)**

One can compute the enveloping curves for any monotonic sequence of peaks. In Fig. 4 we show two series of sequential peaks: \(S_1 = \{1-2-3-4-5-\ldots\}\) and \(S_2 = \{2-6-7-\ldots\}\). Any monotonic sequence of peaks corresponds to the set of eigenvalues \(\lambda_k\) constructed on the basis of Farey sequence [38]. For example, as shown below, the peaks in the series \(S_1\) are located at \(\lambda_k = -2 \cos \frac{\pi k}{k+1}\), \((k = 1, 2, \ldots)\), while the peaks in the series \(S_2\) are located at \(\lambda_{k'} = \ldots\)
-2 \cos \frac{\pi^k}{2^{k-1}}, (k' = 2, 3, ...). Positions of peaks obey the following rule: let \{\lambda_{k-1}, \lambda_k, \lambda_{k+1}\} be three consecutive monotonically ordered peaks (e.g., peaks 2–3–4 in Fig. 4), and let
\[
\lambda_{k-1} = -2 \cos \frac{\pi p_{k-1}}{q_{k-1}}, \quad \lambda_{k+1} = -2 \cos \frac{\pi p_{k+1}}{q_{k+1}}
\]
where \(p_k\) and \(q_k\) \((k = 1, ..., N)\) are coprimes. The position of the intermediate peak, \(\lambda_k\), is
\[
\lambda_k = -2 \cos \frac{\pi p_k}{q_k}, \quad \frac{p_k}{q_k} = \frac{p_{k-1}}{q_{k-1}} \oplus \frac{p_{k+1}}{q_{k+1}} \equiv \frac{p_{k-1} + p_{k+1}}{q_{k-1} + q_{k+1}}
\] (28)

The sequences of coprime fractions constructed via the \(\oplus\) addition are known as Farey sequences. A simple geometric model behind the Farey sequence, known as Ford circles [39], is shown in Fig. 5a. Take the segment \([0,1]\) and draw two circles \(O_1\) and \(O_2\) both of radius \(r = \frac{1}{2}\), which touch each other, and the segment at the points 0 and 1. Now inscribe a new circle \(O_3\) touching \(O_1\), \(O_2\) and \([0,1]\). Where is the position of the new circle along the segment? The generic recursive algorithm is shown in Fig. 5b and constitutes the Farey sequence construction.

\[
\begin{align*}
\text{(a)} & \quad 0_1 \quad 1_1 \quad 1_2 \quad 2_3 \\
\text{(b)} & \quad 0_2 \quad 1_1 \quad 1_2 \quad 1_3 \quad 2_4 \\
& \quad 0_1 = \frac{1}{2} \quad 1_1 = 1 \quad 1_2 = \frac{3}{2} \quad 2_3 &= \frac{5}{3} \\
& \quad 1_2 = \frac{3}{2} \quad 2_4 = \frac{7}{4}
\end{align*}
\]

Consider two examples related to the set of eigenvalues.

- The monotonic series \(S_1\) is can be constructed recursively by taking
\[
\lambda_1 \equiv -2 \cos \frac{\pi}{2} = 0, \quad \lambda_\infty \equiv -2 \cos \frac{\pi}{1} = 2
\]
The next-to-highest peak 2 is located at
\[
\lambda_2 = -2 \cos \left[ \pi \left( \frac{1}{2} \oplus \frac{1}{1} \right) \right] = -2 \cos \frac{2\pi}{3} = 1
\]
Peak 3 is located at
\[
\lambda_3 = -2 \cos \left[ \pi \left( \frac{2}{3} \oplus \frac{1}{1} \right) \right] = -2 \cos \frac{3\pi}{4} = \sqrt{2}
\]
Peak 4 is located at
\[ \lambda_4 = -2 \cos \left[ \pi \left( \frac{3}{4} + \frac{1}{1} \right) \right] = -2 \cos \frac{4\pi}{5} = \frac{\sqrt{5} + 1}{2} \]

etc. The generic expression for \( \lambda_k \) in the series \( S_1 \) reads
\[ \lambda_k = -2 \cos \frac{k\pi}{k + 1}, \quad (k = 1, 2, \ldots) \quad (29) \]

- The monotonic series \( S_2 \) is can be constructed similarly by taking as reference points \( \lambda_1 \) and \( \lambda_2 \). For peak 6 we have
\[ \lambda_6 = -2 \cos \left[ \pi \left( \frac{1}{2} + \frac{2}{3} \right) \right] = -2 \cos \frac{3\pi}{5} = \frac{\sqrt{5} - 1}{2} \]

