Spectrum of the Laplace-Beltrami Operator and the Phase Structure of Causal Dynamical Triangulation

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We propose a new method to characterize the different phases observed in the non-perturbative numerical approach to quantum gravity known as Causal Dynamical Triangulation. The method is based on the analysis of the eigenvalues and the eigenvectors of the Laplace-Beltrami operator computed on the triangulations: it generalizes previous works based on the analysis of diffusive processes and proves capable of providing more detailed information on the geometric properties of the triangulations. In particular, we apply the method to the analysis of spatial slices, showing that the different phases can be characterized by a new order parameter related to the presence or absence of a gap in the spectrum of the Laplace-Beltrami operator, and deriving an effective dimensionality of the slices at the different scales. We also propose quantities derived from the spectrum that could be used to monitor the running to the continuum limit around a suitable critical point in the phase diagram, if any is found.

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I. INTRODUCTION

Causal Dynamical Triangulations (CDT) [1] is a numerical Monte-Carlo approach to Quantum Gravity based on the Regge formalism, where the path-integral is performed over geometries represented by simplicial manifolds called “triangulations”. The action employed is a discretized version of the Einstein-Hilbert one, and the causal condition of global hyperbolicity is enforced on triangulations by means of a space-time foliation.

One of the main goals of CDT is to find a critical point in the phase diagram where the continuum limit can be performed in the form of a second-order phase transition. The phase diagram shows the presence of four different phases [2–7], and the hope is that the transition lines separating some of these phases could contain such a second order critical point. Presently, such phases are identified by order parameters which are typically based on the counting of the total number of simplexes of given types or on other similar quantities (e.g., the coordination number of the vertices of the triangulation). The main motivation of the present study is to enlarge the set of observables available for CDT, trying in particular to find new order parameters and to better characterize the geometrical properties of the various phases at different scales.

One successful attempt to characterize the geometries of CDT has been obtained by implementing diffusion processes on the triangulations [9, 10]. In practice, one analyzes the behavior of random walkers moving around the triangulations: from their properties (e.g., the return probability) one can derive relevant information, such as the effective dimension felt at different stages of the diffusion (hence at different length scales). In this way, estimates of the spectral dimension of the triangulations have been obtained.

In this paper we propose and investigate a novel set of observables for CDT configurations, based on spectral methods, namely, the analysis of the eigenvalues and the eigenvectors of the Laplace–Beltrami (LB) operator. This can be viewed as a generalization of the analysis of the spectral dimension, since the Laplace–Beltrami operator completely specifies the behavior of diffusion processes (see Appendix A for a closer comparison). Still, as we will show in the following, the Laplace–Beltrami operator contains more geometric information than just the spectral dimension.

Nowadays, spectral methods find application in a huge variety of different fields. To remember just a few of them, we mention shape analysis in computer aided design and medical physics [11, 12], dimensionality reduction and spectral clustering for feature selection/extraction in machine learning [13], optimal ordering in the PageRank algorithm of the Google Search engine [14], connectivity and robustness analysis of random networks [15]. Therefore, the application to CDT is just one more application of a well known analysis tool. On the other hand, some well known results which have been established in other fields will turn out to be useful in our investigation of CDT.

In the present paper, we limit our study to the LB spectrum of spatial slices. Among the various results, we will show that the different phases can be characterized by the presence or absence of a gap in the spectrum of the LB operator, as it happens for the spectrum of the Dirac operator in strong interactions, and we will give an interpretation of this fact in terms of the geometrical properties of the slices. The presence/absence of a gap will also serve to better characterize the two
different classes of spatial slices which are found in the recently discovered bifurcation phase \[4, 8\]. Moreover, we will show how the spectrum can be used to derive an effective dimensionality of the triangulations at different length scales, and to investigate quantities useful to characterize the critical behavior expected around a possible second order transition point.

The paper is organized as follows. In Section II, we discuss our numerical setup together with a short review of the CDT approach, summarizing in particular the major features of the phase diagram that will be useful for the discussion of our results. In Section III, we describe some of the most relevant properties of the Laplace-Beltrami operator in general, then focusing on its implementation for the spatial slices of CDT configurations and discussing a toy model where the relation between the LB spectrum and the effective dimensionality of the system emerges more clearly. Numerical results are discussed in Section IV. Finally, in Section V, we draw our conclusions and discuss future perspectives. Appendix A is devoted to a discussion of the relation existing between the spectrum of the LB operator and the spectral dimension, defined by diffusion processes as in Ref. \[10\].

II. A BRIEF REVIEW ON CDT AND NUMERICAL SETUP

It is well known that, perturbatively, General Relativity without matter is non-renormalizable already at the two-loop level \[16\]. Nevertheless, interpreted in the framework of the Wilsonian renormalization group approach \[17\], this really means that the gaussian point in the space of parameters of the theory is not an UV fixed point, as for example it happens for asymptotically free theories. Indeed, Weinberg conjecture of asymptotic safety of the gravitational interaction \[18\] states the existence of an UV non-gaussian fixed point, which makes the theory well defined in the UV (i.e. renormalizable), but in a region of the phase diagram not accessible by perturbation theory. Various non-perturbative methods have been developed in the last decades to investigate this possibility, like Functional Renormalization Group techniques \[19\] or the Monte-Carlo simulations of standard Euclidean Dynamical Triangulations (DT) \[20, 23\] or Causal Dynamical Triangulations, the latter being the subject of this study.

Monte-Carlo simulations of quantum field theories are based on the path-integral formulation in Euclidean space, where expectation values of any observable \(O\) are estimated as averages over field configurations sampled with probability proportional to \(e^{-\frac{S}{\hbar}}\), \(S\) being the action functional of the theory. Regarding the Einstein–Hilbert theory of gravity, the action is a functional of the metric field \(g_{\mu\nu}\), given by

\[
S[g_{\mu\nu}] = \frac{1}{16\pi G} \int d^d x \sqrt{-g} \left( R - 2\Lambda \right)
\]

where \(G\) and \(\Lambda\) are respectively the Newton and Cosmological constants, while the path–integral expectation values are formally written as averages over geometries (classes of diffeomorphically equivalent metrics)

\[
\langle O \rangle = \frac{1}{Z} \int \mathcal{D}[g_{\mu\nu}] \mathcal{O}[g_{\mu\nu}] e^{-\frac{S[g_{\mu\nu}]}{\hbar}},
\]

where \(Z\) is the partition function.

The first step in setting up Monte-Carlo simulations is the choice of a specific regularization of the dynamical variables into play. In the case of gravity without matter fields the only variable is the geometry itself, which can be conveniently regularized in terms of triangulations, namely a collection of simplexes, elementary building blocks of flat spacetime, glued together to form a space homeomorphic to a topological manifold. The simplexes representing (spacetime) volumes in 4–dimensional spaces are called pentachorons, analogous to tetrahedra in 3–dimensional spaces and triangles in 2–dimensional spaces (i.e. surfaces).

Besides the general definition, and at variance with standard DT, triangulations employed in CDT simulations are required to satisfy also a causality condition of global hyperbolicity\(^2\). This is realized by assigning an integer time label to each vertex of the triangulation in order to partition them into distinct sets of constant time called spatial slices, and constraining simplexes to fill the spacetime between adjacent slices (i.e. slices with neighbouring integer labels). The resulting triangulation has therefore a foliated structure\(^3\), and the simplexes can be classified by a (time–ordered) pair specifying the number of vertices on the slices involved (e.g., the pairs \(4, 1\), \((3, 2)\), \((2, 3)\) and \((1, 4)\) classify all spacetime pentachorons). In order to ensure both the simplicial manifold property and the foliated structure at the same time, spatial slices, considered as simplicial submanifolds composed of glued spatial tetrahedra, need to be topologically equivalent. This basically means that triangulations are always geodetically complete manifolds, and topological obstructions (e.g., singularities) can only be realized

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1. For simplicity, we are not including manifolds with boundaries, so there is no Gibbons–Hawking–York term in the action.
2. The global hyperbolicity condition is equivalent to the existence of a Cauchy surface, the strongest causality condition which can be imposed on a manifold \[24\].
3. The main reason for restricting to foliated triangulations is that it allows to define conveniently the analytical continuation from Lorentzian to Euclidean space (see Ref. \[11\] for details). However, simulations without preferred foliation in \(2+n\) dimensions have been build in Ref. \[25\], showing results similar to the foliated case.
in an approximate fashion, with increasing accuracy in the thermodynamic limit (infinite number of simplexes). The numerical results shown in Section IV refer to slices with \(S^3\) topology, but other topologies could be investigated as well (e.g., the toroidal one \([1,8]\)).

In practice, it is convenient, without loss of generality, to impose a further condition, that is fixing the length-squared of every spacelike link (i.e. connecting vertices on the same slice) to a constant value \(a^2\), and the square-length of every timelike link (i.e. connecting vertices on adjacent slices) to a constant value \(-\alpha a^2\). The constant \(\alpha\) takes the role of lattice spacing, while \(\alpha\) represents a genuinely regularization–dependent asymmetry in the choice of time and space discretizations. With this prescription, simplexes in the same class (according to the above definition) not only are equivalent topologically, but also geometrically, so that the expression of the discretized action greatly simplifies. Indeed, at the end of the day, the standard 4–dimensional action employed in CDT simulations with \(S^3\) topology of the slices and periodic time conditions becomes a functional of the triangulation \(\mathcal{T}\), and takes the relatively simple form

\[
S_E = -k_0 N_0 + k_4 N_4 + \Delta (N_4 + N_{41} - 6 N_0),
\]

where \(N_0\) counts the total number of vertices, \(N_4\) counts the total number of pentachorons, and \(N_{41}\) is the sum of the total numbers of type (4,1) and type (1,4) pentachorons, while \(k_4, k_0\) and \(\Delta\) are free dimensionless parameters, related to the Cosmological constant, the Newton constant, and the freedom in the choice of the time/space asymmetry parameter \(\alpha\) (see Ref. [1] for more details).

