Group Field theory and Tensor Networks: 
towards a Ryu-Takayanagi formula in full quantum gravity

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Abstract

We establish a dictionary between group field theory (thus, spin networks and random tensors) states and generalized random tensor networks. Then, we use this dictionary to compute the Rényi entropy of such states and recover the Ryu-Takayanagi formula, in two different cases corresponding to two different truncations/approximations, suggested by the established correspondence.

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I. INTRODUCTION

Background independent approaches to quantum gravity suggest a picture of the microstructure of the universe in which continuum spacetime and geometry disappear and are replaced by discrete and non-spatiotemporal entities. Among them, Loop Quantum Gravity (LQG) \cite{1-5}, the modern incarnation of the canonical quantization programme for the gravitational field, together with its covariant counterpart (spin foam models), and Group Field Theory (GFT) \cite{6-9}, a closely related formalism sharing the same type of fundamental degrees of freedom, identify this microstructure with (superpositions of) spin networks, which are graphs labeled by group-theoretic data. More precisely, in GFT models of quantum gravity spin network states arise as many-body states in a 2nd quantised context, whose kinematics and dynamics are governed by a quantum field theory over a group manifold with quanta corresponding to tensor maps associated to nodes of the spin network graphs. Random combinatorial structures, corresponding both to the elementary building blocks of quantum spacetime and to their interaction processes, become central. The same is true in the related context of random tensor models \cite{10-12}, which, for our present purposes can be seen as a simplified version of GFTs, stripped down of the group-theoretic data, leaving only the combinatorial aspects. Indeed, the random tensors can be understood as the GFT fields considered for the special case of a finite group. For a more detailed account of these three quantum gravity formalisms, and for the many results obtained, we refer to the cited literature. In the following, we will provide more precise definitions of their main ingredients.

Tensor networks, in recent years, have attracted a lot of attention as powerful quantum information tools in the context of condensed matter and, more generally, quantum many-body systems (including quantum field theory). For recent reviews, see \cite{13-14}. Also in this case, we will give precise definitions in the following. Here it suffices to say that tensor networks encode the entanglement properties of many-body systems in their combinatorial structure, in which tensors are connected along a network pattern and identify (the coefficients, in a given basis, of the wave function corresponding to) quantum states of the given system. Born as convenient mathematical tools for numerical evaluations of many-body wavefunctions, which become translatable into graphical manipulations, tensor network techniques have found an amazing number of applications: from the classification of exotic phases of quantum matter (e.g. topological order) \cite{15,16} to new formulation of
the non-perturbative renormalization of interacting quantum field theories \[17\]-\[19\], down to realizations of the AdS/CFT correspondence \[20\]-\[23\].

Despite their disparate origin, it should be clear already from our sketchy description that the type of mathematical structures identified by quantum gravity approaches and used in the theory of tensor networks are very similar. And consequently, it is very natural to try to put the two frameworks in more direct contact. This is the main goal of the present article. Indeed, the structural similarity had been noted before \[24\]-\[27\], and also exploited in the context of renormalization of spin foam models treated as lattice gauge theories \[28\]-\[31\]. The last set of works, in particular, has already shown how fruitful tensor network techniques can be for quantum gravity models.

Before we start presenting our results, we want to offer some motivations for our work, both from the quantum gravity perspective and from the tensor network side.

From the quantum gravity point of view, the general motivation is clear. Tensor networks provide a host of tools and results that could find useful application in quantum gravity; in particular they may become central tools in the renormalization analysis of GFT models \[32\]-\[37\], in addition to their mentioned role in the renormalization analysis of spin foams models \[38\]-\[40\]. And such renormalization analyses are, in turn, the main avenue for solving the crucial problem of the continuum limit in such formalism.

More specifically, tensor networks are very effective in taking into account and controlling the entanglement properties of quantum states in many-body systems. This is exactly the language in which GFT deals with quantum gravity states; moreover, in GFT, the very connectivity of spin network states, encoded in the links of the underlying graphs, is associated with entanglement between the fundamental quanta constituting them (associated to nodes) \[41\]. One example of this type of application, as we show in this paper, is the computation of entanglement entropy in spin network states and relate LQG with holography, which was also the subject of a number of other works in the LQG/GFT literature \[27\]-\[32\]-\[55\].

Further, the identification of the true (interacting) vacuum state of a quantum gravity theory, in absence of any space-time background or preferred notions of energy, is a difficult matter even at the purely conceptual level, leaving aside the formidable technical challenges. One possible criterion, suited to this context, is to look for states which maximize entanglement, by some measure (e.g. entanglement entropy). In this respect, to reformulate the kinematics and dynamics of GFT and LQG states in terms of tensor networks, and to do
the same for their renormalization, seems a promising strategy.

Finally, recent results in the application of tensor networks to AdS/CFT \[21\] suggest that this application would be fruitful even within the conventional perspective of canonical quantum gravity (including LQG). From this perspective, in fact, the task of quantum gravity is the construction of the space of quantum states of the gravitational field which satisfy the (quantum counterpart of the) Hamiltonian constraint encoding the dynamics of quantised GR. A number of results in AdS/CFT suggest that a static AdS space-time, which we expect to be one such state, at the quantum level, satisfies the Ryu-Takayanaki (RT) formula \[20\] for the entanglement entropy, which is very efficiently computed (as we also show in this paper) via random tensor network techniques \[23\]. One is led to conjecture that this may be a general properties of physically interesting quantum states of the gravitational field, and so far no counterexample to this conjecture has been found. This prompts the search, by the same techniques, for similar states in canonical quantum gravity.

From the perspective of the theory of tensor networks, one general good point of dwelling into the correspondence with quantum gravity states should also be obvious. This identifies a new domain of applications, of truly fundamental nature, for techniques and ideas which have already proven powerful in others. Indeed, we expect that a number of key results obtained via tensor network techniques, most notably holographic mappings and indications of new topological phases in many-body systems, can be reproduced in this new context, with deep implications. In perspective, it is here that one will be able to test the suggestion that quantum information has a truly foundational role to play in our understanding of physical reality.

More practically, a number of techniques have been developed, and many results obtained, concerning the dynamics of GFT and spin-network states, also thanks to the many related developments in the theory of random tensors, and our dictionary proves that the GFT formalism provides a natural definition of the dynamics of random tensor networks. Specifically, it means that the many results in GFT can help dealing with general (non-Gaussian) probability distributions over random tensor networks, as well as offering new takes on more standard problems, like entropy calculations, in tensor network theory. In fact, we offer some examples of these applications in the following.

In this paper, we do not target the more ambitious objective of a calculation of the RT formula for the entanglement entropy in the full quantum gravity formalism of group
field theory. Having established the general dictionary between group field theory states and (generalized) random tensor networks, we content ourselves with reproducing the RT formula, along the lines of the derivation given in [23] in two new cases: for group field theory states corresponding to generalized tensor networks, but only using a group field theory dynamics in the simplest approximation and dealing only with averages over the tensor functions associated to the network nodes, rather than treating the full tensor network as a group field theory observable; for the simple truncation of group field theory states corresponding to spin networks with fixed spin labels. We leave a more complete and comprehensive analysis for forthcoming work.

The paper is organized as follows. In the next section, we summarize the basic elements of spin network states and of their embedding in the GFT formalism, as well as the definition of tensor networks. Having done so, we define the precise correspondence between GFT states and tensor networks, showing how the first generalizes and provides a Fock space setting for the second. In the following section, we derive the $N$th Rényi entropy using GFT techniques, in the group representation and for a generalized tensor network, but without taking advantage of the full GFT formalism; next, we compute the same Rényi entropy and derive the RT formula from a purely spin-network perspective, seen as a truncation of more general GFT states. This is meant to be a clear example of how the same problem can be fruitfully approached from both sides of the correspondence. Finally, in the last section, we discuss one key universality result from the theory of random tensors, which extends to GFTs, and which could have direct impact on the applications of random tensor networks. We end up with a summary of our results.

II. GROUP FIELD THEORY AND TENSOR NETWORKS

A $d$-dimensional GFT is a combinatorially non-local field theory living on $(d$ copies of) a group manifold [6–9]. Due to the defining combinatorial structure, the Feynman diagrams $\mathcal{F}$ of the theory are dual to cellular complexes, and the perturbative expansion of the quantum dynamics defines a sum over random lattices of (a priori) arbitrary topology. A similar lattice interpretation can be given to the quantum states of the theory. For GFT models where appropriate group theoretic data are used and specific properties are imposed on the
states and quantum amplitudes, the same lattice structures can be understood in terms of simplicial geometries. The associated many-body description of such lattice states can be given in terms of a tensor network decomposition. The corresponding (generalized) tensor networks are thus provided with a field theoretic formulation and a quantum dynamics (and, in specific models, with additional symmetries). In this section, after a brief introduction to the GFT formalism, we detail this correspondence between GFT states and (generalized) tensor networks.