Peak 7 is located at
\[ \lambda_7 = -2 \cos \left[ \pi \left( \frac{1}{2} + \frac{3}{5} \right) \right] = -2 \cos \frac{4\pi}{7} \]

etc. The generic expression for \( \lambda_{k'} \) in the series \( S_2 \) reads
\[ \lambda_{k'} = -2 \cos \frac{k'\pi}{2k' - 1}, \quad (k' = 2, 3, \ldots) \quad (30) \]

As follows from \( (27) \), the function \( \rho_{\text{lin}}(\lambda) \) is nonzero only at \( \lambda_k \). In Fig. 6 we show the set of eigenvalues \( \lambda_n(k) = -2 \cos \frac{nk\pi}{n+1} \) for \( n = 1, \ldots, 50 \). Every point designates some eigenvalue, \( \lambda_k \) (see \( (24) \)); the points along each curve correspond to different values \( k \) for a specific value \( n \). The collection of eigenvalues in each horizontal line, coming from different \( n \), gives the multiplicity of corresponding eigenvalue \( \lambda_k \) and contributes to the height of a peak in the spectral density \( \rho_{\text{lin}}(\lambda) \).

The heights of peaks in any monotonic sequence (like \( S_1 \) or \( S_2 \)) can be computed directly from \( (27) \) by summing geometric series along each horizontal line in Fig. 6. For example, for the sequences \( S_1 \) and \( S_2 \) we have:

\[
\begin{align*}
S_1: \\
\lambda_1 &= -2 \cos \frac{\pi}{2} \\
\lambda_2 &= -2 \cos \frac{2\pi}{3} \\
\lambda_k &= -2 \cos \frac{\pi k}{k+1} \\
\rho^{S_1}(\lambda_k) &= \sum_{s=1}^{\infty} q^{(k+1)s-1} = \frac{q^k}{1-q^{k+1}} \quad (k = 1, 2, \ldots)
\end{align*}
\]

and

\[
\begin{align*}
S_2: \\
\lambda_2 &= -2 \cos \frac{2\pi}{3} \\
\lambda_3 &= -2 \cos \frac{3\pi}{5} \\
\lambda_{k'} &= -2 \cos \frac{\pi k'}{2k'-1} \\
\rho^{S_2}(\lambda_{k'}) &= \sum_{s=1}^{\infty} q^{(2k'-1)s-1} = \frac{q^{2k'-2}}{1-q^{2k'-1}} \quad (k = 2, 3, \ldots)
\end{align*}
\]
Figure 6: Family of 50 curves \( w_n(k) = -2 \cos \frac{n \pi k}{n+1} \). Each curve corresponds to a particular \( n = 1, \ldots, 50 \), the points mark values \( k = 1, \ldots, n \). Each horizontal line is the multiplicity of the corresponding root and contributes to the height of the peak in the spectral density.

Note that the expressions (31)–(32) are tightly linked to so-called visibility diagram (our notion is slightly modified with respect to a visibility diagram defined in [40]). Consider the square lattice of integer points \((m, n)\) and add a weight \( q_m \) to each vertical row. Emit rays at rational angles \( \alpha_{p,q} = \arctan \frac{p}{q} \) from the point \((0, 0)\) within the wedge \([\pi/4, \pi/2]\) in the positive direction as shown in Fig. 7.

Let us sum up the weights, \( q_m \), of integer points along each ray emitted at the rational slope \( \frac{p}{q} \). For example, consider the sequence of slopes \( \tan \alpha_{1,k} = \frac{1}{k} \) with \( k = 1, 2, \ldots \) and \( \tan \alpha_{2k',2k' - 1} = \frac{k'}{2k' - 1} \) shown in Fig. 7. One can verify that these geometric series exactly reproduce the sequences \( S_1 \) and \( S_2 \) in Eqs. (31)–(32) at corresponding \( k \) and \( k' \).