We want to stress that, even if CDT configurations are defined by means of triangulations, the ultimate goal of the approach is to perform a continuum limit in order to obtain results describing continuum physics of quantum gravity. Therefore, the specific discretization used in CDT must be meant as artificial, becoming irrelevant in the continuum limit. For this reason, simplexes should not be considered as forming the physical fabric of spacetime: eventually, one would like to find a critical point in the parameter space where the correlation length diverges and the memory about the details of the fine structure is completely lost.

In standard CDT simulations, configurations are sampled using a Metropolis–Hastings algorithm [26], where local modifications of the triangulation at a given simulation time (i.e. insertions or removals of simplexes) are accepted or rejected according to the probability induced by the action in Eq. (3) and complying with the constraints discussed above.

Unlike usual lattice simulations of quantum field theories, the total spacetime volume of CDT triangulations changes after a Monte Carlo update. In order to take advantage of finite size scaling methods (i.e. extrapolation of results to the infinite volume limit), it is convenient to control the volume by performing a Legendre transformation from the parameter triple \((k_4, k_0, \Delta)\) to the triple \((V, k_0, \Delta)\), where the parameter \(k_4\) is traded for a target volume \(V\). In practice, this is implemented by a fine tuning of the parameter \(k_4\) to a value that makes the total spacetime or spatial volumes\(^5\) fluctuate with mean around a chosen target volume (respectively \(\mathcal{N}_4\) or \(\mathcal{N}_{41}\)), and adding to the sample only configurations whose total volume lies in a narrow range around the target one. Moreover, a (weak) spacetime volume fixing to a target value \(\mathcal{N}_4\) can be enforced, for example, by adding a term to the action of the form \(\Delta S = \epsilon (N_4 - \mathcal{N}_4)^2\), where \(\epsilon\) quantifies how much large volume fluctuations are suppressed. A relation similar to the latter holds for fixing the total spatial volume (substituting \(N_{41}\) with \(N_4\)). Fixing a target total spatial volume \(V_{S,tot} = \mathcal{N}_4 \mu\), one can investigate the properties of configurations sampled at different values of the remaining free parameters \(k_0\) and \(\Delta\).

The general phase structure of CDT which is found in the \(k_0 - \Delta\) plane is thoroughly discussed in the literature [1, 5, 6, 27]. Here we will only recall some useful facts. Four different phases have been identified, called \(A, B, C_{ds}\) and \(C_b\), as sketched in Fig. 1, where for the two \(C\) phase the labels \(dS\) and \(b\) stand respectively for de Sitter and bifurcation. At a qualitative level, configurations in the different phases can be characterized by the distribution of their spatial volume \(V_S(t)\), which counts the number of spatial tetrahedra (spatial volume \(V_S\)) in each slice as a function of the slice time \(t\). For configurations in the \(B\) phase, the spatial volume is concentrated almost in a single slice, leaving the other slices with a minimal volume\(^6\). For both the \(C_{ds}\) and \(C_b\) phase, the spatial volume is peaked at some slice–time but then, unlike the case of the \(B\) phase, falls off more gently with \(t\), so that the majority of the total spatial volume is localized in a so-called “blob” with a finite time extension; also in this case, slices out of this blob have a minimal volume. Finally, configurations in phase \(A\) are characterized by multiple and uncorrelated peaks in the spatial volume distribution.

From these observations, it is apparent that \(C_{ds}\) and \(C_b\) are the only physically relevant phases. Indeed, the average spatial volume distribution in the \(C_{ds}\) phase is in good agreement with the prediction for a de Sitter Uni-

\(^4\) A series of steps is needed in order to obtain the CDT action in Eq. (3): Regge discretization of the continuous action, computation of volumes and dihedral angles, Wick rotation and the use of topological relations between simplex types.

\(^5\) The total spatial volume of a configuration is the number of spatial tetrahedra, which equals \(\frac{24}{5}\) by elementary geometrical arguments.

\(^6\) Triangulations with \(S^3\) topology, like spatial slices in \(3 + 1\) standard CDT simulations, must have at least 5 tetrahedra.
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versus, having a $S^4$ geometry after analytical continuation
to the Euclidean space [28]. The bifurcation phase, in
stead, is characterized by the presence of two different
classes of slices which alternate each other in the slice
time $t$ [5, 6].

The transition lines between the different phases
(dashed lines in Fig. 1) have been investigated by means
of convenient observables. Regarding the $B$-$C_b$ and
$A$-$C_{dS}$ transition lines, the definitions employed are
based on the observation that changes in the qualitative
behavior of the spatial volume distribution function
$V_S(t)$ occur for almost constant values of $\Delta$ or $k_0$
respectively, suggesting quantities conjugated to them in
the action [3] as candidate order parameters: namely
$\text{conj}(\Delta) \equiv (N_A + N_{dS} - 6N_0)/N_4$ for the $B$-$C_b$
transition and $\text{conj}(k_0) \equiv N_0/N_{dS}$ for the $A$-$C_{dS}$ transition.
Finite size scaling computations using these observables
as order parameters suggest a first-order nature for the
$A$-$C_{dS}$ transition, while the $B$-$C_b$ transition appears to be of second-order [27].

The definition of observables employed as order parameters for the $C_b$-$C_{dS}$ transition is more involved [5, 6]: in the $C_b$ phase, one of the two classes of spatial slices is characterized by the presence of vertices with very high coordination number; also in this case there are hints for a second-order transition, even if results might depend on the topology chosen for the spatial slices [7].

Global counts of simplexes, like those entering the
definitions of $\text{conj}(\Delta)$ and $\text{conj}(k_0)$, are not sufficient to clearly distinguish the different geometrical properties of the various phases. From this point of view, the spectral dimension $D_S(\tau)$ (see Appendix A for more details) is probably one of the few useful probes available
up to now to probe the geometrical structure of CDT configurations. It is basically a measure of the effective dimension of the geometry at different stages of the diffusion process, it has permitted to demonstrate that, in the bulk of configurations in the de Sitter phase, the spectral dimension tends to a value $D_s \simeq 4$ for large diffusion times [10]. In the following, we will show how the analysis of the spectrum of the LB operator, which is discussed in the following section, permits to access new classes of observables, and how some clear characteristic differences among the various phases emerge in this way.

The code employed for this study is an home–made
implementation in C++ of the standard CDT algorithm
discussed in Ref. [1], which was checked against many
of the standard results which can be found in the CDT
literature. We performed simulations with parameters
chosen as shown in Fig. 1 by points marked with a star
symbol and reported also in Table I for later convenience. The assignment of simulation points to the
different phases refers to the total volumes fixed in our
runs ($N_{dS} = 40k$ and $80k$).

![FIG. 1: Sketch of the phase diagram CDT in 4d and with spherical topology of spatial slices. The results shown in the present paper have been obtained from simulations running at the points marked by a star symbol *.
](image)

TABLE I: Coordinates ($k_0$ and $\Delta$) for the simulation points chosen, as shown in Figure 1 and the phases in which they are contained. Some of the points are labeled also by a letter for later convenience. The assignment of simulation points to the different phases refers to the total volumes fixed in our runs ($N_{dS} = 40k$ and $80k$).

| $k_0$ | $\Delta$ | Phase |
|-------|---------|-------|
| 2.2   | -0.2    | $B$   |
| 2.2   | -0.05   | $B$   |
| 2.2   | 0.022   | $B$   |
| 2.2   | 0.05    | $C_b$ |
| 2.2   | 0.1     | $C_b$ |
| 2.2   | 0.15    | $C_b$ |
| 2.2   | 0.3     | $C_{dS}$ |
| 2.2   | 0.45    | $C_{dS}$ |
| 2.2   | 0.6     | $C_{dS}$ |
| 5     | 0.6     | $A$   |

III. THE LAPLACE-BELTRAMI OPERATOR

The LB operator, usually denoted by the symbol $-\Delta$, is the generalization of the standard Laplace operator. Its specific definition depends on the underlying space and on the algebra of functions on which it acts. For a generic smooth Riemannian manifold $(\mathcal{M}, g_{\mu\nu})$ the Laplace-Beltrami operator acts on the algebra of smooth
functions $f \in C^\infty(M)$ in the form \[29]:
\[
-\Delta f = -\frac{1}{\sqrt{|g|}} \partial_{\mu}(\sqrt{|g|} g^{\mu\nu} \partial_{\nu} f) = -g^{\mu\nu} (\partial_{\mu} \partial_{\nu} - \Gamma_{\mu\nu}^{\alpha} \partial_{\alpha}) f,
\]
where $g$ is the metric determinant, $g^{\mu\nu}$ is the inverse metric and $\Gamma_{\mu\nu}^{\alpha}$ are the Christoffel symbols.

It is easily shown that $-\Delta$ is invariant with respect to isometries. Furthermore, since it is positive semi-definite, a set of eigenvectors $B_M$ solving the eigenvalue problem $-\Delta f = \lambda f$ is an orthogonal basis for the algebra $C^\infty(M, \mathbb{R})$; in the following we will refer to such sets as spectral bases, which, for convenience and without loss of generality, we will always consider orthonormal. A spectral basis can then be used to define the Fourier transform of spatial slices, by means of their associated dual graphs.