A. Group Field Theory

Let $G$ denote an arbitrary semi-simple Lie group; in the following, we assume for simplicity that $G$ is compact, but the framework can easily be generalized to the non-compact case. A group field $\varphi$ is a complex function defined on a number of copies of the group manifold $G$:

$$\varphi : G^d \rightarrow \mathbb{C}$$

$$g_i \mapsto \varphi(g_i)$$

where we use the shorthand notation $g_i$ for the set of $d$ group elements $\{g_1, g_2, \cdots, g_d\}$.

The GFT field can be also seen as an infinite-dimensional tensor, transforming under the action of some (unitary) group $U^{\times d}$, as:

$$\varphi(g_1, \ldots, g_d) \rightarrow \int [dg_i] U(g_i', g_i) \cdots U(g_d', g_d) \varphi(g_1, \ldots, g_d),$$

and

$$\varphi^*(g_1, \ldots, g_d) \rightarrow \int [dg_i] U^\ast(g_i', g_i) \cdots U^\ast(g_d', g_d) \varphi^*(g_1, \ldots, g_d)$$

$$\text{for } \int dg_i U(g_i', g_i)^\ast U(g_i, \tilde{g}_i) = \delta(g_i', \tilde{g}_i).$$

This requires the $d$ arguments of the GFT field to be labeled and ordered. We will see in the following how one can decompose the same field into finite-dimensional tensors; in this finite-dimensional case, the correspondence with tensor network formalism will be evident, and it will also be evident then in which sense GFTs provide a generalization of it.

The GFT dynamics is defined by an action, at the classical level, and a partition function at the quantum level. The combinatorial structure of the pairing of field arguments in the GFT interactions is part of the definition of a GFT model. An interesting class of models
6[12] is defined by the requirement that the interaction monomials are \textit{tensor invariants}, i.e. that GFT fields are convoluted in such a way as to produce an invariant under the above mentioned (unitary) transformations[4].

Another class of GFT models is instead based on the requirement that the Feynman diagrams of the theory are simplicial complexes, which in turn requires the interaction kernels to have the combinatorial structure of d-simplices. This class of models is also the one on which model building for 4d quantum gravity has focused on, producing models whose Feynman amplitudes have the form of simplicial gravity path integrals and spin foam models [6–9], and, more generally, lattice gauge theories. This involves an additional symmetry requirements on the GFT fields and interactions, which will play a crucial role in the following.

In this simplicial case, the GFT action has the general form

$$S_d[\varphi] = \frac{1}{2} \int d g_i d g_i' \varphi(g_i) K(g_i g_i'^{-1}) \varphi(g_i') +$$

$$+ \frac{\lambda}{d+1} \int \prod_{i \neq j = 1}^{d+1} d g_{ij} V(g_{ij} g_{ji}'^{-1}) \varphi(g_{ij}) \cdots \varphi(g_{d+1j}) ,$$

where $dg_i$ is an invariant measure on $G$ and we use the notation $\varphi(g_{ij}) = \varphi(g_{12}, \cdots, g_{1d+1})$. $K$ is the kinetic kernel, $V$ the interaction kernel, $\lambda$ a coupling constant for the $d+1$-degree homogeneous interaction. The two kernels satisfy the invariance properties

$$K(h g_i g_i'^{-1} h') = K(g_i g_i') ,$$

$$V(h_i g_{ij} g_{ji}'^{-1} h_j^{-1}) = V(g_{ij} g_{ji}'^{-1}) \quad \forall h, h', h_i \in G .$$

This implies that the action is invariant under the gauge transformations $\delta \varphi(g_i) = \tilde{\varphi}(g_i)$, where $\tilde{\varphi}$ is any function satisfying

$$\int_G d h \tilde{\varphi}(h g_1, \cdots, h g_d) = 0 .$$

This symmetry is gauge fixed if one restricts the field $\varphi$ to satisfy

$$\varphi(h g_i) = \varphi(g_i) .$$

1 Such invariants are in one to one correspondence with \textit{colored d-graphs} $B$ constructed as follows: for each GFT field (resp. its complex conjugate) draw a white (resp. black) node with $d$ outgoing links each labeled by $d$ different colors, then connect all links in such a way that a white (resp. black) node is always connected to a black (resp. white) node and that only links with the same color can be connected.
The action is also invariant under the global symmetry

\[ \varphi(g_1, \cdots, g_d) \rightarrow \varphi(g_1 h, \cdots, g_d h). \]

GFT’s Feynman diagrams define cellular complexes \( \mathcal{F} \) weighted by amplitudes assigned to the faces, edges and vertices of the dual two-skeleton otubularf a chosen triangulation of a d dimensional topological spacetime \( \mathcal{M}_\mathcal{F} \). As mentioned, their Feynman diagram evaluations reproduce the associated amplitudes of a spin foam model, or, in different variables, of a simplicial gravity path integral \([56–58]\), providing a generalisation of the lattice formulation of gravity à la Regge, with an accompanying sum over lattices, generalising matrix models for 2d gravity to any dimension \([6–12]\).

Let us give some more detail on the construction, to clarify the above points. A specific theory, with a specific related Feynman cellular complex, is completely defined by the choice of the kernels. Let’s consider the simplest case, consisting in the choice

\[ K(g_i, g_i') = \int_G d h \prod_i \delta(g_i g_i'^{-1} h), \]

\[ V(g_{ij} g_{ji}^{-1}) = \int_G \prod_i d h_i \prod_{i < j} \delta(h_i g_{ij} g_{ji}^{-1}, h_j^{-1}) \]

where \( \delta(\cdot) \) is the delta function on \( G \) and the integrals ensure the gauge invariance defined in (5), and let us restrict to the case of dimension \( d = 3 \). To keep track of the combinatorics of field arguments in the kernels, it is useful to represent the Feynman diagram as a stranded graph. The field \( \varphi \) has three arguments, so each edge of a Feynman diagram comprises three strands running parallel to it. Four edges meet at each vertex and the form of the interaction \( V \) in (9) forces the strands to recombine as in Figure 1.

The three strands running along the edges can be understood to be dual to a triangle and the propagator \( K \) gives a prescription for the gluing of two triangles. At the vertex, four triangles meet and their gluing via \( V \) form a tetrahedron. With this interpretation the Feynman diagram of a GFT is clearly dual to a triangulated 3d simplicial complex (which will be generically a singular pseudo-manifold) and this is true in any dimension \([59–61]\).

The quantum states of the theory can be given a similar combinatorial characterization in terms of graphs and dual cellular complexes, as it should be already intuitive in the above example, in which GFT fields themselves are associated to triangles. We will not detail this aspect of the formalism.
FIG. 1. Correspondence between Feynman diagram and triangulation: Each strand of the graph forms a closed loop which can be interpreted as the boundary of a 2d disk. These data are enough to reconstruct a topological 2d complex $F$, the vertices and edges of this complex correspond to vertices and edges or the Feynman graph, the boundary of the faces of $F$ correspond to the strands of the Feynman graph.

**B. Fourier modes of the group field as tensor fields**

As a function on a group $G$, the field $\varphi$ can be decomposed in terms of unitary irreducible representations $(\rho, V_\rho)$ of $G$ using the Peter-Weyl theorem, $L^2(G) \simeq \bigoplus \rho V_\rho \otimes V_\rho^*$, giving

$$\varphi(g) = \sum_\rho d_\rho \text{Tr} [\hat{\varphi}_\rho \rho^{ab}(g)] \quad (10)$$

Here, $d_\rho \in \mathbb{N}$ is the dimension of the representation $\rho : G \to \text{Aut}(V_\rho)$, the indices $a, b = 1, \ldots, d_\rho$ are matrix indices associated to the matrix $\rho(g)$ representing the group element $g$, and $\hat{\varphi}_\rho \in V_\rho \otimes V_\rho^* \simeq \text{End}(V_\rho)$ is the matrix Fourier coefficient of the function $\varphi$. In other words, each $\hat{\varphi}_\rho$ is a rank $d_\rho = N$ matrix.