Two parametrically defined enveloping curves, namely \((\lambda_k, \rho^{S_1}(k))\) for the sequence \( S_1 \) \((k = 1, 2, \ldots)\) and \((\lambda_{k'}, \rho^{S_2}(k'))\) for the sequence \( S_2 \) \((k' = 2, 3, \ldots)\), are shown in Fig. 4a by dashed lines. Equations (31)–(32) allow to find the singular tails of distributions \( \rho^{S_1}(\lambda) \) as \( \lambda \to 2 \) and \( \rho^{S_2}(\lambda) \) as \( \lambda \to 0 \). Expand now the parametrically defined curves in (31)–(32) at \( k \to \infty \) and \( k' \to \infty \):

\[
S_1 : \lambda = -2 \cos \frac{\pi k}{k+1} \bigg|_{k \to \infty} \to 2 - \frac{\pi^2}{12}, \quad \rho^{S_1}(\lambda_k) \bigg|_{k \to \infty} \to q^k
\]

\[
S_2 : \lambda' = -2 \cos \frac{\pi k'}{2k' - 1} \bigg|_{k' \to \infty} \to \frac{\pi}{2k'}, \quad \rho^{S_2}(\lambda_{k'}) \bigg|_{k' \to \infty} \to q^{2k'}
\]

Expressing \( k \) and \( k' \) in (33) in terms of \( \lambda \) and \( \lambda' \), we get

\[
k \to \frac{\pi}{\sqrt{2} - \lambda}, \quad k' \to \frac{\pi}{2\lambda'}
\]

Asymptotics (34) together with (33) permit us to find the singular tails of the eigenvalue distribution \( \rho_{lin}(\lambda) \) at \( \lambda \to 2 \) and \( \lambda' \to 0 \):

\[
\rho_{lin}^{S_1}(\lambda \to 2) \to q^{\pi/\sqrt{2} - \lambda}, \quad \rho_{lin}^{S_2}(\lambda' \to 0) \to q^{\pi/\lambda'}
\]
The edge singularity $\rho_{\text{lin}}^S(\lambda \to 2)$ at $\lambda \to 2$ in Eq. (35) of the eigenvalue distribution reproduces the corresponding Lifshitz tail of the one-dimensional Anderson localization in random Schrödinger operator [41, 42]: $\rho(E) \sim e^{-E^{d/2}}$, where $E = 2 - \lambda$ and $d = 1$. The appearance of the edge singularity $\rho(E) \sim e^{-1/\sqrt{E}}$ in our situation is purely geometric, it is not relied on any entropy-energy-balance consideration like optimal fluctuation [41].

V. SPECTRAL DENSITY OF BI-DIAGONAL MATRICES IN THE LIMIT $q \to 1$ AND THE DEDEKIND $\eta$-FUNCTION

A. Limiting properties of the Dedekind $\eta$-function via duality relation

Recall the definition of the Dedekind $\eta$-function [43]:

$$\eta(z) = e^{\pi iz/12} \prod_{n=0}^{\infty} (1 - e^{2\pi inz}); \quad z = x + iy \quad (y > 0)$$  \hspace{1cm} (36)

Consider now the following function

$$f(z) = A^{-1} |\eta(z)|(|\text{Im } z|)^{1/4}; \quad A = \left| \eta \left( \frac{1}{2} + i \frac{\sqrt{3}}{2} \right) \left( \frac{\sqrt{3}}{2} \right)^{1/4} \right| \approx 0.77230184...$$  \hspace{1cm} (37)

This function satisfies the following duality relation

$$f \left( \left\{ \frac{m}{k} \right\} + iy \right) = f \left( \left\{ \frac{s}{k} \right\} + \frac{i}{k^2 y} \right); \quad ms - kr = 1; \quad \{m, s, k, r\} \in \mathbb{Z}; \quad y > 0$$  \hspace{1cm} (38)
where \( \{ \frac{m}{k} \}, \{ \frac{s}{k} \} \) denote fractional parts of corresponding quotients. Two particular identities follow from (38):

\[
f \left( \frac{k}{k + 1} + iy \right) = f \left( \frac{1}{k + 1} + \frac{i}{(k + 1)^2 y} \right)
\]

\[
f \left( \frac{k'}{2k' - 1} + iy \right) = f \left( \frac{2}{2k' - 1} + \frac{i}{(2k' - 1)^2 y} \right)
\]

(39)

Using (39) we can obtain the asymptotics of \( \eta(z) \) when \( y \to 0^+ \). Take into account the relation of Dedekind \( \eta \)–function with Jacobi elliptic functions:

\[
\vartheta'_1(0, e^{\pi iz}) = \eta^3(z)
\]

(40)

where

\[
\vartheta'_1(0, e^{\pi iz}) = \left. \frac{d\vartheta_1(u, e^{\pi iz})}{du} \right|_{u=0} = e^{\pi iz/4} \sum_{n=0}^{\infty} (-1)^n (2n + 1) e^{\pi i n(n+1)z}
\]