In the form of Eq. (8) the geometric role of eigenvectors described by the heat equation
\[
\partial_t u(x, x_0; t) - \Delta u(x, x_0; t) = 0.
\]
We can expand the solution in a spectral basis $B_M = \{e_n | \lambda_n \in \sigma_M, \lambda_{n+1} \geq \lambda_n \}$ associated to the spectrum of (increasingly ordered) eigenvalues $\sigma_M = \{\lambda_n\}$
\[
u(x; t) = \sum_{n=0}^{[\sigma_M|-1]} u_n(t) e_n(x),
\]
so that Eq. (5) is transformed (by orthogonality) in a set of decoupled equations
\[
\partial_t u_n(t) = -\lambda_n u_n(t) \ \forall n,
\]
\[
\Rightarrow u(x; t) = \sum_{n=0}^{[\sigma_M|-1]} e^{-\lambda_n t} u_n(0) e_n(x).
\]
In the form of Eq. (8) the geometric role of eigenvectors in the diffusion process is evident: $\lambda_n$ represents the diffusion rate of the mode $e_n(x)$, so that the smallest eigenvalues are associated to eigenvectors along the slowest diffusion directions and vice versa. In this specific sense, the spectrum $\sigma_M$ encodes information about the characteristic scales of the manifold, while the set of eigenstates $B_M$ identifies all the possible diffusion modes, and forms a basis for the algebra of functions on the manifold. Similar considerations can be applied to the problem of wave propagation on the manifold, where the heat equation is replaced by the wave equation; this is the reason behind the famous idea of “hearing the shape of a drum” \[30].

The definition of the Laplace–Beltrami operator can be extended easily to more general algebras, like the graded algebra of differential forms or the algebra of functions on a graph \[31, 32\], the latter being of particular importance in our discussion, since, as discussed below, it allows us to implement straightforwardly the spectral analysis on CDT spatial slices, by means of their associated dual graphs. A undirected graph $G$ \[33\] is formally a pair of sets $(V, E)$, where $V$ contains vertices, which assume the role of lattice sites, whereas the set of edges, $E \subset V \times V$, is a symmetric binary relation on $V$ encoding the connectivity between vertices in the form of ordered pairs of vertices $\{(v_i, v_j)\}$. The reason why, in this first study, we choose to apply spectral methods to analyze the geometry of spatial slices only is that spatial tetrahedra have all link lengths equal to the spatial lattice size $a$, so that the distance between their centers is equal for any adjacent tetrahedra; therefore, it is possible to represent faithfully spatial slices by dual undirected and unweighted graphs, where the vertex set is the set of tetrahedra, and the edge set is the adjacency relation between tetrahedra. The algebra on which the Laplace–Beltrami operator acts can be taken as that of the real-valued functions $f : V \to \mathbb{R}$, which can be represented as the vector space $\mathbb{R}^N$ (where $N = |V|$), once an ordering of the vertices $i \to v_i \in V \ \forall i \in \{0, 1, \ldots, N - 1\}$ has been arbitrarily chosen, without loss of generality\[7\]. In this representation the Laplace–Beltrami operator becomes formally a matrix, named Laplace matrix, and defined as:
\[
L = D - A,
\]
where $D$ is the (diagonal) degree matrix such that the element $D_{ii} \equiv |\{e \in E | v_i \in e\}|$ counts the number of vertices connected to the vertex $v_i$, while $A$ is the symmetric adjacency matrix such that the element $A_{ij} = \chi_E(\{v_i, v_j\})$ is 1 only if the vertices $v_i$ and $v_j$ are connected (i.e. $\{v_i, v_j\} \in E$) and zero otherwise.

For instance, the graph associated with a one-dimensional hypercubic lattice with $N$ sites and periodic boundary conditions corresponds to $D = 2 \cdot 1_{N \times N}$, and $A_{ij} = \delta_{i, (j+1) \mod N} + \delta_{i, (j-1) \mod N}$, while the Laplace matrix can be read off as the lowest order approximation to the Laplace–Beltrami operator estimated by evaluating functions on lattices sites:
\[
-\Delta f(x_i) = -\frac{d^2 f}{dx_i^2}(x_i) = \frac{2f_i - f_{i+1} - f_{i-1}}{a^2} + O(a),
\]
where $a$ is the lattice spacing and $f_n \equiv f(x_{(n \mod N)})$.

Notice that, since any tetrahedron of CDT spatial slices is adjacent to exactly 4 neighboring tetrahedra, the dual graphs are 4-regular (i.e. each vertex has degree 4), so that the adjacency matrix suffices to compute eigenvalues and eigenvectors ($L = 4 \cdot \mathbb{I} - A$), and furthermore it is sparse. In practice, we build and save the graphs
associated to each slice in the adjacency list representation. Being already a memory-efficient storage for the adjacency matrix of the graph, these structures can be directly fed to any numerical solver optimized for the computation of eigenvalues and eigenvectors of sparse, real and symmetric matrices. The spectra and eigenvectors analyzed in the present paper have been obtained using the ‘Armadillo’ C++ library [34] with Lapack, Arpack and SuperLU support for sparse matrix computation.

By solving the eigensystem for the LB spectrum, we can easily obtain eigenvectors as a side product. Even if the spectrum of a graph does contain much geometric information, still alone it is not capable to completely characterize geometries, but only classes of isospectral graphs. Conversely, the joint combination of eigenvalues and eigenvectors yields complete information on the graph\(^8\), but decomposed in a way useful for the analysis of geometries.

1. Spectral gap and connectivity

As argued above, geometric information about the large scales comes from the smallest eigenvalues and associated eigenvectors. The 0-th eigenvalue has a topological character, and in the general case its multiplicity tells us how many connected components the graph is composed of, but for connected graphs its role is trivial and uninteresting.

Arguably the most interesting eigenvalue is the first (non-zero) \(\lambda_1\), which, depending on the context, is called spectral gap or algebraic connectivity. The latter name comes from the observation that the larger the spectral gap \(\lambda_1\), the more the graph is connected.

A measure of connectivity for a compact Riemannian manifold \(\mathcal{M}\) is given by the Cheeger isoperimetric constant \(h(\mathcal{M})\) defined as the minimal area of a hypersurface \(\partial A\) dividing \(\mathcal{M}\) into two disjoint pieces \(A\) and \(\mathcal{M} \setminus A\)

\[
h(\mathcal{M}) \equiv \inf_{\partial A} \frac{\text{vol}(\partial A)}{\text{vol}(A)\text{vol}(\mathcal{M} \setminus A)},
\]

where the infimum is taken over all possible connected submanifolds \(A\).

For a graph \(G = (V, E)\), the Cheeger constant is usually defined by

\[
h(G) \equiv \min \left\{ \frac{|\partial A|}{|A|} \right\} \text{subject to } |A| \leq \frac{|V|}{2},
\]

where \(\partial A\) is the set of edges connecting \(A\) with \(V \setminus A\). The relation between the Cheeger constant and the spectral gap for a graph \(G\) where all vertices have exactly \(d\) neighbours is encoded in the Cheeger’s inequalities

\[
\frac{1}{2} \lambda_1 \leq h(G) \leq \sqrt{2d\lambda_1}.
\]

This property of the spectral gap is interesting for the analysis of geometries of slices in CDT, since, as we will see in the next section, it highlights different behaviors for the various phases.

B. Eigenvalue distribution and a toy model

When one considers the whole spectrum of the LB operator, two particularly interesting quantities are the density \(\rho(\lambda)\), defined so that \(\rho(\lambda)\,d\lambda\) gives the number of eigenvalues found in the range \([\lambda, \lambda + d\lambda]\), and its integral \(n(\lambda)\), which gives the total number of eigenvalues below a given value \(\lambda\).

Both functions can be defined for single configurations (spatial slices) or can be given as average quantities over the Euclidean path integral ensemble. As we shall see, the latter quantity, \(n(\lambda)\), will prove particularly useful to characterize the properties of triangulations at different scales. It is an increasing function of \(\lambda\) and its inverse is simply the \(n\)-th eigenvalue \(\lambda_n\). We will usually show

\[^8\text{Recall that, by the spectral theorem, the LB matrix can be decomposed as } L = U\Lambda U^\dagger, \text{ where } \Lambda \text{ is the diagonal matrix of eigenvalues, and } U \text{ is the matrix with corresponding eigenvectors as columns. The adjacency matrix, which defines the graph, can be simply obtained as minus the off-diagonal part of the LB matrix.}\

\[^9\text{This observation is not restricted to Laplace matrices of graphs, but applies to the spectra of the Laplace operator in a general space.}\

\[^{10}\text{The more general case of graphs with multiple connected components can be easily treated by studying its components individually: the Laplace matrix can be put in block-diagonal form, one block for each component, so that its spectrum is the union of the individual spectra, and its eigenspaces are direct sums of the individual eigenspaces.}\

\[ n(\lambda) = \frac{\omega_d}{(2\pi)^d} V^{d/2} \lambda^{d/2} \]  

where \( V \) is the volume of the manifold (which is assumed to be finite, with or without a boundary), \( d \) is its dimensionality, and \( \omega_d \) is the volume of the \( d \)-dimensional ball of unit radius. As we shall better discuss below, Weyl law, even if asymptotic, is generally expected to hold with a good approximation in the range of \( \lambda \) for which one is not sensitive to the specific infrared properties (i.e. shape, boundaries and/or topology) of the manifold. How violations to the Weyl law emerge and how they can be related to a sort of effective dimension at a given scale will be one of the main points of our discussion.