Let us consider, as a specific example, the same decomposition for the case of $d = 3$, with $G = SU(2)$. The unitary irreps of $SU(2)$, $\mathbb{V}^j$, are labeled by the spin $j \in \mathbb{N}/2$. Using the right invariance property of the field, one obtains the following decomposition

$$\varphi(g_1, g_2, g_3) = \sum_{\{j\}} \text{Tr} \left[ \varphi_{m_1, m_2, m_3}^{(j)} \left( \prod_i \sqrt{d_j} D_{m_1, n_1}^{j_i} (g_i) \right) \tilde{t}_{m_1, m_2, m_3}^{(j)} \right] \quad (11)$$

where $d_j$ is the dimension, $D^j(g) \in \text{End}(\mathbb{V}^j)$ the group matrix element and $\tilde{t}_{m_1, n_2, n_3}^{(j)} \in \text{Hom}_G(\mathbb{V}^{j_1} \otimes \mathbb{V}^{j_2} \otimes \mathbb{V}^{j_3}, \mathbb{C})$ is the three-valent intertwiner operator (related to the Clebsch-Gordan map $\Psi_{j_1 j_2}^{j_3} : \mathbb{V}^{j_1} \otimes \mathbb{V}^{j_2} \to \mathbb{V}^{j_3}$). We used the shorthand notation $\{j\}$ for the set of spin labels $(j_1, j_2, j_3)$. 
The fields $\varphi^{\{j\}}_{m_1,m_2,m_3}$ result from the contraction of the Fourier transformed GFT fields $\hat{\varphi}_{\{j\}}$ with the intertwiner tensor imposing the gauge symmetry at the vertex. \(^2\)

$$\varphi^{\{j\}}_{\{m\}} = \sum_{\{k\}} \hat{\varphi}_{\{m\};\{k\}} \iota^{\{k\}}_{\{j\}} \prod_i \sqrt{d_{ji}}. \quad (12)$$

The Fourier transformed fields depend on the (discrete) representation space labels of the Lie group in question. Thus, generically Fourier transformed GFT fields are tensors of some rank $d$, $\varphi_{\{m\}_j}$ with discrete indices $\vec{m}_j = \{m_1, \ldots, m_d\}$. \(^3\)

In (11), such tensors are contracted with the spin network basis tensors

$$S^{\{j\}}_{\{m\}} = \left( \prod_i \sqrt{d_{ji}} D_{ji}^{\vec{m}_i \vec{n}_i} (g_i) \right) \tilde{\iota}^{\{j\}}_{\{n\}}, \quad (13)$$

encoding the properties of the vertex of the spin network graph dual to the (d-1)-dimensional triangulation that can be associated to the GFT states.

### C. Group Field Single Particle States

Functions $\varphi(g_i)$ can also be understood as single particle wave functions for quanta corresponding to single open vertices of a spin network graph (in fact, they also label coherent states of the GFT field operator, which define the simplest condensate states of the theory [55, 62, 63]).

Let us define these ‘single-particle’ quantum states as

$$|\varphi\rangle = \int_{G^d} dg_i \varphi(g_i) |g_i\rangle \quad (14)$$

where $dg_i \equiv dg_1 dg_2 \ldots dg_d$ is the Haar measure on the group manifold $G^d$, invariant under the gauge transformation, and the vectors $|g_1\rangle \ldots |g_d\rangle$ provide a basis on the respective infinite dimensional spaces $\mathbb{H} \simeq L^2[G]$.

The single particle state $|\varphi\rangle$ is then defined in $\mathbb{H}^{\otimes d}$. Moreover we require $|\varphi\rangle$ to be normalized (this is of course not the case for the classical GFT fields or the GFT condensate wavefunctions):

$$\langle \varphi | \varphi \rangle = \int dg_i \overline{\varphi(g_i)} \varphi(g_i) = 1. \quad (15)$$

\(^2\) This is the standard factorization of a symmetric tensor into a degeneracy tensor with all the degrees of freedom and a structural tensors (the Clebsch-Gordan coefficients) completely determined by the symmetry group G (Wigner-Eckart theorem) [23].

\(^3\) To regularize some quantities, especially at the dynamical level, it may be necessary to impose a (large) cut-off $N$ in the range of the representation indices.
Considering the case of $G = SU(2)$, we can decompose the basis $|g\rangle$ into the unitary irreducible representation of $SU(2)$ as

$$|g\rangle \equiv \sum_{j,m,n} \sqrt{d_j} D_{mn}^j(g) |j, n, m^\dagger\rangle$$  \hspace{1cm} (16)

and vice versa

$$|j, n, m^\dagger\rangle = \int_{SU(2)} dg \sqrt{d_j} D_{mn}^j(g) |g\rangle.$$  \hspace{1cm} (17)

In particular, the tensor decomposition given in (11) holds at the quantum level, hence defining the quantum fields $\varphi^{(j)}_{m_1, m_2, m_3}$ as actual tensors states.

Tensors in (13) define the $SU(2)$-invariant single vertex spin network wave functions (in group representation)

$$\psi_\chi(g_i) = \langle \chi | g_i \rangle = \left( \prod_i \sqrt{d_j} D_{m_i, n_i}^j(g_i) \right) \bar{\gamma}_{\{j\}}^{\{i\}},$$  \hspace{1cm} (18)

The basis vector $|\chi\rangle = |j, m, i\rangle$ denotes the standard $SU(2)$ spin network basis (labelled by spins and angular momentum projections associated to their $d$ open edges, and intertwiner quantum numbers).

### D. Many-Body Description and Tensor Network States

We now describe the quantum states of the formalism, emphasizing their many-body structure, following [64].

Consider a $d$-valent graph formed by $V$ disconnected components, each corresponding to a single gauge invariant $d$-valent vertex and $d$ 1-valent vertices, thus having $d$ edges.\(^4\) We refer to this type of disconnected components as open spin network vertices.

To such a graph we can associate a generic wavefunction given by a function of $d \times V$ group elements,

$$\Phi(g_i) = \Phi(g_1, ..., g_d, g_1, ..., g_d, ..., g_v, ..., g_v)$$  \hspace{1cm} (19)

\(^4\) One could work instead with the larger Hilbert spaces of non-gauge invariant states $L^2[G^{d \times V}]$ without imposing any gauge symmetry at the vertices of spin network graphs, and consider this condition as part of the dynamics. The above construction would proceed identically, with the same final result, but with the basis of single-vertex states now given by the above functions without the contraction of representation function with a $G$-intertwiner.
FIG. 2. A tensor network Γ is a set of tensors whose indices are contracted according to a network pattern. A network pattern can be always represented as a graph, given by a set of nodes (n) and links (ℓ) connecting nodes. A link is called an internal link when it connects two different nodes; while it is called a boundary link when it connects only one node. The number of links that connect to a node is called the valence of the node.

defined on the group space $G^{d \times V} / G^V$ ($V$ copies of $G^d$, quotiented by the isotropy group of the single particle function $\varphi_v(g_i)$ at the each vertex); here the index $a$ runs over the set of vertices, while the index $i$ still runs over the links attached to each vertex).

These functions are exactly like many-particles wave functions for point particles living on the group manifold $G^d$, and having as classical phase space $(T^*G)^d$ (which is also the classical phase space of a single open spin network vertex or polyhedron).