(41)

Rewrite (39) as follows:

\[
\left| \eta \left( \frac{k}{k + 1} + iy \right) \right| = \frac{1}{\sqrt{(k + 1)y}} \left| \eta \left( \frac{1}{k + 1} + \frac{i}{(k + 1)^2 y} \right) \right|
\]

(42)

(Since the computations are identical for both equations in (39), we do not reproduce the derivation for the second one, and will write for it the final answer only). Applying (40)–(41) to (42), we get:

\[
\left| \eta \left( \frac{k}{k + 1} + iy \right) \right| = \frac{1}{\sqrt{(k + 1)y}} e^{\frac{\pi i}{12(1 + k)^2}} \sum_{n=0}^{\infty} (-1)^n (2n + 1) e^{\pi i n(n+1)z} (1/3)
\]

(43)

Equation (43) enables us to extract the leading asymptotics of \( \left| \eta \left( \frac{k}{k + 1} + iy \right) \right| \) in the \( y \to 0^+ \) limit. Noting that every term of the series in (43) for any \( n \geq 1 \) and for small \( y \) (i.e. for \( y \ll (k + 1)^{-2} \)) converges exponentially fast, we have

\[
\left| \eta \left( \frac{k}{k + 1} + iy \right) \right| \quad \text{as} \quad y \to 0^+ \quad \Rightarrow \quad \frac{1}{\sqrt{(k + 1)y}} e^{-\frac{\pi}{12(k + 1)^2}};
\]

(44)

Similarly, we get the asymptotics for \( \left| \eta \left( \frac{k'}{2k' - 1} + iy \right) \right| \) at \( y \to 0^+ \) (valid for \( y \ll (2k' - 1)^{-2} \)). Thus, we have finally:

\[
\sqrt{-\ln \left| \eta \left( \frac{k}{k + 1} + iy \right) \right|} \quad \text{as} \quad y \to 0^+ \quad \Rightarrow \quad \frac{1}{\sqrt{k + 1}} \sqrt{\frac{\pi}{12y}}
\]

(45)

In (45) we have dropped out the subleading terms at \( y \to 0^+ \), which are of order of \( O(\ln y) \).
B. Tails of bi-diagonal matrices at $q \to 1$ and relation with Dedekind $\eta$-function

Expressions (31)–(32) allow to analyze the tails of the distribution $\rho_{\text{lin}}(\lambda)$ when $q \to 1$ and $k$ is fixed. Namely, taking the limit $q \to 1$ in (31)–(32), we arrive at:

$$
\rho_{1}^{S_{1}}(\lambda k) \bigg|_{q \to 1} = \frac{q^{k}}{1 - q^{k+1}} \to \frac{1}{(k + 1)(1 - q)}
$$

$$
\rho_{1}^{S_{2}}(\lambda k') \bigg|_{q \to 1} = \frac{q^{2k'-2}}{1 - q^{2k'-1}} \to \frac{1}{(2k' - 1)(1 - q)}
$$

(46)

These expressions lead us to the following asymptotics of tails of the distribution $\rho(\lambda)$ at $\lambda \approx 2 - \frac{\pi^{2}}{k^{2}} |_{k \gg 1} \to 2$ and $\lambda' \approx \frac{\pi}{2k'} |_{k \gg 1} \to 0$:

$$
\rho_{1}^{S_{1}} \bigg|_{q \to 1} (\lambda \to 2) \to \frac{\sqrt{2 - \lambda}}{\pi(1 - q)}
$$

$$
\rho_{1}^{S_{2}} \bigg|_{q \to 1} (\lambda' \to 0) \to \frac{\lambda'}{\pi(1 - q)}
$$

(47)

Note the essential difference with (35). It is easy to see that all tails of intermediate monotonic sequences have scaling behavior as in the sequence $\rho_{1}^{S_{2}}(\lambda)$.

Return now to (45) and substitute for $k$ and $k'$ the asymptotic expansions $k(\lambda) \approx \frac{\pi}{\sqrt{2 - \lambda}}$ and $k'(\lambda') \approx \frac{\pi}{\lambda}$ (see (34)). Since near the spectrum edges, $k \gg 1$ and $k' \gg 1$, we get

$$
\sqrt{-\ln |\eta(\lambda + iy)|} \bigg|_{\lambda \to 2, y \to 0} \to \frac{\sqrt{2 - \lambda}}{\sqrt{12\pi y}} \quad \text{and} \quad \sqrt{-\ln |\eta(\lambda + iy)|} \bigg|_{\lambda' \to 0, y \to 0} \to \frac{\lambda}{\sqrt{12\pi y}}
$$

(48)

The tails in (48) exactly match the tails of the spectral density $\rho^{A}(\lambda)$ and $\rho^{B}(\lambda)$ in (47) if we make an identification $1 - q = \sqrt{12\pi y}$.