In the following we shall consider the LB spectrum computed on discretized manifolds. It is therefore useful to start by analyzing a simplified and familiar model, consisting of a regular and finite 3-dimensional cubic lattice, with respectively \( L_x \), \( L_y \) and \( L_z \) sides along the \( x \), \( y \) and \( z \) directions. All lattice sites are connected with 6 nearest neighbors sites, with periodic boundary conditions in all directions: this is therefore the discretized version of a 3-dimensional torus. The Laplacian operator can be simply discretized on this lattice and its eigenvectors coincide with the normal modes of a corresponding system of coupled oscillators: they are plane waves having wave number \( \vec{k} = (k_x, k_y, k_z) \), with \( k_i = 2\pi m_i / L_i \) and \( m_i \) integers such that \(-L_i/2 < m_i \leq L_i/2\), so that

\[ \lambda_{\vec{m}} = 4\pi^2 \left( \frac{m_x^2}{L_x^2} + \frac{m_y^2}{L_y^2} + \frac{m_z^2}{L_z^2} \right). \]  

Determining \( n(\lambda) \) for a given \( \lambda \) now reduces to counting how many vectors \( \vec{m} \) exist such that \( \lambda_{\vec{m}} \leq \lambda \). That corresponds to finding the triplets of integer numbers, i.e. the cubes of unit side, within the ellipsoid of semi-axes \( R_i = \lambda^{1/2} L_i / (2\pi) \), with the constraint that \(-L/2 < m_i \leq L/2 \forall i \). The latter constraint expresses the particular (cubic) discretization that we have adopted for the 3-dimensional torus, i.e. the structure of the system at the UV scale: if \( \lambda \) is low enough so that \( R_i < L_i \forall i \), then we are not sensitive to such scale. On the other hand, the discretized structure of the eigenvalues expresses the finiteness of the system, i.e. the properties of the system at the IR scale: if we have also \( R_i \gg 1 \forall i \) then we are not sensitive to such scale either, and the counting reduces approximately to estimate the volume of the ellipsoid, so that

\[ n(\lambda) \simeq \frac{4\pi}{3} R_x R_y R_z = \frac{4\pi}{3} \frac{L_x L_y L_z}{(2\pi)^3} \lambda^{3/2}, \]  

which is nothing but Weyl law for \( d = 3 \).

In Fig. 2 we show the exact distribution of \( \lambda_n \) as a function of \( n/V \), for various choices of \( L_x \), \( L_y \) and \( L_z \). The tick line represents the Weyl law prediction, \( \lambda = 6\pi^2 (n/V)^{2/3} \). When \( n/V \to 1 \), all systems show similar deviations from the law, which are related to the common structure at the UV scale. The Weyl law is a very good approximation for lower values of \( n/V \), as expected, and actually down to very small values of \( n/V \) for the symmetric lattice where \( L_x = L_y = L_z = 50 \).

For the asymmetric lattices, instead, some well structured deviations emerge at low \( n/V \), where \( \lambda \) follows a Weyl-like power law which is typical of lower dimensional models and can be easily interpreted as follows. For the lattice with \( L_x = L_y = l = 15 \) and \( L_z = 600 \), one
does not find any eigenvalue with \( m_x \neq 0 \) and \( m_y \neq 0 \) as long as \( \lambda < 4\pi^2/l^2 \simeq 0.175 \), therefore in this range the distribution of eigenvalues is identical to that of a one-dimensional system, for which \( \lambda \propto (n/V)^{1/2} \); for \( \lambda > 4\pi^2/l^2 \) also eigenvalues for which \( m_x \) and/or \( m_y \) are non-zero appear, and their distribution goes back to the standard 3-dimensional Weyl law. Making a wave-mechanics analogy, at low energy only longitudinal modes are excited, while transverse modes are frozen until a high enough energy threshold is reached. The point where one crosses from one power law behavior to the other brings information about the size of the shorter transverse scale. Similar considerations apply to the lattice \( L_x = 3 \), \( L_y = 75 \) and \( L_z = 600 \), which has three different and well separated IR scales: in this case one sees a one-dimensional power law for small \( n/V \), which first turns into a two-dimensional one as modes in the \( y \)-direction start to be excited, and finally ends up in a standard 3d Weyl law when also modes with \( m_x \neq 0 \) come into play.

The argument above can be rephrased at a more general level. Suppose we have a \( D \)-dimensional manifold where \( d \) “transverse” dimensions are significantly shorter than the other \( D - d \) “longitudinal” dimensions, with a typical transverse scale \( l \). As long as one considers small eigenvalues, the modes in the transverse directions will not be excited, so that the counting of eigenvalues will be given by the Weyl law for \( D - d \) dimensions, i.e. \( n(\lambda) = \omega_{D-d}(V/l^d)\lambda^{(D-d)/2} \). The change from one regime to the other will take place when the transverse directions get excited for the first time, i.e. at \( \lambda \simeq \pi^2/l^2 \) (the actual prefactor depends on the details of the shorter dimension), which corresponds to \( n \propto V^{-D} \), with a proportionality constant which depends only on the details of the short transverse scales and is independent of the details of the longer scales. Therefore, different manifolds, sharing the same structure at short scales associated with an effective dimensional reduction, lead to a distribution \( \lambda_n \) where the change from one power law behavior to the other takes place at the same point in the \((n/V)\lambda\) plane, where \( V \) is the global volume of the manifold. The value of \( n/V \), being proportional to \( l^{-D} \), brings information about the size of the short scale.

To better illustrate the concepts above, in Fig. 3 we show the distribution of \( \lambda_n \) as a function of \( n/V \) for three different choices of \( L_x, L_y \) and \( L_z \). The curves obtained for \((L_x,L_y,L_z) = (3,75,600)\) and \((L_x,L_y,L_z) = (3,75,1200)\) go exactly onto each other: their short scale structure is the same and the function \( n(\lambda) \) just differs for different number of modes which are counted along the large dimension \( L_z \), however this difference disappears when one considers the scaling variable \( n/V \), leading to a perfect collapse. The collapse instead is not perfect when one considers the lattice \((L_x,L_y,L_z) = (3,15,600)\), which has a different “intermediate” scale: moving from large to small \( n/V \), the turning point from dimension 3 to dimension 2 is the same as for the two other lattices, however the turning point from dimension 2 to dimension 1 takes place earlier, because \( L_y \) is shorter.

The possible examples which one can discuss within the toy model are quite limited. For instance, one cannot consider the case in which there are points where the manifold branches into multiple connected ramifications, something which in general can lead to an increase, instead of a decrease, of the effective dimension. However, extrapolating the arguments given above, we can conjecture the following. \( D \)-dimensional manifolds having different overall volumes and shape, but sharing a similarity in the structures which are found at intermediate and short scales, will lead to similar (i.e. collapsing onto each other) curves when \( \lambda_n \) is plotted against \( n/V \), \( V \) being the total volume of the manifold. Moreover, the power law taking place at a given value of \( n/V \) will give information about the effective dimensionality \( d_{EFF} \) of the manifold at a scale of the order \((n/V)^{-1/D} \), with

\[
\frac{2}{d_{EFF}} = \frac{d\log \lambda}{d\log(n/V)} .
\]

This kind of information is similar to what is obtained by implementing diffusive processes to measure the spectral dimension.

IV. NUMERICAL RESULTS

In this section we present results regarding mostly the spectrum of the LB operator defined on spatial slices, while a detailed discussion regarding the eigenvectors is postponed to a forthcoming study. We performed the analysis on spatial slices of configurations in each phase; in particular, almost all the results shown come from simulations running deep into each phase, at the points circled and labeled by a letter in Fig. 4 and in Table 4.

While the total spatial volume has been fixed in each simulation to a target value, the spatial volume of single slices, \( V_S \), can vary greatly from one slice to the other (apart from phase \( B \)). That will permit us to access the dependence of the spectrum on \( V_S \), an information that will be very important for many aspects. As discussed above, each spatial slice will be associated with a 4-regular undirected graph, with each vertex of the graph corresponding to a spatial tetrahedron. For this reason, it will be frequent in the following discussion to borrow concepts and terminology from graph theory.

We will first look at the low lying part of the spectrum, show how the transition from one phase to the other can be associated to the emergence of a gap in the spectrum, and discuss what that means in terms of the geometrical properties of the triangulations. We will then turn to results regarding the whole spectrum and show how one can obtain information on the effective dimension of the geometry at different scales. Finally, we will describe two methods to visualize graphs and apply them to show the appearance of spatial slices.
A. The low lying spectrum and the emergence of a gap

Apart from the zero eigenvalue, \( \lambda_0 = 0 \), the remaining eigenvalues will fluctuate randomly from one configuration to the other and, moreover, their distribution will depend on \( V_S \) in a well defined way that we are going to discuss later on. As an example, in Fig. 4 we show the distribution of \( \lambda_1 \) and \( \lambda_3 \) on a set of around \( 3 \times 10^3 \) slices of approximately equal volume \( V_S \simeq 2300 \) and in \( C_{dS} \) phase. Therefore, while the spectrum of each spatial slice is intrinsically discrete, because of the finite number of vertices making up the associated graph, it makes sense to define a continuous distribution \( \rho(\lambda) \), assigned so that \( \rho(\lambda) d\lambda \) gives back the number of eigenvalues which are found on average in the interval \( [\lambda, \lambda + d\lambda] \). In general \( \rho(\lambda) \) will be a function of the bare parameters chosen to sample the triangulations and, for fixed parameters, of the spatial volume \( V_S \) of the chosen slice.

In Figs. 5 and 6 we show the low lying part of the distribution \( \rho(\lambda) \) obtained from simulations performed respectively in the \( C_{dS} \) and \( B \) phases, selecting in each case three different ranges of spatial volumes \(^{11}\). In order to focus just on the low part of the spectrum, we have limited the input for \( \rho \) to just the first few eigenvalues in each case (\( n \leq 100 \)).

A striking difference between the two phases emerges. In the \( B \) phase there is a gap \( \Delta \lambda = \lambda_1 \simeq 0.1 \) which does not disappear and is practically constant as the spatial volume \( V_S \) grows, i.e. as one approaches the thermodynamical limit. This gap is absent in the \( C_{dS} \) phase, where the distribution of the first 100 eigenvalues is instead more and more squeezed towards \( \lambda = 0 \) as \( V_S \) grows. The presence or absence of a gap in the spectrum is a characteristic which distinguishes different phases in many different fields of physics: think for instance of Quantum Chromodynamics, where the absence/presence of a gap in the spectrum of the Dirac operator distinguishes between the phases with spontaneously broken/unbroken chiral symmetry. Let us discuss what is the meaning of the gap in our context.