Accordingly, a state $|\Phi\rangle \in \mathbb{H}_V \simeq L^2[G^{d \times V} / G^V]$ can be conveniently decomposed into products of single-particle (single-vertex) states,

$$\Phi(g_i^a) = \langle g_i^a | \Phi \rangle = \sum_{\chi_1, i = 1 \ldots V} \varphi_{\chi_1 \ldots \chi_V} \psi_{\chi_1}(g_i) \cdots \psi_{\chi_V}(g_i)$$  \hspace{1cm} (20)

While the above decomposition is completely general, a special class of states can be constructed in direct association with a graph or network Γ. The association works as follows. Start from the d-valent graph with $V$ disconnected components (open spin network vertices) to which a generic $V$-body state of the theory is associated. A partially connected d-valent graph can be constructed by choosing at least one edge $i$ in a vertex $a$ and gluing it to one edge $j$ of the vertex $b$, i.e. joining the two edges along their 1-valent vertices. The final graph will be fully connected if all edges have been glued. Each pair of glued edges $\{a_i, b_j\}$ will identify a link $L$ of the resulting (partially) connected graph. In the spin representation, i.e. in terms of the basis of functions $\psi_{\chi_1}(g_i) \cdots \psi_{\chi_V}(g_i)$, the gluing is implemented by the identification of the spin labels $j_i^a$ and $j_j^b$ associated to the two edges being glued and by the contraction of the corresponding vector indices $m_i^a$ and $m_j^b$. In other
words, the corresponding wave functions for closed graphs can be decomposed in a basis of closed spin network wave functions, obtained from the general product basis by means of the same contractions:

$$\Phi_\Gamma(g^\alpha) = \langle g^\alpha | \Phi_\Gamma \rangle = \sum_{\chi_a, a=1,\ldots,V} \Phi^{j_1\ldots j_V}_\Gamma \left( \prod_{L \in \Gamma} \delta_{j_a, j_b} \delta_{m^a_l, m^b_l} \right) \psi_{\chi_1}(g_l) \ldots \psi_{\chi_V}(g_l)$$

(21)

where the coefficients of the wave function can in turn be understood as the resulting of considering generic coefficients \( \phi^{\chi_1\ldots\chi_V} \) and contracting them with some choice of functions \( M_{n^a_l n^b_l}^{j_a j_b} \)

$$\Phi^{j_1\ldots j_V}_\Gamma = \phi^{\chi_1\ldots\chi_V} \left( \prod_{L \in \Gamma} \delta_{j_a, j_b} M_{n^a_l n^b_l}^{j_a j_b} \right),$$

(22)

where the contraction is left implicit.

For fixed \( \{j\} \), each resulting contraction scheme of tensors (each identified by a set of labels \( \chi \)) defines a tensor network state.

In the group representation, the gluing amounts to considering wave functions with a specific symmetry under simultaneous group translation of the arguments associated to the edges being glued:

$$\Phi_\Gamma(g_a) = \Phi_\Gamma(g^1, \ldots, g^d h^d_{1,\ldots,V}, g_2, \ldots, g_{d'}, \ldots, g_V h^d_{1,\ldots,V}, \ldots, g_V).$$

(23)

In the end, given a tensor network with graph \( \Gamma \), the \( \Phi^{j_1\ldots j_V}_\Gamma \) defined above will contain all the information about the combinatorics of the quantum geometry state.

A further special case corresponds to those states for which the coefficients \( \phi^{\chi_1\ldots\chi_V} \) themselves have a product form, i.e. can be decomposed in terms of tensors. In this case, as it is for the spin network wave functions, the coefficients \( \Phi^{j_1\ldots j_V}_\Gamma \) can be obtained as a tensor trace

$$\Phi^{j_1\ldots j_V}_\Gamma = \text{Tr}(\bigotimes_L M \bigotimes_v \varphi^{\{j\}(v)} \{m\})$$

(24)

again, in the case of fully connected graphs \( \Gamma \) (otherwise, some angular momentum labels will remain on the left had side, corresponding to the edges that have not been glued). In lattice theory, we would say that the network \( \Gamma \) (fixed \( \{j\} \)) provides a tensor network decomposition of the tensor state \( \Phi^{j_1\ldots j_V}_\Gamma \).

The equivalence of a special class of GFT states with the lattice tensor network states, and the sense in which GFT states generalise them, can be further elucidated by the following example.
E. Link state as a gluing operation

A tensor \( \hat{T} \) is a multidimensional array of complex numbers \( \hat{T}_{\lambda_1,\ldots,\lambda_d} \in \mathbb{C} \). The rank of tensor \( \hat{T} \) is the number \( d \) of indices. The size of an index \( \lambda \), denoted \( d_{|\lambda|} \), is the number of values that the index \( \lambda \in \mathbb{N} \) takes [65].

Analogously, at the quantum level, to each leg of the tensor one associates a Hermitian inner product space \( \mathcal{H}_D \), with dimension \( D \) given by the size of the indices \( \lambda \in \{1, 2, \ldots, d_{|\lambda|} = D\} \). Given an orthonormal basis \( |\lambda\rangle \), in \( \mathcal{H}_D \), a covariant tensor of rank \( d \) is a multilinear form on the Hilbert space of the vertex \( T: \mathcal{H}_D^\otimes d \to \mathbb{C} \). Hence a tensor state is written as

\[
|T\rangle = \sum_{\lambda_1,\ldots,\lambda_d} \hat{T}_{\lambda_1,\ldots,\lambda_d} |\lambda_1\rangle \otimes \cdots \otimes |\lambda_d\rangle
\]  

(25)

where \( \hat{T}_{\lambda_1,\ldots,\lambda_d} \equiv T(\lambda_1, \ldots, \lambda_d) \) denote the components in the canonical dual tensor product basis.

A tensor network is generally given by a set of \( d \)-valent vertices \( v \), corresponding to rank \( d \) tensors. In particular, a state corresponding to a set of unconnected vertices is written as a tensor product of individual vertex states

\[
|T_N\rangle \equiv \bigotimes_n |T_n\rangle
\]  

(26)

Individual vertex states are glued by links. To each end of a link we associate a Hilbert space \( \mathcal{H}_D \). The Hilbert space of the link \( \ell \) is then \( \mathcal{H}_\ell = \mathcal{H}_D^\otimes 2 \) and a link state can be written as

\[
|M\rangle = M_{\lambda_1\lambda_2} |\lambda_1\rangle \otimes |\lambda_2\rangle
\]  

(27)

where we choose to take the link states \( |M\rangle \) to be generically entangled[7]. In general, the entanglement of the links will encode the information on the connectivity of the graph. Two nodes are connected if their corresponding states contract with a link state,

\[
\hat{T}_{12} \equiv \langle M| |T_1\rangle |T_2\rangle = T_{\lambda_1,\ldots,\lambda_v}^{(1)} M_{\lambda_v\lambda_0} T_{\lambda_1,\ldots,\lambda_u}^{(2)} \bigotimes_{i \neq a}^v |\lambda_i\rangle \otimes \bigotimes_{j \neq b}^u |\lambda'_j\rangle
\]  

(29)

5 One can observe it by defining a density matrix \( \rho_M \equiv |M\rangle \langle M| \) and tracing out one of the Hilbert space, without losing generality, tracing out \( \mathcal{H}_D \) of \( |\lambda_2\rangle \), then computing the von Neumann entropy of the reduced density matrix \( \rho_1 \equiv \text{Tr}_2 \rho_M = M^\dagger M \). The entropy \( S = \text{Tr} \rho_1 \ln \rho_1 \) is non-zero unless \( M_{\lambda_1\lambda_2} \) can split as \( M_{\lambda_1\lambda_2} = A_{\lambda_1} B_{\lambda_2} \). For simplicity, in the next sections we will often assume that the link state is maximally entangled, i.e.

\[
|M\rangle = \frac{1}{\sqrt{D}} \delta_{\lambda_1\lambda_2} |\lambda_1\rangle \otimes |\lambda_2\rangle.
\]  

(28)
Notice that if $|M\rangle$ was a non-entangled state, the connection would be trivial, i.e. the two nodes would be practically disconnected and the corresponding state could be written as a tensor product of two states,

$$\hat{T}_{12} = T_{\lambda_1\ldots\lambda_n}^{(1)}A_{\lambda_1} \otimes \ldots \otimes A_{\lambda_n} \otimes B_{\lambda_1\lambda_2\ldots\lambda_n}^{(2)}|\lambda_1\rangle \otimes \ldots \otimes |\lambda_n\rangle = |T_1T_2\rangle$$

Then given a network $\mathcal{N}$ with $N$ nodes and $L$ links, the corresponding state is

$$|\Psi_{\mathcal{N}}\rangle \equiv \bigotimes_{\ell}^{L} \langle M_\ell | \bigotimes_{n}^{N} |T_n\rangle$$

Because all links are contracted with nodes, $|\Psi_{\mathcal{N}}\rangle$ is then in the Hilbert space associated to the boundary links of the network, which is denoted as $\mathbb{H}_{\partial\mathcal{N}}$. $|\Psi_{\mathcal{N}}\rangle$ is a state in $\mathbb{H}_{\partial\mathcal{N}}$.