The same analysis can be carried out for any monotonic sequence of peaks of the spectral density depicted in Fig. 4. This brings us to the central conjecture of this work, which uncovers the number theoretic nature of the spectral density of Poissonian ensemble of large random bi-diagonal symmetric matrices.

**Conjecture.** Let $\rho_{\text{lin}}(\lambda, q)$ be the spectral density of ensemble of $N \times N$ bi-diagonal symmetric random Bernoulli matrices with the Poissonian distribution of nonzero matrix elements defined in Eqs. (22)–(23). For $N \to \infty$ we have the relation:

$$
\lim_{q \to 1^{-}} \frac{\rho_{\text{lin}}(\lambda, q)}{\sqrt{-\ln |\eta(\frac{1}{\pi} \arccos(-\lambda/2 + i(1-q)^{2}/12\pi)|}} = 1
$$

(49)

where $\eta$ is the Dedekind $\eta$-function defined in (36). One can say, that Eq. (49) establishes the link between spectral density of random one-dimensional Schrödinger-type operators with the limiting values of modular function at rational points on the real axis.

Empirical evidence in favor of the above conjecture is presented in Fig. 8 where we plot three functions:
Fig. 8a: Numerically generated spectral density of ensemble of $N \times N$ ($N = 3000$) bi-diagonal Bernoulli random matrices $B$ (see (22)) with the Poissonian distribution of “1” at $q = 0.98$;

Fig. 8b: Numerically computed eigenvalue density $\rho(\lambda, q)$ given by Eq. (27) for matrices of size $N = 3000$ and $q = 0.98$;

Fig. 8c: Analytic expression given by the right-hand side of (49) at $y = \frac{(1-q)^2}{12\pi} \approx 10^{-5}$.

Figure 8: Spectral density $\rho_{\text{lin}}(\lambda)$ of ensemble of bi-diagonal Poissonian random matrices (shown only for $\lambda \geq 0$): a) direct numeric simulation; b) numeric visualization of Eq. (27); c) analytic construction provided by Eq. (49).

VI. DISCUSSION

A. Ultrametricity and rare-event statistics

We investigated the spectral density of a sub-ensemble of sparse symmetric Bernoulli matrices near the percolation point. We showed that in the vicinity of the percolation point linear chains constitute the overwhelming majority of clusters—about 95%. This led us to study the spectral density of ensemble of linear chains only. The advantage of this consideration is the possibility of exact treatment which allowed us to uncover the number-theoretic structure of the corresponding density of states.

In a more practical setting, our analysis is applicable to investigation of polydisperse solutions of linear polymers at any concentrations with Poissonian distribution in chain lengths, where the ultrametricity could be directly seen in the absorption spectra. One tacit message coming from our work is that experimenting with biological activity of highly diluted solutions of biologically active substances, one should pay attention to a very peculiar structure of background noise originated from the rare-event statistics of dissolved clusters. In order to make conclusion about any biological activity of regarded chemical substance, the signal from background noise should be clearly identified.
As concerns exact results, we showed that the spectral density, $\rho_{\text{lin}}(\lambda)$, in the ensemble of bi-diagonal random symmetric matrices with the Poissonian distribution of matrix elements demonstrates a very peculiar ultrametric structure.

The derivation of the shape of the enveloping curve for the spectral density, $\rho_{\text{lin}}(\lambda)$, of random Shrödinger-type operators presented by random symmetric bi-diagonal matrices, is based on simultaneous localization of eigenvalues, $\lambda_k = 2 \cos \frac{\pi}{k} (k = 1, 2, \ldots)$, constituting the principal sequence, and of their multiplicativity (degeneracy), $\rho(\lambda_k) = q_k^{1/2}$ ($0 < q < 1$). Eliminating $k$ from the parametric dependence ($\lambda_k, \rho(\lambda_k)$), one gets the signature of a Lifshitz tail for Anderson localization at the edge of the spectrum, $\rho(\lambda \rightarrow 2) \sim q^{-\pi/\sqrt{2-\lambda}}$, typical for the one-dimensional Schrödinger random operators. We emphasize that in our case the derivation is purely geometric.