Graphs which maintain a finite gap as the number of vertices goes to infinity are known as expander graphs \(^{37}\) and play a significant role in many fields, e.g., in computer science. They are characterized by a high connectivity, i.e. the boundary of every subset of vertices is generically large. Such a high connectivity is usually as-

\(^{11}\) For the \( B \) phase, different spatial volumes correspond actually to different simulations with different constraint on \( N_{41} \), since in this case most of the spatial volume is contained in one single slice.
associated with a degree of randomness, i.e. lack of order, in the connections between vertices: for instance, random regular graphs are expanders with high probability [38]. The strict relation between the high connectivity and the presence of a finite gap in the spectrum is also encoded in Cheeger’s inequalities, see Eq. (13).

The property which is maybe most relevant to our context is the fact that the diameter of an expander, defined as the maximum distance between any pair of vertices, does not grow larger than logarithmically with total number of vertices [39, 40]. Therefore, in this phase the spatial slices do not develop a well defined geometry, since the size (diameter) of the Universe remains small as the volume tends to infinity, a fact described also in previous CDT studies in terms of a diverging Hausdorff dimension. This fact can be easily interpreted in terms of diffusive processes: as argued above (see Section III), the value of the spectral gap, $\lambda$, can be interpreted as the inverse of the diffusion time of the slowest mode; the fact that the time to diffuse through the whole Universe stays finite means that its size is not growing significantly.

On the contrary, according to the arguments discussed in Section III B for a graph representing a standard manifold having a finite effective dimension on large scales, one expects that the number of eigenvalues found below any given $\lambda$ should grow proportionally to the volume $V_S$, $n \propto V_S \lambda^{d_{	ext{eff}}/2}$, see for instance Eq. (17). That means that the gap must go to zero as $V_S \to \infty$ and, moreover, that a finite normalized density of eigenvalues, $\rho(\lambda)/V_S$, must develop around $\lambda = 0$. Instead, as it will be shown in more detail below, the presence of a spectral gap for slices in the $B$ phase indicates that the effective dimension is indeed diverging at large scales, in agreement with the high connectivity property.

As an independent check, we computed the maximum distance from a randomly chosen vertex to all other vertices in the graph (a quantity usually called the eccentricity of the vertex), iterating the procedure for 200 different starting vertices and for each slice in the $\mathcal{C}_{40}$ and $B$ phases. The maximum eccentricity in a graph corresponds to its diameter, so the eccentricity of a random vertex is actually a lower bound to the diameter. Therefore the results, which are shown in the form of a scatter plot in Fig. 7, are consistent with a diameter which, for sufficiently large volumes, grows as a power law of $V_S$ in phase $\mathcal{C}_{40}$, while on the contrary it seems to reach a constant or to grow at most logarithmically in the $B$ phase.

The properties of slices in phase $A$ are quite similar to those found in phase $\mathcal{C}_{40}$, i.e. one has evidence for a finite density of eigenvalues around $\lambda = 0$ in the large $V_S$ limit, even if the distribution of slice volumes is significantly different from that found in phase $\mathcal{C}_{40}$. An example of the distribution of the first 30 eigenvalues in this phase is reported in Fig. 8.

Instead, the spectra of slices in the bifurcation phase $C_b$ need a separate treatment. Indeed, it is well known that the bulk of the configurations are made up of two

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12 The distance between a pair of vertices is defined as the length of the shortest path (i.e. the geodesic) connecting the two vertices.

13 A finite density of eigenvalues around $\lambda = 0$ is a condition stronger than the simple absence of a gap. Indeed, one might have situations in which isolated quasi-zero eigenvalues develop, while the continuous part of the spectrum maintains a gap: think for instance of two expander graphs connected by a thin bottleneck.
FIG. 9: Spectral gap $\lambda_1$ as a function of the slice-time for single configurations in $C_b$ and $C_{dS}$ phases with total spatial volume $V_{S,tot} = 40k$ and with the slice-time of maximal slice shifted to zero. Only slices in the bulk (with volume $V_S \geq 200$) have been shown.

FIG. 10: Averages of $\lambda_1, \lambda_20$ and $\lambda_{100}$ as a function of the slice-time for configurations in $C_b$ and $C_{dS}$ phases, where the slice-time of maximal slices has been shifted to zero. Only slices in the bulk (with volume $V_S \geq 200$) have been shown.

separate classes of slices, which alternate each other in slice-time and have different properties: it is reasonable to expect that this is reflected also in their spectra. This is indeed the case, as can be appreciated by looking at Fig. 9, where we report the value of $\lambda_1$ obtained on the different slices (i.e. at different Euclidean times) for a typical configuration sampled in the $C_b$ phase, and compare it to a similar plot obtained for the $C_{dS}$ phase. For an easier comparison, the time coordinates of the slices have been relabeled in each case so that the slice with the largest volume corresponds to $t_{slice} = 0$; moreover, we restricted to the bulk of configurations (i.e. we chose slices with $V_S > 200$). Contrary to the $C_{dS}$ phase, in the $C_b$ phase $\lambda_1$ changes abruptly from one slice to the other, with small values alternated with larger ones, differing by even two order of magnitudes. This striking difference, which emerges even for single configurations, is even more clear as one considers the whole ensemble: Fig. 10 shows the average of $\lambda_1$, $\lambda_{20}$ and $\lambda_{100}$ for configurations in the $C_b$ and $C_{dS}$ phases, with slice times relabeled as before. In the $C_{dS}$ phase $\lambda_1$ changes smoothly with $t_{slice}$, and this change is mostly induced by the corresponding change of the slice volume, while in the $C_b$ phase the alternating structure is visible also for higher eigenvalues, even if somewhat reduced and limited to the central region as $n$ grows.

Therefore, we conclude that the alternating structure of spatial slices is apparent and well represented in the low-lying spectra: slices in the bulk of $C_b$ phase configurations can be separated in two distinct classes by the value of their spectral gap, while in the $C_{dS}$ phase there is no sharp distinction apart from a volume-dependent behavior connected to an observed Weyl-like scaling, which will be discussed in more detail in Section IV.B.

In order to get a better perspective on these results, in Fig. 11 we show the eigenvalues $\lambda_n$, with $n = 1, 20, 100$, plotted against the volume of the slice on which they are computed, for the slices of all configurations sampled in the $C_b$ phase (in particular at the simulation point labeled $\tilde{c}$). Slices with volumes larger than a given $V_S$, which we call bifurcation volume$^{14}$, divide in two distinct classes characterized by $\lambda_n$ taking values in well separated ranges. It is interesting that such bifurcation volume depends on $n$: that also explains why in Fig. 10 the alternating behavior of higher order eigenvalues (e.g., $\lambda_{100}$) drops off earlier than lower order ones, since spatial volumes get smaller far from the slice with maximal volume and then their volume becomes less than the bifurcation one at that order. That actually means that the alternating slices found in the $C_b$ phase only differ for the low lying part of the LB spectrum, while for high enough eigenvalues order they are not distinguishable; high eigenvalues mean small scales, hence we expect that the alternating slices have the same small scale structures and only differ at large scales. We will come back on this point later on.

In view of the close similarity with the properties of slices found respectively in the $C_{dS}$ in the $B$ phase, we assign to the two classes of slices the names $dS$-type (low spectral gap) and $B$-type (high spectral gap). Looking again at Fig. 11 we notice that, for sufficiently large volumes, the two classes populate only specific volume ranges. Furthermore, the maximal slice in the $C_b$ phase is typically observed to be of $B$-type, with a volume ranging in a narrow interval which is separated from the volumes of the other slices. This alternating distribution of spatial volumes has been indeed one of the first signals of the

$^{14}$ It is interesting no notice that the $C_b$ phase can be called bifurcation phase for many different reasons.
presence of the new phase \[3, 4].

A summary of the characterization of the phase diagram of CDT according to the (zero or non-zero) gap of the LB operator as an order parameter is reported in Table II. To conclude the discussion about the gap, it is interesting to consider how the distribution of \(\lambda_1\) changes across the different phases. To this purpose, in Fig. 12 we show a scatter plot of \(\lambda_1\) for different values of \(\Delta\) at fixed \(k_0 = 2.2\): darker points corresponds to more frequent values of \(\lambda_1\). As \(\Delta\) increases, the gap in the \(B\) or \(B\)-type slices progressively reduces and approaches zero at the point where one enters the \(C_{dS}\) phase. A gap in the spectrum is a quantity which has mass dimension two (as the LB operator), i.e. an inverse length squared; if future studies will show that the drop to zero takes place in a continuous way, that will give evidence for a second order phase transition with a diverging correlation length.

B. Scaling and spectral dimension

As one expects, and as it emerges from some of the results that we have already shown, the typical values obtained for the \(n\)-th eigenvalue of the LB operator on spatial slices, \(\lambda_n\), scale with the volume \(V_S\) of the slice, and in a different way for the different phases. As an example, in Fig. 13 we show the average values obtained for \(\lambda_n\) (for a few selected values of \(n\)), as a function of the volume, in the \(C_{dS}\) phase.

In order to better interpret this scaling, and inspired by the discussion reported in Section III B, in the following we will consider how \(\lambda_n\) depends on the variable \(n/V_S\). To show that this may indeed be illuminating, in Fig. 14 we report \(\lambda_n\) as a function of \(n/V_S\) for four spatial slices, which have been randomly picked from an ensemble produced in the \(C_{dS}\) phase and have quite different

| \(A\)  | \(B\)  | \(C_b\) | \(C_{dS}\) |
|-------|-------|--------|---------|
| no-gap | gap   | gap    | no-gap  |

TABLE II: Characterization of the phase diagram of CDT according to the zero or non-zero gap of the LB operator as an order parameter.
Notice that of structures at intermediate common scales. They have quite different extensions, they show the same kind of scaling and, in view of the discussion in Section III B, can be interpreted in this way: despite the fact that the slices of phase $A$ and $B$ that we have divided all spatial slices in small volume bins, and then averaged $\lambda_n$ for each $n$ over the slices of each bin: such averages are reported in the figure against $n/V_S$: average eigenvalues are reported with error bars, which however are too small to be visible.