The above structure can be identified also for the special GFT states mentioned at the end of the previous subsection, which are formed by generalised $L^2(G^d)$ functions associated to the nodes of the network. In this case, the analogous of the generic link state in (27), which is also the group counterpart of the gluing operators associated in the spin representation to the matrices $M$, can be defined in as the convolution functional

$$\langle M_{g_\ell} | \equiv \int dg_1dg_2 \ M(g_1^\dagger g_\ell g_2) \ (g_1| \otimes |g_2) \in \mathbb{H}^{\oplus 2},$$

where the functions $M(g)$ are assumed to be invariant under conjugation $M(g) = M(hgh^{-1})$. When a link $\ell$ connects two nodes, say $a$ and $b$, the corresponding state $\langle M_{g_\ell} |$ contracts with states $|\varphi_a\rangle$ and $|\varphi_b\rangle$

$$\langle M_{g_\ell} | \varphi_a \rangle |\varphi_b\rangle = \int dg_1dg_2dg_a^bg_b^b M(g_1^\dagger g_\ell g_2) \ \varphi_a(g_1, g_a^a) \varphi_b(g_2, g_b^b) |g_1^a\rangle |g_2^b\rangle ,$$

where we have singled out, among the arguments of the vertex wave functions $\varphi$ the ones affected by the gluing operation. In these terms, the open $d$-valent tensor network graph $\Gamma$ with $V$ vertices, can be written as

$$|\Phi_{\Gamma}^{g_\ell}\rangle \equiv \bigotimes_{\ell \in \Gamma} \langle M_{g_\ell} | \bigotimes_{n}^{V} |\varphi_n\rangle = \int dg_\theta \ \Phi_{\Gamma}(g_\ell, g_\theta) |g_\theta\rangle$$

where the $\{g_\theta\}$ denote the group elements on the open links.

The role of the link state in tensor network, thus, is naturally generalised by the convolution function, defined for the group field variables. This is due to the fact that the group
fields $\varphi(g_i)$ on $G^d$ can be interpreted as rank $d$ tensors, with indices spanning the group space $G$, and associated Hilbert space (for each index) being $L^2(G)$. The multiparticle state given in (23) can then be interpreted as a tensor state with indices $g_0$ and rank given by the number of open links of the spin network graph.

F. Link function in spin decomposition

As showed in II D, many-body state can also be decomposed into spin representations. Suppose $M(g_1^\dagger g_2 g_\ell g_2)$ can be written as

$$M(g_1^\dagger g_\ell g_2) = \sum_{jmn} d_j M_{mn}^j D_{mn}^j(g_1^\dagger g_\ell g_2)$$

Then, as a simple example, the state $\langle M | \varphi_a \rangle \langle \varphi_b \rangle$ can be written in terms of $\varphi_{\ell n}^j$, $i_m$ and $M_{mn}^j$ as

$$\langle M | \varphi_a \rangle \langle \varphi_b \rangle = \int dg_1 dg_2 dg_\ell d_{g_1^\dagger} M(g_1^\dagger g_\ell g_2) \varphi_a(g_1, g_\ell^a) \varphi_b(g_2, g_\ell^b) | g_{1i} \rangle | g_{2i} \rangle$$

Graphically, the last line can be presented as

$$\langle M | \varphi_a \rangle \langle \varphi_b \rangle = \sum_{jmnkl} \sum_{i_a i_b} M_{mn}^l D_{kl}^l(g_\ell) | j_{ai} \rangle | j_{bi} \rangle$$

The case of ordinary, finite-dimensional tensors is obtained if we pass from a Lie group to a discrete group.

Let us consider, as a basic example, the case of a field defined on the discrete $n$th cyclic group $Z_n$. Given the nonempty set $X = \{ \vec{\lambda} | \vec{\lambda} = (\lambda_1, \ldots, \lambda_d), \lambda_k \in Z_n \}$. The function $\varphi : X \rightarrow \mathbb{R}$ (or $\mathbb{C}$) is a real or complex valued function on $X$ and we indicate by

$$\varphi_{\vec{\lambda}} \equiv \varphi(\vec{\lambda}).$$

the value of $\varphi$ on the set of $d$ elements $\vec{\lambda}$. The function $\varphi_{\vec{\lambda}}$ can be interpreted as a tensor with $d$ discrete indices $\varphi_{\lambda_1, \ldots, \lambda_d}$, where $\lambda \in \{1, 2, \ldots, \text{dim}(Z_n)\}$. 

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6 The case of ordinary, finite-dimensional tensors is obtained if we pass from a Lie group to a discrete group.

7 Notice that we are introducing the bold font for vectorial quantities, in order to shorten the notation in spin representation.
From the graphic equation, one can immediately observe that the upper part is an open tensor network $|\Phi^j\rangle$, given by the tensor trace of a collection of tensors

\begin{equation}
\phi_{m}^{ji} \equiv \sum_{n} \bar{t}_{nm} \phi_{nm}^{j} \tag{40}
\end{equation}

for each node and matrices $M_{nm}^{j}$ for each link.

G. Dictionary

We summarize the established dictionary between group field theory states and generalized random tensor networks in terms of two synthetic tables. The correspondence between group field theory and tensor network description is summarized in Table A:
| Table A | Group Fields | Tensors |
|---------|--------------|---------|
| group basis | $|g_i\rangle \in \mathbb{H} \simeq L^2|G|$ | $|\lambda_i\rangle$, $\lambda_i = 1, \ldots, D$ in $\mathbb{H}_D$ index basis |
| one particle state | $|\varphi\rangle = \int_{G^d} dg_i \varphi(g_i) |g_i\rangle$ | $|T_n\rangle = \sum_{\{\lambda_i\}} T_{\{\lambda_i\}} |\lambda_i\rangle \in \mathbb{H}_n = \mathbb{H}_D^{\otimes d}$ tensor state |
| gluing functional | $\langle M_{g_1}| = \int d g_1 d g_2 \ M(g_1^* g_2) \langle g_1| \langle g_2|$ | $|M\rangle = M_{\lambda_1\lambda_2} |\lambda_1\rangle \otimes |\lambda_2\rangle \in \mathbb{H}_\ell = \mathbb{H}_D^{\otimes 2}$ link state |
| multiparticle state | $|\Phi_T\rangle \in \mathbb{H}_V \simeq L^2[G^{d \times V}/G^V]$ | $|\Psi_N\rangle$ tensor network state |
| product state convolution | $|\Phi^{\otimes}_T\rangle \equiv \bigotimes_{\ell \in T} \langle M_{g_\ell}| \bigotimes_n |\varphi_n\rangle$ | $|\Psi_N\rangle \equiv \bigotimes_\ell^L \langle M_{\ell}| \bigotimes_n^N |T_n\rangle \in \mathbb{H}_{\partial N}$ tensor network decomposition |
| randomness | $\frac{1}{Z} d\nu(\varphi)$ | $T^U_\mu \equiv (UT^0)_\mu$ random tensor state |

The generalisation of tensor networks in terms of GFT states is evident in the spin-j decomposition of the latter $\varphi(g_i) = \sum_j \text{Tr}[\varphi^j_{\{m\}} \left( \prod_i \sqrt{d_{i}} D_{m_i, n_i}^j(g_i) \right) \tilde{\varphi}^j_{\{n\}}]$. Once we turn off the sum over all possible $j$s, fix the representation labels and ask them to be equal, generically Fourier transformed GFT fields $\varphi^j_{\{m\}}$, are tensors of single rank $d$, with discrete indices $m_i = \{m_1, \ldots, m_d\}$ spanning a finite dimensional space. The equivalence is summarized in table B:
In the following sections, with the longer-term goal of a full understanding and computation of the Ryu-Takayanagi (RT) formula \cite{20} in the field-theoretic GFT context, we are going to use the inputs provided by the established dictionary to investigate the holographic RT formula for the case of networks of combinatorial tensor group fields described by means of the GFT formalism and spin network techniques, along the lines proposed for the case of Random Tensor Networks by \cite{23}.