Analyzing the limit $q \rightarrow 1$, we have demonstrated that the function $\rho_{\text{lin}}(\lambda)$ shares number-theoretic properties connected with the structure of the Dedekind $\eta$–function near the real axis at rational points. Intriguingly, the limiting behavior of the Dedekind function near real axis at rational points has tantalizing connections with Vassiliev knot invariants [44] and with quantum invariants of 3-manifolds [45]; a new concept of quantum modular forms was recently suggested [46] as a potential explanation.

B. Manifestation of the Dedekind $\eta$-function in natural science

Finally, we mention two other physical systems exhibiting a connection with the $\eta$ function. In Ref. [47] it has been argued that buckling of a leaf of some plants (like lettuce or spinach) is related with the isometric embedding of hyperbolic graphs (Cayley trees) into the 3D Euclidean space. This buckling is described by a function, $g(z)$, (Jacobian of transformation), which defines the deformation of the hyperbolic graph in its projection from the 3D Euclidean space into the complex plane (see also the Section 5.1 of [48]).

For a regular 3-branching Cayley tree, the function $g(z)$ can be expressed via the Dedekind $\eta$-function [47, 49]. It has been noted in [49] that properly normalized function $g(z)$, namely, $f(z) = A^{-1}|\eta(z)|/(\text{Im } z)^{1/4}$, defined in Eq. (37), can be regarded as a continuous tree isometrically embedded into the half-plane $\text{Im } z > 0$. The constant $A$ in (37) is chosen to set the maximal value of the function $f(z)$ to 1: $0 < f(z) \leq 1$ for any $z$ in the upper half-plane $\text{Im } z > 0$. The function $f(z)$, being the modular function, has the following property [49]: $f(z)$ has local maxima equal to 1 at the vertices of the 3-branching Cayley tree (and only of them) isometrically embedded into the upper half-plane $\text{Im } z > 0$. The cut of the function $f(z)$, shown in Fig. 9b, is equivalent to the Farey construction.

Another well-known manifestation of number theory in natural science arises in the context of phyllotaxis [50]. The connection of the cell packing with the Farey sequences has been observed long time ago. It was not clear, however, why nature selects the Fibonacci sequence among other possible Farey sequences. A tantalizing answer to this question has been given by L. Levitov, see [51]. He proposed an “energetic” approach to the phyllotaxis,
Figure 9: Cut \((f(z) \in [0.985, 1.0])\) of the contour plot of the function \(f(z)\) (see (37)): a) the representation in the upper half-plane, \(z = x + iy\); b) the representation in the unit disc obtained by the conformal mapping of the upper half-plane to the unit disc.

suggesting that the development of a plant is connected with an effective motion along the geodesics on the surface associated with the energetic relief of growing plant.

Intriguingly, the energy relief discussed in [51] coincides with the relief of the function \(f(z)\) depicted in Fig. 9. The black arcs drawn in Fig. 9a,b are parts of semicircles orthogonal to the boundaries and represent the geodesics of the surface \(f(z)\), which are open level lines (by the symmetry, there are only two such level lines in Fig. 9a and three in Fig. 9b). The best rational approximation of the boundary point, \(\frac{\sqrt{5} - 1}{2}\), at which the solid geodesics in Fig. 9a terminates, is given by the continued fraction expansion

\[
\frac{\sqrt{5} - 1}{2} = \frac{1}{1 + \frac{1}{1 + \frac{1}{1 + \ldots}}},
\]

(50)
cut at some level \(k\). Cutting (50) at sequential \(k\), we get the set of fractions constituting the interlacing Fibonacci sequences, \(\{F_k\}\), in nominators and denominators:

\[
\frac{1}{1}, \frac{1}{1'}, \frac{2}{2'}, \frac{5}{3'}, \frac{8}{5'}, \frac{F_{k-1}}{F_k'}, \frac{F_k}{F_{k+1}}.
\]

Thus, the normalized \(\eta\)-function, \(f(z)\), plays a role of the energy relief of a growing plant.

One can say that the Fibonacci numbers are the benchmarks, which fix the connection between ultrametric and hyperbolic geometry, since, on one hand they appear as discrete symmetries of the hyperbolic space, and on the other hand, being a subset of \(p\)-adic numbers, they uniquely parameterize the distances in the ultrametric space [52].
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