Each phase has its own characteristic profile. The profiles of phases $A$ and $C_{dS}$ are quite similar and differ by tiny deviations; in particular, in both cases one has that $\lambda_n \to 0$ as $n/V_S \to 0$, which is an equivalent way to state the absence of a gap in the spectrum. Instead, the profile of phase $B$ is significantly different and characterized by the fact that $\lim_{n/V_S \to \infty} \lambda_n \neq 0$, in agreement with the presence of a gap. In Fig. 15 we do not report any data regarding the $C_3$ phase, which is discussed separately because of the particular features that we have already illustrated above.

Following the discussion in Section III B each scaling profile can be associated with a running effective dimensionality $d_{EFF}$ of the spatial triangulations at a scale of the order $(n/V_S)^{-1/3}$; that can be done by taking the logarithmic derivative of $\lambda_n$ with respect to $n/V_S$, see Eq. (17). For this reason, in Fig. 16 we report $d_{EFF} = 2 \log(n/V_S)/d\log \lambda_n$, which has been computed numerically by taking the average derivative of the profile over small bins of the variable $n/V_S$.

At very small scales, both the $A$ and the $C_{dS}$ phase are effectively 3-dimensional. However, going to larger scales (smaller $n/V_S$), the effective dimension decreases, going down to values around $d_{EFF} \sim 1.5$, which is approximately the same large scale dimensionality observed by diffusion processes [41]. The crossover between the

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**FIG. 14:** Plot of $\lambda_n$ against its volume-normalized order $n/V_S$, for four randomly selected slices with volumes $V_S \simeq 500, 1000, 2000, 3000$, taken from configurations sampled deep into the $C_{dS}$ phase (simulation point c) with total spatial volume $V_{S,tot} = 40k$.

**FIG. 15:** Averages of $\lambda_n$ versus $n/V_S$ computed in bins of $n/V_S$ with size $2/V_{S,max}$ for slices taken from configurations sampled deep into the $A$, $B$ and $C_{dS}$ phases (simulation points $a$, $b$ and $c$). The volume is fixed to $V_{S,tot} = 40k$ for configurations in $A$ and $C_{dS}$ phase, and to $V_{S,tot} = 8k$ for configurations in $B$ phase.

**FIG. 16:** Running dimension obtained from the logarithmic slope $m$ of the curves shown in Fig. 15 as $2/m$ (see Section III B and Eq. (17)), computed over bins of different ranges of $n/V_S$ and for configurations sampled in phases $C_{dS}$, $A$ and $B$. The curve associated to the $B$ phase is diverging for $n/V_S \to 0$ (it is around 30 for $n/V_S \sim 10^{-4}$), but part of it has been omitted from the plot, to improve the readability of the curves obtained for the other two phases.

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15 Notice that $n/V_S$ can take values in the range $(0,1)$ (recall that $\lambda_0 = 0$ is excluded from our discussion), while the maximum eigenvalue $\lambda$ is always bounded by $2k = 8$, that is twice the degree of vertices in the $k$-regular graph.
two regimes takes place for \( n/V_S \) in the range \( 0.1 - 0.4 \), meaning that typical structures of lower dimensionality develop, with a transverse dimension of the order of just a few tetrahedra.

Actually, the plot of \( d_{EFF} \) shows a difference between phase \( A \) and phase \( C_{dS} \), which was not clearly visible before: contrary to phase \( A \), in phase \( C_{dS} \) the effective dimensionality seems to slowly grow again as one approaches larger and larger scales. This slow grow can be interpreted as a progressive ramification of the lower dimensional structures, i.e. as a hint that it has a fractal-like nature.

The effective dimensionality has a completely different behavior in phase \( B \): it is smaller than 3 (\( d_{EFF} \approx 2.5 \)) on small scales, then starts growing and diverges at large scales. This is due to the fact that \( d \log \lambda_n/d \log (n/V_S) \rightarrow 0 \) as \( n/V_S \rightarrow 0 \), because of the presence of the gap, and, on the other hand, the diverging dimensionality can be interpreted in terms of the fact that the diameter of the slice grows at most logarithmically with \( V_S \). Also the low dimensionality observed at small scales can be interpreted in terms of the large connectivity of the associated graphs: each tetrahedron has 4 links to other tetrahedra, some of these links can, in some sense, not “local”, i.e. they are a shortcut to reach directly some otherwise “far” tetrahedron; then, the probability that a couple of neighbouring tetrahedra are adjacent to a common tetrahedron gets smaller and leads to a lower effective dimensionality at short scales.

Regarding the properties of the slices found in the bifurcation phase \( C_0 \), on the basis of what we have shown and discussed in Section [IV.A] we have decided to perform a separate analysis for the different classes of spatial slices. In Fig. 17 we report \( \lambda_n \) vs. \( n/V_S \) for slices according to their relative position with respect to the central largest \( B \)-type slice (which corresponds to \( t_{slice} = 0 \)). The differences between the two classes is clearly visible also from the scaling profiles, which resemble, especially for large scales, those found in the \( B \) and in the \( C_{dS} \) phase for \( B \)-type and \( dS \)-type slices, respectively.

However, one striking feature emerges: at small scales, in particular for \( n/V_S \gtrsim 0.1 \), the scaling profiles coincide almost perfectly. We conclude that, at such scales, the two classes of slices present strong similarities, despite the completely different large scale behavior. Hints of this fact were already discussed in Section [IV.A] Such similarities are likely induced by the causal structure connecting adjacent spatial slices in CDT triangulations.

**C. Running scales and the search for a continuum limit**

The analysis of the scaling profiles reported above permits to identify well defined scales, in terms of the parameter \( n/V_S \), where something happens, like a change in the effective dimensionality of the system. Such scales are given in units of the elementary lattice spacing of the system, i.e. the size of a tetrahedron.

On the other hand, the possible presence of a second order critical point, where a continuum limit can be defined for Quantum Gravity, implies that the lattice spacing should run to zero as the bare parameters approach the critical point. This running of the lattice spacing should be visible by the corresponding growth of the value, determined in lattice units, of some physical scale. This is a standard approach in lattice field theories, where one usually considers correlations lengths which are the inverse mass of some physical state.

One of the major challenges in the CDT program is to identify and determine physical scales which could pro-
vide such kind of information and thus give evidence that the lattice spacing is indeed running. Promising steps in this direction have been already done by means of diffusive processes, where the scale is fixed by the diffusion time, both in CDT [9] and in DT [21]. Here we propose that LB spectra and the observed scaling profiles may be helpful in this direction, and that a careful study of how such profiles change as a function of the bare parameters could provide useful information.

A possible second order point is believed to separate the $C_b$ from the $C_{ds}$ phase, therefore it makes sense to analyze how the profiles change in both phases when moving towards the supposed phase transition, and if the observed changes can be associated to any running scale. A growth in the scale associated to some particular feature of the scaling profile means that its location moves to smaller values of $n/V_S$.

As an example, in Fig. 18 we report the scaling profiles obtained for slices in phase $C_b$ and $\Delta = 0.15$, which is closer to the phase boundary with respect to the case $\Delta = 0.10$, which has been discussed previously and is reported in Fig. 17. An appreciable difference between the two cases is that the region where the profiles of $B$-type and $dS$-type coincide is larger (i.e. extends to smaller $n/V_S$) for $\Delta = 0.15$. From a quantitative point of view, one finds that the approximate value of $n/V_S$ where the profiles start differing by more than 5% is around 0.13 for $\Delta = 0.10$ and around 0.074 for $\Delta = 0.15$. In other words, there is a scale up to which $B$-type and $dS$-type slices are similar to each other, and such scale grows as one approaches the $C_b$-$C_{ds}$ phase transition.

In a similar way, one can look at how the scaling profiles found in the $C_{ds}$ phase change as one approaches the phase transition from the other side. Such scaling profiles are reported in Fig. 19. The short-scale region, and in particular the point where the effective dimension starts changing, seems not sensible to the change of $\Delta$. However the small $n/V_S$ (large-scale) region changes, with the profile undergoing an overall bending towards the left: notice that this implies a change in the effective dimensionality observed at the largest scales, which indeed, for $\Delta = 0.3$, is $d_{\text{EFF}} \gtrsim 2$.

Finally, as we have already stressed above, the gap itself, which for $B$-type slices seems to approach zero as one gets closer to the $C_b$-$C_{ds}$ phase transition (see Fig. 12) could be interpreted in terms of a diverging correlation length if the behavior is proved to be continuous.

The reported examples are only illustrative of the fact that the LB spectrum can provide useful scales which could give information on the nature of a possible continuum limit. Such program should be carried on more systematically by future studies, in particular by approaching the $C_b$-$C_{ds}$ phase transition more closely.

![Fig. 19: Averages of $\lambda_n$ versus $n/V_S$, computed in bins of $n/V_S$ with size $2/V_{S,max}$, for slices taken from configurations sampled in the $C_{ds}$ phase, with $k_0 = 2.2$ and different values of $\Delta$. The total spatial volume of each configuration is $V_{S,tot} = 40k$.](image)

**D. Fine structure of the full spectrum**

In this section we will show some details regarding the full distribution of eigenvalues (i.e. over the whole spectrum) in the different phases. Figs. 20, 21 and 22 show the normalized distribution of eigenvalues for spatial slices with volumes in selected ranges, and for simulations performed deep into the phases $C_{ds}$, $A$ and $B$ respectively.