In the tensor network generalisation of the gauge gravity duality \cite{66}, the RT formula strongly supports a general relation between entanglement and geometry, in turn leading to the suggestion that the whole of spacetime geometry can be understood as emergent from (quantum information-theoretic) properties of non-spaciotemporal quantum building blocks. Of course, this last suggestion has a life on its own and it has been brought forward in many different contexts \cite{63, 67, 74}. In this sense, our work provides further steps towards the calculation of the RT formula within a complete quantum gravity setting, a concrete and

| Table B | GFT network | Spin Tensor Network | Tensor Network |
|---------|-------------|---------------------|---------------|
| node    | $\varphi(\vec{g})$ | $\varphi_j^{(m)}$ | $T_{(\mu)}$ |
|         | $= \varphi(g_1, g_2, g_3, g_4)$ | $\propto \sum_{\{k\}} \varphi_j^{(m)}(g_1, g_2, g_3, g_4)$ | |
| link    | $M(g_1^\dagger g_2 g_3 g_4)$ | $M_{\lambda_1 \lambda_2}$ | |
| sym     | $\varphi(h \vec{g}) = \varphi(\vec{g})$ | $\prod_{s} D_m^i D_{m'}^i (g)_{m'_m} = i_{m_1 \ldots m_v}$ | $\prod_{s} U_{\mu_1 \mu_2} T_{\mu_1 \ldots \mu_v}$ |
| state   | $|\Phi_g^j\rangle \equiv \otimes_{\ell} \langle M_{g_\ell} | \otimes_n |\psi_n\rangle$ | $|\Psi^j\rangle \equiv \otimes_{\ell} \langle M^j_\ell | \otimes_n |\phi_{n_\ell}^{j_n}\rangle$ | $|\Psi^N\rangle \equiv \otimes_{\ell} \langle M^N_\ell | \otimes_n |T_n\rangle$ |
| indices | $g_i \in G$, $|g_i\rangle \in \mathbb{H} \simeq L^2[G]$ | $m_i \in \mathbb{H}_j$, SU(2) spin-$j$ irrep. | $\mu_i \in \mathbb{Z}_n$, $n$th cyclic group |
| dim     | $\infty$ | dim $\mathbb{H}_j = 2j + 1$ | dim $\mathbb{Z}_n = n$ |
general indication of the holographic character of gravity, which goes beyond the AdS/CFT
gauge gravity duality framework.

III. RYU-TAKAYANAKI FORMULA FOR A GFT TENSOR NETWORKS

The starting point of our analysis is the state $|\Psi_\Gamma\rangle$, corresponding to an open network
graph where each node is dressed with a group field generalised tensor. Because of the
field theoretic description, we can see the network as a random tensor network and use
the established correspondence to apply standard path integral formalism to evaluate the
expectation values of entropies and other tensor observables. In particular, then, our goal
consists in investigate the holographic entanglement properties of the GFT network by means
of techniques recently applied to the study of the holographic behaviour for Random Tensor
Networks [23], building on the dictionary we have established between the two languages.
This calculation is not in the full GFT setup, i.e. the state $|\Psi_\Gamma\rangle$ is not treated, in the
calculation of the averaging over random (generalised) tensors, as an $n$-point function of
a given GFT. This more complete calculation is postponed to a future analysis. Still, we
apply several techniques from GFT and generalized the calculations in [23] based on our
dictionary:

1. Tensors are generalized to group fields, from a finite dimensional object to a square
   integrable $L^2$ function, mapping from group manifolds to the complex numbers $\mathbb{C}$.

2. A gauge symmetry of the group field associated to each vertex as a vertex wave function
   is introduced in order to fit our setup more to the context of the quantum gravity
   theory.

3. The average over the $N$-replica of the wave functions (generalised tensors) associated
to each network vertex is reinterpreted as a $N$-point correlation function of a (simple)
   GFT model, which turns the averaged Rényi entropy into an amplitude in GFT.

The last point can be seen as an approximation of a more complete calculation in which
the (average over the) whole tensor network is understood as a GFT N-point function, and
computed as such. This more complete calculation based on the full GFT setup is being
explored [76]. We believe that the leading term of the entropy, at least for the entanglement
entropy, would not be changed.
FIG. 3. Boundary \( \partial \mathcal{N} \) of network \( \mathcal{N} \) divided into two parts \( A \) and \( B \).

Given our tensor network state, \( |\Psi_N\rangle \equiv \bigotimes_{E}^L \langle M_E| \bigotimes_{n}^N |T_n\rangle \in \mathcal{H}_{\partial \mathcal{N}} \), we start by considering a bipartition of the boundary Hilbert space,

\[
\mathcal{H}_{\partial \mathcal{N}} = \mathcal{H}_A \otimes \mathcal{H}_B
\]

associated to the definition of two – a priori non adjacent – subregions \( A \) and \( B \) of the boundary (see Figure 3).

A measure of the entanglement between the two subsystems is given by the von Neumann entropy of the reduced density matrix of the subsystem, either \( A \) or \( B \), defined by partial tracing over the full system Hilbert space. Focussing on subsystem \( A \), for \( \rho \equiv |\Psi_N\rangle \langle \Psi_N| \), we have

\[
\rho_A = \text{Tr}_B(\rho),
\]

and the entanglement entropy between \( A \) and \( B \) is given by the von Neumann entropy

\[
S_{\text{EE}}(A) = -\text{Tr}[\hat{\rho}_A \ln \hat{\rho}_A],
\]

where now

\[
\hat{\rho}_A \equiv \frac{\rho_A}{\text{Tr}\rho}
\]

is the normalized reduced density matrix.

In order to calculate \( S_{\text{EE}}(A) \), due to the technical difficulty in computing the von Neumann entropy, we need to make use of the standard replica trick. Contracting \( N \) copies of the reduced density matrix \( \rho_A \) and taking the logarithm of the trace of \( \rho_A^N \), one obtains the \( N \)th-order Rényi entropy

\[
S_N(A) = -\frac{1}{N-1} \ln \text{Tr} \hat{\rho}_A^N.
\]

22
The above formula is easier to compute and coincides with the von Neumann entropy of region $A$ in the limit $N \to 1$

$$S_{\text{EE}}(A) = \lim_{N \to 1} S_N(A)$$  \hspace{1cm} (46)

A. $N$th Rényi entropy for a GFT random tensor network

We focus now on the case of the $N$th Rényi entropy for a bipartite GFT state $|\Psi_\Gamma\rangle$ with support on a generic open graph $\Gamma$. We divide the boundary $\partial \Gamma$ of the graph $\Gamma(V_\Gamma, L_\Gamma, L_{\partial \Gamma})$, with $V_\Gamma$ nodes, $L_\Gamma$ internal links and $L_{\partial \Gamma}$ boundary links, into two parts, called $A$ and $B$. The $N$th Rényi entropy between $A$ and $B$ is given by

$$e^{(1-N)S_N} = \frac{Z_N}{Z_0^N}$$  \hspace{1cm} (47)

with $Z_0^N \equiv (\text{Tr}\rho)^N$, $Z_N \equiv \text{Tr}\rho_A^N = \text{Tr}[\rho^\otimes N P(\pi_A^0;N,d)]$ and the network density matrix $\rho$ defined as

$$\rho = |\Psi_\Gamma\rangle \langle \Psi_\Gamma| = \text{Tr}_\ell \left[ \bigotimes_n |M_\ell\rangle \langle M_\ell| \bigotimes_n |\psi_n\rangle \langle \psi_n| \right] \equiv \text{Tr}_\ell \left[ \bigotimes_n \rho_\ell \bigotimes_n \rho_n \right].$$  \hspace{1cm} (48)

Here, for convenience, we use the equivalence of the trace of the reduced density with the result of the trace over the action of the permutation operator $P(\pi_A^0;N,d)$ on the full $\rho^N$ for

$$P(\pi_A^0;N,d) = \prod_{s=1}^N \delta_{\mu^{(s+1)d)}(\mu^{(s)d})}$$  \hspace{1cm} (49)

with $d$ is the dimension of the Hilbert space in the same region $A$.

Given the random nature of the tensor network, we look for the typical value of the entropy. Analogously to the case considered in \cite{23}, the variables $Z_N$ and $Z_0$ are easier to average than the entropy, since they are quadratic functions of the network density matrix $\rho$. In particular, the entropy average can be expanded in powers of the fluctuations $\delta Z_N = Z_N - \mathbb{E}(Z_N)$ and $\delta Z_0^N = Z_0^N - \mathbb{E}(Z_0^N)$, so that

$$\mathbb{E}(S_N(A)) = -\mathbb{E} \left( \log \frac{\mathbb{E}(Z_N)}{\mathbb{E}(Z_0^N)} + \delta Z_N \right)$$

$$= -\log \frac{\mathbb{E}(Z_N)}{\mathbb{E}(Z_0^N)} + \text{fluctuations}$$  \hspace{1cm} (50)

---

8 For $N = 2$, e.g., the cyclic group $S_2$ only has two elements: the identity $\mathbb{1}$ and swap operator $F$, so that $P(\pi_{A_1}^0,2,d) \equiv F(A)$. Then, $Z_2 = \text{Tr}[\rho^\otimes 2 F(A)] = \rho_{A_1 A_2 B_1 B_2} F(A_{A_1 A_2 B_1 B_2})$ and

$Z_0^2 = \text{Tr}[\rho^\otimes 2] = \rho_{A_1 A_2 B_1 B_2} F(A_{A_1 A_2 B_1 B_2})$. 