The $A$ and the $C_{ds}$ phase present a detailed non-trivial fine structure which is very similar. Even if we are not interested, at least in the present context, to provide a detailed interpretation of the full spectrum, we notice that such fine structure is mostly relative to eigenvalues which are of order 1 or larger, hence associated to typically small scales; this is confirmed by the fact that, contrary to the low part of the spectrum, such fine structure is almost left invariant by changing the volume of the slice. For instance, it can be noticed that the distributions are sharply peaked around the integer values $\lambda = 4, 5, 6$; indeed, by inspecting the spectra of single configurations and the associated eigenvectors, we observed that these integer eigenvalues often occur with high multiplicity and can be associated to the presence of recurrent regular short-scale structures and to very localized eigenvectors.

The normalized distribution in the case of configurations in the $B$ phase does not show particular features, other than the already discussed presence of a spectral gap. The distribution looks in general more regular in this phase, even if some of the peaks around integer values are still present, but much reduced in amplitude.
FIG. 20: Normalized distribution of all the eigenvalues for slices with volume in the range $V_S \in [2000, 2500]$ for configurations deep into the $C_{ds}$ phase (simulation point $c$), and with total spatial volume $V_{S,\text{tot}} = 40k$. 

FIG. 21: Normalized distribution of all the eigenvalues for slices with volume in the range $V_S \in [200, 400]$ for configurations deep into the $A$ phase (simulation point $a$), and with total spatial volume $V_{S,\text{tot}} = 40k$. 

FIG. 22: Normalized distribution of all the eigenvalues of the maximal slices for configurations deep into the $B$ phase (simulation point $b$), and with spatial volume about $V_S \approx 8k$.

E. Visualization of spatial slices

We have seen how each spatial slice of the triangulations can be associated to a graph with non-trivial properties, i.e. what is usually called a complex network. There are different methods to visualize a complex network, some of them already considered in previous studies (see, e.g., Ref. [21]), here we will briefly discuss only two of them: Laplace embedding [42] and spring embedding [43, 44]. The former makes use of the eigenvectors associated to the smallest eigenvalues, which are already computed by solving the eigenvalue problem, while the latter is based on a mapping of the graph to a system of points connected by springs: as we are going to discuss, the two methods are strictly related, however spring embedding proves more useful to give an intuitive picture of the short-scale structures.

The underlying idea, common to both methods, is to represent any graph $G = (V, E)$ in a $m$-dimensional Euclidean space by finding a set of $m$ independent functions $\{\phi_n(v_i)\}_{n=1}^{m}$ which act as coordinates for each vertex $v_i \in V$, in such a way that vertices with smaller graph distance have coordinates with values as closer as possible. The “closeness” can be defined in many ways, consisting in solving different optimization problems, and that makes the two methods different. We will use the notation $\vec{\phi}_n \equiv (\phi_n(v_i))_{i=1}^{V}$ for each $n = 1, \ldots, m$.

1. Laplace embedding

The optimization problem for Laplace embedding [42] consists in minimizing the following functional of the coordinate functions $\{\phi_n\}$:

$$E_{LB}[\phi] \equiv \frac{1}{2} \sum_{n=1}^{m} \sum_{(v_i, v_j) \in E} \left( \phi_n(v_i) - \phi_n(v_j) \right)^2 = \frac{1}{2} \sum_{n=1}^{m} \vec{\phi}_n^T L \vec{\phi}_n. \quad (18)$$

subject to the constraints $\vec{\phi}_n \cdot \vec{\phi}_k = \delta_{n,k}$ and $\vec{\phi}_n \cdot \vec{1} = 0$ for each $n, p = 1, \ldots, m$, where $\vec{1}$ is the uniform vector with unit coordinates and $L$ is the matrix representation of the LB operator. It is straightforward to prove that a solution to this constrained optimization problem is given by the set of the first $m$ eigenvectors $\{\vec{e}_n\}_{n=1}^{m}$ of the Laplace–Beltrami matrix, where we excluded the 0-th mode $\vec{e}_0 = \frac{1}{\sqrt{|V|}} \vec{1}$ (second constraint) and ordered eigenvectors without multiplicity (i.e. $\vec{e}_n$ is associated to $\lambda_n$ and $\lambda_n \leq \lambda_{n+1}$).
For example, the coordinates associated to each vertex $v \in V$ in a 3-dimensional Laplace embedding are the values of the first 3 eigenvectors on that vertex, that is $v \mapsto (e_1(v), e_2(v), e_3(v)) \in \mathbb{R}^3$. Fig. 23 shows the 3-dimensional Laplace embedding of a typical slice in the bulk of a configuration deep in the $C_{dS}$ phase (simulation point c). The geometry seems to be made up of filamentous structures, but that really means that the first 3 eigenvectors, describing the slowest modes of diffusion, are not capable of describing short scale structures inside the filaments. However, they efficiently describe the largest scale geometry, which in the $C_{dS}$ case is non-trivial and unexpected.

2. **Spring embedding**

The optimization problem that has to be solved for spring embedding of an unweighted undirected graph $G = (V, E)$ consists in the energy minimization of a system of ideal springs with fixed rest length $l_0$ and embedded in $\mathbb{R}^m$, with extrema connected in the same way as the links of the abstract graph $G$. Having assigned coordinates $\{\phi_n(v_i)\}_{n=1}^m$ to each abstract vertex of the graph $v_i \in V$, the potential energy of the system is defined as:

$$\mathcal{E}_S[\phi] = \frac{1}{2} \sum_{(v_i, v_j) \in E} \left( l_0 - \sqrt{\sum_{n=1}^m (\phi_n(v_i) - \phi_n(v_j))^2} \right)^2$$

(19)

In the limit $l_0 \to 0$, the functional $\mathcal{E}_S$ becomes equal to $\mathcal{E}_{LB}$ but with no constraint, so that the solution would collapse to the trivial solution, bringing all vertices to the same point, in this limit. On the other hand, for $l_0 > 0$, the springs will push vertexes apart from each other and help resolving even the shortest-scale structures, which are not visible with Laplace embedding.

The simplest algorithm to find a (local) minimum is to initialize the coordinates of each vertex to a random value, and then relax the system of springs by performing a gradient descent. Fig. 24 shows the spring embedding of the same slice represented by Laplace embedding in Fig. 23. The large scale structure is well represented by both methods, but spring embedding permits to better discern short-scale structures at the finest level.

Such representations of the spatial slices are illuminating to understand the properties of the LB spectrum for $C_{dS}$ slices. The slices are extended objects, i.e., one finds vertexes which are far apart from each other, implying the existence of slow diffusion modes and a continuum of quasi-zero eigenvalues for large $V_S$. On the other hand, the large scale structure is made of lower-dimensional
substructures, which have a typical transverse size of the order of a few vertexes, and which often branch, making the overall spectral dimension (i.e. the diffusion rate) fractional at large scales.

For comparison, Fig. 25 shows the spring embedding of a typical slice in the $B$ phase. The high connectivity of the graph, which is clearly visible from the figure, does not permit the development of extended large scale structures, so that diffusion modes maintain always fast and a finite gap remains even in the $V_S \to \infty$ limit.

V. DISCUSSION AND CONCLUSIONS

In this work we have investigated the properties of the different phases of CDT that can be inferred from an analysis of the spectrum of the Laplace-Beltrami operator computed on the triangulations. The present exploratory study has been limited to the properties of spatial slices: those can be associated to regular graphs where each vertex is linked to 4 other vertexes. Let us summarize our main results and further discuss them:

i) We have shown that the different phases can be characterized according to the presence or absence of a gap in the spectrum, which therefore can be considered as new order parameter for the phase diagram of CDT. In particular, a gap is found in the $B$ phase, while for the $A$ and the $C_{ds}$ phases one finds a non-zero density of eigenvalues around $\lambda = 0$ in the thermodynamical (large spatial volume $V_S$) limit. The $C_b$ phase, instead, shows the alternance of spatial slices of both types (gapped and non-gapped): that better characterizes the nature of the alternating structures already found in previous works [5, 6], which for this reason we have called $B$-type and $dS$-type slices.

The presence or absence of a gap in the spectrum is a characteristic which distinguishes different phases in many different fields of physics: think for instance of Quantum Chromodynamics, where the absence/presence of a gap in the spectrum of the Dirac operator characterizes the phases with spontaneously broken/unbroken chiral symmetry.

In this context, the presence of a gap tells us that the spatial slices are associated to expander graphs, characterized by a high connectivity. That can be interpreted geometrically as a Universe with an infinite dimensionality at large scales, with a diameter which grows at most logarithmically in the thermodynamical limit; a small diameter in the phases with a gap is consistent with the findings of previous studies and is supported by a direct computation (see Fig. 7). On the contrary, the closing of the gap can be interpreted as the emergence of a Universe with a standard finite dimensionality at large scales. It is interesting to notice that the value of the gap which is found seems to change continuously as one moves from the $B$ to the $C_b$ phase, and approaches zero as the $C_{ds}$ phase is approached.

ii) We have shown that the spectrum can be characterized by a well defined scaling profile: the $n$-th eigenvalue, $\lambda_n$, is a function of just the scaling variable $n/V_S$. The profile is different for each phase and characterizes it; moreover, from the profile one can deduce information on the effective dimensionality $d_{EFF}$ of the system at different scales, which generalize a similar kind of information gained by diffusion processes.

The $C_{ds}$ and the $A$ phase share a similar profile, corresponding to $d_{EFF} \simeq 3$ at short scales, which then drops to $d_{EFF} \simeq 1.5$ for $n/V_S \lesssim 0.1$. At larger scales, the two phases show a different behavior, with $d_{EFF}$ which keeps decreasing as $n/V_S$ decreases in the $A$ case, while in the $C_{ds}$ phase it starts growing again at large scales. Slices in the $B$ phase, instead, show an effective dimensionality which, in agreement with their high connectivity, seems to diverge in the large scale limit.