23
As showed in [23], for large enough bond dimensions $D$, as a direct consequence of the concentration of measure phenomenon [75], the statistical fluctuations around the average value are exponentially suppressed. Therefore, it is possible to approximate the entropy with high probability by the averages of $Z_N$ and $Z_0^N$,

$$e^{(1-N)E(S_N)} \sim \frac{E(\text{Tr} \rho_A^N)}{E(\text{Tr} \rho)^N} = \frac{E(\text{Tr}[\rho^N \mathcal{P}(\pi^0_A; N, d)])}{E(\text{Tr} \rho)^N} = \frac{\text{Tr} \left[ \bigotimes \rho^N \otimes \mathcal{E}(\rho^N) \mathcal{P}(\pi^0_A; N, d) \right]}{\text{Tr} \left[ \bigotimes \rho^N \otimes \mathcal{E}(\rho^N) \right]}.$$

(51)

In order to get the typical Rényi entropy one needs then to compute $E(Z_N)$ and $E(Z_0^N)$ separately. The average over the tensor fields can be carried out before taking the partial trace, since the latter is a linear operation. Therefore, the key step consists in computing the quantity

$$E(\rho_n^N) = E[(|\psi_n\rangle \langle \psi_n|^N) = E \left[ \left( \int \prod_a d_{\psi_n} d_{\bar{\psi}_n} \psi_n(g_a) \bar{\psi}_n(g_a) |g_a\rangle \langle g_a| \right)^N \right],$$

(52)

hence, eventually, the expectation value of $N$ copies of the network wavefunction,

$$E \left[ \prod_a \psi_n(g_a) \bar{\psi}_n(g_a) \right].$$

(53)

where $d_{\psi} \equiv \prod_i d_{g_i}$, $\psi(g) \equiv \psi(g_1, \cdots, g_4)$ and $g$ is independent from $\bar{g}$, which denotes the arguments of $\bar{\psi}$.

Now, we define the averaging operation $E[\cdots]$ via the path integral of a generic group field theory model

$$E[f(\psi, \bar{\psi})] \equiv \int [D\psi][D\bar{\psi}] f(\psi, \bar{\psi}) e^{-S[\psi, \bar{\psi}]}$$

(54)

where $S[\psi, \bar{\psi}]$ is the action of the given model of interest,

$$S[\psi, \bar{\psi}] = \int d\psi d\bar{\psi} \bar{\psi}(g) \mathcal{K}(g, g) \psi(g) + \lambda S_{\text{int}}[\psi, \bar{\psi}] + cc,$$

(55)

the first term on the right hand side defining the kinetic term of the model. In the following calculation, we consider the particular case where

$$\mathcal{K}(g, g) = \delta(g^\dagger g)$$

(56)

which thus implies a free part of the action of the simple form [9]

$$S_0[\psi, \bar{\psi}] = \int d\psi \bar{\psi}(g) \psi(g)$$

(57)

9 Notice that several GFT models of quantum gravity [6][4] can be put in this form.
We further assume that the coupling constant $\lambda$ is much smaller than 1, so the path integral $\mathbb{E}[f[\psi, \bar{\psi}]]$ can be perturbatively expanded in powers of $\lambda$

$$\mathbb{E}[f[\psi, \bar{\psi}]] = \int [\mathcal{D}\psi][\mathcal{D}\bar{\psi}] f[\psi, \bar{\psi}] e^{-S_0[\psi, \bar{\psi}]} \left(1 + \lambda S_{\text{int}}[\psi, \bar{\psi}] + \mathcal{O}(\lambda^2)\right)$$

$$\equiv \mathbb{E}_0[f[\psi, \bar{\psi}]] + \mathcal{O}(\lambda) \quad .$$

This is the regime of validity of the so-called spin foam expansion, seen from within the GFT formalism [1–9]. In the following calculation, we will only focus on the leading term $\mathbb{E}_0[f[\psi, \bar{\psi}]]^{10}$.

Because of the gauge symmetry $\psi(hg) = \psi(g)$, the gauge equivalent paths in the above path integral have to be removed (via gauge fixing). In order to do so, we first introduce the following notation: if $g = (g_1, g_2, g_3, g_4)$, then

$$[g] \equiv g^{-1} = (1, g_1^{-1}g_2, g_1^{-1}g_3, g_1^{-1}g_4) \quad .$$

Then, we insert the delta functional $\delta[\psi(g) - \psi([g])]$ constraint into the path integral, so that the average becomes

$$\mathbb{E}_0[f[\psi, \bar{\psi}]] \equiv \int [\mathcal{D}\psi][\mathcal{D}\bar{\psi}] f[\psi, \bar{\psi}] \delta[\psi(g) - \psi([g])] e^{-f \int dg \bar{\psi}(g)\psi(g)} \quad .$$

Since this equation is simply the expectation value of $f[\psi, \bar{\psi}]$ in the free group field theory, we can immediately give the expectation value of (53) via Wick theorem:

$$\mathbb{E}_0\left[\prod_{a}^{N} \psi(g_a)\bar{\psi}(g_0)\right] = C \sum_{\pi \in S_N} \prod_{a}^{N} \delta\left([g_a][g_\pi(a)]^\dagger\right) \quad ,$$

$$= C \sum_{\pi \in S_N} \int \prod_{a}^{N} dh_a \prod_{a}^{N} \delta\left(h_ag_a[g_\pi(a)]^\dagger\right) \quad ,$$

where $g$ is independent from $g$, $\delta([g][g]^\dagger) = \prod_{s=2}^{4} \delta\left(g_1^\dagger g_s g_s^\dagger g_1\right)$ and $\delta(hgg') = \prod_{s=1}^{4} \delta\left(hg_s g_s^\dagger\right)$.

In the second equality, we re-introduce the gauge symmetry by inserting integrals of $h_a \in SU(2), N = 1, 2, \cdots N$ into the delta functions such that $g_{sa}$ on each leg of the node are on an equal footing, unlike $g_1 = \mathbb{1}$ in the gauge fixing procedure. So in the following calculation, the network is without gauge fixing, i.e. all integrals of $g$ have to be performed.

$^{10}$ This, in turn, means that, from the point of view of the quantum gravity model, the quantum gravity dynamics is imposed only to the extent in which it is captured by the kinetic term in the GFT action.
Denote now $\prod_a^{N} \delta \left( h_a g_{a} \mathcal{g}^{(a)} \right)$ as

$$\mathbb{P}_h(\pi) \equiv \prod_a^{N} \delta \left( h_a g_{a} \mathcal{g}^{(a)} \right) = \prod_{a}^{N} \prod_{s=1}^{4} \delta \left( h_s g_{sa} g_{s\pi(a)} \right) \equiv \prod_{s}^{4} \mathbb{P}_s^{\pi}(\pi),$$

(62)

where $h$ denotes the set of $h_a$, $a = 1, \ldots, N$. When $h_a = \mathbb{1}$ for all $a$ from 1 to $N$,

$$\mathbb{P}_1(\pi) = \prod_a^{N} \delta \left( g_a \mathcal{g}^{(a)} \right) = \mathbb{P}(\pi; N, D^4) = \prod_{s}^{4} \mathbb{P}_s^{\pi}(\pi; N, D^4)$$

(63)

where $\mathbb{P}(\pi; N, D^4)$ and $\mathbb{P}_s^{\pi}(\pi; N, D^4)$ are the representations of $\pi \in \mathcal{S}_N$ on $\mathbb{H}^\otimes 4$ and $\mathbb{H}$, respectively.