An interesting feature has been found for the two different and alternating (in Euclidean time) classes of spatial slices in the $C_b$ phase: despite the different overall structure, they share an identical profile at small length scales, which is likely induced by the causality condition imposed on triangulations and is therefore an essential property of CDT. The profiles remain identical up to characteristic length scale above which they start to diverge, as expected since one class presents a gap and the other does not.
iii) We have proposed that the scaling profiles might be used to identify particular length scales which change as a function of the bare parameters, and thus could serve as possible probes of the running to the continuum limit, if any. Among those, we have found of particular interest the characteristic length scale up to which the alternating slices found in the $C_b$ phase share the same profile: we have seen that such length grows as one approaches the boundary with the $C_{dS}$ phase. On the other side of the boundary, also the profiles of the slices in $C_{dS}$ phase show a modification at large scales as the $C_b$ phase is approached, leading in particular to a growing effective dimensionality.

Along these lines, one could conjecture that, if a second order critical point is really found between the two phases, at such a point the different profiles found in the $C_b$ phase could merge at all scales and coincide with the profile from the $C_{dS}$ phase. Such a critical point would also been characterized by the vanishing of the gap for the $B$-type slices of the $C_b$ phase. Moreover, it would be interesting to test what the effective dimensionality found at large scales would be at the critical point: is it possible that, just on the critical point where a continuum limit can be defined, the effective dimensionality of spatial slices goes back to $D = 3$ at all physical scales?

The present work can be continued along many directions. First of all, the region around the transition between the $C_b$ and the $C_{dS}$ phase should be studied in much more detail than what done in the present exploratory work, to see if some of the conjectures that we have made above can be put on a more solid basis. In addition, a careful study of the critical behavior around the transition of the spectral gap, which is the new order parameter introduced in this study, could provide information about the universality class to which the continuum limit, if any, belongs. Of course, it could well be that one finds a first order transition, i.e. a sudden jump in the gap and in other properties, but then one should perform simulations for lines corresponding to different $k_0$ to see if the first order terminates at some critical endpoint.

We have not considered yet the information which can be gained by inspecting the eigenvectors of the LB operator, that will be done in a forthcoming study. In particular, it will be interesting to consider and analyze their localization/delocalization properties, in a way similar to what has been done in similar studies for the spectrum of the Dirac operator in QCD [45, 46].

It will be interesting to extend the study of the spectrum to the full triangulations, i.e. not just for spatial slices. That will require some implementative effort: unlike spatial tetrahedra (which are all identical), pentachorons can have edges with different Euclidean lengths, and therefore a regular graph representation does not describe the geometry faithfully. Nevertheless, the Laplace-Beltrami operator for general triangulated manifolds would have a well defined representation in the formalism of Finite Elements Method, as discussed and applied for example in Refs. [11] [12].

Finally, it would be interesting to apply spectral methods also to other implementations of dynamical triangulations, like the standard Euclidean Dynamical Triangulations (DT) where no causality condition is imposed. The implementation in this case would be straightforward, as for the spatial slices of CDT, i.e. given in terms of regular indirected graphs. We plan to address the issues listed above in the next future.

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Appendix A: Heat-kernel expansion and spectral dimension

Here we describe how to compute the spectral dimension of a graph $G$ from the spectrum of its LB matrix using the heat-kernel expansion. Later we will apply the definition to slices in $C_{dS}$ phase, making a comparison between the standard definition of spectral dimension (i.e. via diffusion processes), and the one obtained by the spectrum.

Let us consider the fundamental solution to the diffusion equation on a $k$-regular connected graph $G$ with LB matrix $L$:

$$
\begin{align*}
\partial_t K_{v,v_0}(t) &= -\frac{1}{k} \sum_{v' \in V} L_{v,v'} K_{v',v_0}(t) \\
K_{v,v_0}(0) &= \delta_{v,v_0}. 
\end{align*}
$$

Discretizing time with unit steps $\Delta t = 1$ [10], Eq. (A1) becomes the equation for random walk on the graph, where $K_{v,v_0}(\tau)$ is the probability that a random walker starting from the vertex $v_0$ at time $t = 0$ is found at the vertex $v$ at time $t = \tau$. The return probability $Z_{v_0}(t) = K_{v_0,v_0}(t)$ is the probability that a random walker comes back to the starting vertex $v_0$ after $t$ steps. Averaging over all starting vertices, the return probability reduces to $Z(t) = \text{Tr} K(t)$, the heat-kernel trace. In practice, the diffusion is performed only for a random subset of starting vertices, from which the return probability is then estimated as an average over explicit diffusion processes.

However, $Z(t)$ can also be computed using the spectrum of the LB matrix using the heat-kernel expansion for the solution to Eq. (A1):

$$
K_{v,w}(t) = \frac{1}{|V|} \sum_{n=0}^{|V|-1} e_n(v) e_n(w) e^{-\lambda_n t/k},
$$

(A2)
where \( \{\lambda_n\} \) and \( \{\tilde{e}_n\} \) are the eigenvalues and associated eigenvectors of the LB matrix of the graph \( G \). Notice that the terms in Eq. (A2) corresponding to larger eigenvalues are more suppressed for increasing times than terms corresponding to smaller ones. In particular, for times \( t \gg k/\lambda_1 \), the only surviving term is given by the 0-th eigenvalue, and the probability distribution tends to be uniformly distributed amongst all vertices: 

\[
\lim_{t \to +\infty} K_{v,v_0}(t) = \frac{1}{|V|} \quad \forall v, v_0 \in V \quad \text{(assuming a single connected component)}.
\]

The return probability, obtained from the spectrum, then takes the form

\[
Z(t) = \text{Tr} K(t) = \sum_{v \in V} K_{v,v}(t)
= \frac{1}{|V|} \sum_{n=0}^{|V|-1} e^{-\lambda_n t/k}, \quad (A3)
\]

where we used the decomposition in Eq. (A2) and the orthonormality of eigenvectors.

The return probability \( Z(t) \) can be nicely interpreted as a statistical partition function, for its formal analogy with the concept in statistical physics: the diffusion time takes here the role of the inverse temperature, while the eigenvectors and their associated eigenvalues take the role of microstates and their associated energies respectively.

In the case of a compact smooth manifold \( M \), for which the Laplace–Beltrami spectrum \( \{\lambda_n\}_{n=0}^{\infty} \) is countable but unbounded, the averaged return probability density \( Z(t) \) has the following asymptotic expansion for \( t \to 0^+ \):

\[
Z(t) = \frac{1}{\text{vol}(M)} \sum_{n=0}^{\infty} e^{-\lambda_n t} = (4\pi t)^{-\frac{\dim(M)}{2}} \frac{1}{\text{vol}(M)} \left( \sum_{i=0}^{t-1} c_i \sqrt{t} + O(t^{\frac{1}{2}}) \right). \quad (A4)
\]

The return probability for unidimensional random walks is \( 1/\sqrt{4\pi t} \), so it is reasonable for a smooth manifold to locally decompose the random motion along the \( \dim(M) \) directions and get the return probability as a product of independent unidimensional return probabilities. In the case of random walks on \( \mathbb{R}^d \) the return probability density equal to \( Z(t) = (4\pi t)^{-d/2} \), so one can infer the value of coefficients: \( c_0 = \text{vol}(M) \) and \( c_i = 0 \forall i \geq 1 \).

 Corrections to the \( t^{-\frac{\dim(M)}{2}} \) behavior must be due to the geometric properties characterizing the manifold under study. For example, the first three coefficient have a geometrical interpretation, as discussed by McKeane and Singer:

\[
c_0 = \text{vol}(M), \quad c_1 = -\frac{\sqrt{\pi}}{2} \text{area}(\partial M), \quad c_2 = \frac{1}{3} \int_M R - \frac{1}{6} \int_{\partial M} J, \quad (A5-A7)
\]

where \( \partial M \) is the (possibly empty) boundary of the manifold \( M \), \( R \) is the scalar curvature of the manifold and \( J \) is the mean curvature of the boundary.

 We expect that similar results hold for graphs approximating manifolds, but a first difficulty can be easily detected as shown by the following argument. At a time \( t \) only eigenvalues \( \lambda \lesssim \frac{1}{t} \) contribute to the sum in Eq. (A4), but for \( t \to 0^+ \) the full unbounded spectrum of the smooth manifold tends to contribute. The spectrum of a graph \( G \), however, is bounded by the largest eigenvalue, so that here the expansion in Eq. (A4) is not numerically reliable for times \( t \lesssim (\lambda_{|V|-1})^{-1} \). Nevertheless one can plot the return probability as a function of time and get an estimate of the dimension \( d \) by extrapolation to \( \tau \to 0^+ \) using the definition of what is called spectral dimension:

\[
D_S(\tau) \equiv -\frac{2}{d \log t} \frac{d \log Z}{d \log t} \bigg|_{t=\tau}. \quad (A8)
\]

Fig. [26] shows the comparison between the estimates of spectral dimension obtained employing explicit diffusion processes (Eq. (A1)) integrated with step size \( \Delta t = 1 \) and the spectrum of the Laplace–Beltrami matrix on graphs associated to spatial slices in \( C_{DS} \) phase: we applied Eq. (A8) using the average of the return probability \( Z(t) \) computed on each slice having volume in the range \( 2000-22000 \), and, for the definition via diffusion, averaging the return probability also over 200 iterations of diffusion processes starting from randomly selected vertices.
the slice. Using the definition via diffusion, at small diffusion times the return probability, and therefore also the spectral dimension, is highly fluctuating due to the short scale regularity of the tetrahedral tiling of the space (a phenomenon already discussed in Refs. [1][10]): this is not present in the definition via the spectrum, where a bump is observed instead. For larger diffusion times ($\tau \gtrsim 100$) the curves obtained using both methods agree even using only the lowest 5% part of the spectrum, which confirms that this regime represents indeed the large scale behavior. Here we observe a spectral dimension $D_s \approx 1.5$ for the spatial slices of configurations in $C_{48}$ phase. This fact, already observed in literature using diffusion processes [11], seems compatible also with the observations obtained from large scale scaling relations for the eigenvalues discussed in Section [IV.E].

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