Then, $Z_N$ and $Z_0^N$ become

$$Z_N \approx C_{\mathcal{V}} \sum_{\pi_n \in \mathcal{S}_N} \int \prod_n \text{d}h_n \text{Tr} \left[ \bigotimes_\ell \rho_N^{\ell} \bigotimes_n \mathbb{P}_{h_n(\pi_n)} \mathbb{P}(\pi_0; N, d) \right]$$

$$\equiv C_{\mathcal{V}} \sum_{\pi_n \in \mathcal{S}_N} \int \prod_n \text{d}h_n \mathcal{N}_{\mathcal{A}}(h_n, \pi_n)$$

(64)

$$Z_0^N = C_{\mathcal{V}} \sum_{\pi_n \in \mathcal{S}_N} \int \prod_n \text{d}h_n \text{Tr} \left[ \bigotimes_\ell \rho_N^{\ell} \bigotimes_n \mathbb{P}_{h_n(\pi_n)} \right]$$

$$\equiv C_{\mathcal{V}} \sum_{\pi_n \in \mathcal{S}_N} \int \prod_n \text{d}h_n \mathcal{N}_{0}(h_n, \pi_n),$$

(65)

which means that $Z_N$ and $Z_0^N$ correspond to summations of the networks $\mathcal{N}_{\mathcal{A}}(h_n, \pi_n)$ and $\mathcal{N}_{0}(h_n, \pi_n)$ where at each node $n$ we have a contribution $\mathbb{P}_{h_n(\pi_n)}$ and at each link $\ell$ we have a contribution $\rho_N^{\ell}$. The only difference between these two networks is the boundary condition: where $Z_N$ is defined with $\mathbb{P}(\pi_\mathcal{A}; N, d)$ on $\mathcal{A}$ of $\partial \Gamma$ and $\mathbb{P}(\mathcal{1}; N, d)$ on $\overline{\mathcal{A}}$ of $\partial \Gamma$, and $Z_0^N$ is defined with $\mathbb{P}(\mathcal{1}; N, d)$ for all boundary region $\partial \Gamma$.

Since at each node $\mathbb{P}_{h_n(\pi_n)}$ is decoupled among the incident legs, because of (62), the value of the networks $\mathcal{N}_{\mathcal{A}}(h_n, \pi_n)$ and $\mathcal{N}_{0}(h_n, \pi_n)$ can be written as products factorised over links:

$$\mathcal{N}_{\mathcal{A}}(h_n, \pi_n) = \prod_{\ell \in \mathcal{A}} \mathcal{L}_{\ell}(\pi_n, \pi_{n'}; h_n, h_{n'}) \prod_{\ell \in \mathcal{A}} \mathcal{L}_{\ell}(\pi_n, \pi_\mathcal{A}; h_n) \prod_{\ell \in \mathcal{A}} \mathcal{L}_{\ell}(\pi_n, \mathcal{1}; h_n)$$

(66)

$$\mathcal{N}_{0}(h_n, \pi_n) = \prod_{\ell \in \Gamma} \mathcal{L}_{\ell}(\pi_n, \pi_{n'}; h_n, h_{n'}) \prod_{\ell \in \partial \Gamma} \mathcal{L}_{\ell}(\pi_n, \mathcal{1}; h_n)$$

(67)

Because the $\mathcal{L}_{\ell}$ on the boundary are special cases of the $\mathcal{L}_{\ell}$ in the graph $\Gamma$, it is enough to calculate the $\mathcal{L}_{\ell}$ on the internal links. In general, $\mathcal{L}(\pi, \pi', h, h')$ can be written as a trace of a modified representation of a permutation group element $\varpi \equiv (\pi')^{-1} \pi$ as

$$\mathcal{L}(\pi, \pi'; h, h') = \text{Tr} \left[ \mathbb{P}_h(\pi) \rho_N^{\ell} \mathbb{P}_{h'}(\pi) \right] = \text{Tr} \left[ \mathbb{P}_H \left( (\pi')^{-1} \pi \right) \right] \equiv \text{Tr} \left[ \mathbb{P}_H (\varpi) \right],$$

(68)
where

\[ H = \left\{ H_a \mid H_a \equiv (h_{\omega(a)})^\dagger h_a, \ \forall a = 1, \ldots, N \right\} \]  

(69)

\[ \text{FIG. 4. } \mathcal{L}(\pi, \pi', h, h') \]

When \( \pi = \pi' \), we have \( \omega = 1 \) and \( H = (h')^\dagger h \), and then

\[ \mathcal{L}(\pi, \pi'; h, h') = \text{Tr} \left[ P_h(\pi) \rho_{\pi} P_{h'}(\pi) \right] = \text{Tr} \left[ P_{(h')^\dagger h} (1) \right] \]

(70)

\[ = \prod_a \int d g_a d g'_a d g_{\pi(a)} d g'_{\pi(a)} \delta \left( h_a g_a g_{\pi(a)}^\dagger \right) \delta \left( g_{\pi(a)} g'_{\pi(a)}^\dagger \right) \times \]

\[ \times \delta \left( h'_a g'_a g'_{\pi(a)}^\dagger \right) \delta \left( g'_a g'_a^\dagger \right) \]

\[ = \prod_a \delta \left( (h'_a)^\dagger h_a \right) = \prod_a \delta \left( H_a \right) \]

(71)

The above equation can be depicted graphically as in Fig.5

\[ \text{FIG. 5. } \mathcal{L}(\pi, \pi, h, h') \]

When \( \pi \neq \pi' \), we have

\[ \mathcal{L}(\pi, \pi'; h, h') = \text{Tr} \left[ P_H (\omega) \right] \]

(72)

In order to perform the computation, it is necessary to use some facts about the permutation group \( S_N \), which we recall briefly, before proceeding.

- Any element \( \omega \in S_N \) can be expressed as the product of disjoint cycles \( \mathcal{C}_i \)

\[ \omega \equiv \prod_i \chi(\omega) \mathcal{C}_i \]  

(73)
where \( 1 \leq \chi(\varpi) \leq N \) is the number of cycles in \( \varpi \), which is 1 when \( \varpi \) is a 1-cycle and is \( N \) only when \( \varpi = 1 \). For instance, the permutation \( \varpi = \{3241\} \) can be expressed as a product of two cycles \((134)(2)\), in which \( \varpi(1) = 3, \varpi(3) = 4, \varpi(4) = 1 \) and \( \varpi(2) = 2 \). \((132)\) is a 3-cycle, because there are three elements in the cycle. We denote the number of elements in the cycle \( \mathcal{C}_i \) as \( r_i \), which is also called the length of the cycle. We also have \( \sum_i r_i = N \). Although the cycles \( \mathcal{C}_i \) commute with each other, we order the cycles such that

\[
1 \leq \cdots \leq r_i \leq r_{i+1} \leq \cdots \leq N .
\]

We denote \( a^i_k \), where \( k \) is from 1 to \( r_i \), the elements of \( \mathcal{C}_i \), and then we furthermore assume that \( \varpi(a^i_k) = a^i_{k+1} \).

Thus, the cycle can be written as

\[
\mathcal{C}_i = (a^i_1 a^i_2 \cdots a^i_{r_i}) .
\]

• The trace of \( \mathbb{P}_H(\varpi) \) can be expressed as the product of the traces of the individual cycles \( \mathcal{C}_i \)

\[
\text{Tr}[\mathbb{P}_H(\varpi)] = \prod_i \text{Tr}[\mathbb{P}_H(\mathcal{C}_i)] .
\]

Using the definition of \( \mathbb{P}_H \), one can immediately obtain the trace of the cycle \( \mathcal{C}_i \) as

\[
\text{Tr}[\mathbb{P}_H(\mathcal{C}_i)] = \int \prod_{k=1}^{r_i} dg_{a_k^i} \delta \left( H_{a_k^i} g_{a_k^i} g_{a_k^i}^\dagger \right) = \delta \left( \prod_k H_{a_k^i} \right) ,
\]

where

\[
\prod_{k=1}^{r_i} H_{a_k^i} \equiv H_{a_{r_i}} \cdots H_{a_2} H_{a_1} .
\]

Then the trace of \( \mathbb{P}_H(\varpi) \) is

\[
\mathcal{L}(\pi, \pi'; h, h') = \text{Tr}[\mathbb{P}_H(\varpi)] = \prod_i \delta \left( \prod_k H_{a_k^i} \right) .
\]

• On the boundary of \( \mathcal{N}_0 \) and \( B \) of \( \mathcal{N}_A \), \( \mathcal{L}(\pi, \mathbb{1}; h, h') \) is a very special case of \( \mathcal{L}(\pi, \pi'; h, h') \) where \( \pi' = \mathbb{1} \) and \( h' = \mathbb{1} \)

\[
\mathcal{L}(\pi, \mathbb{1}; h) \equiv \mathcal{L}(\pi, \mathbb{1}; h, \mathbb{1}) = \text{Tr}[\mathbb{P}_h(\pi)] = \prod_i \delta \left( \prod_k h_{a_k^i} \right) .
\]
On the boundary $A$ of $\mathcal{N}_A$, $\mathcal{L}(\pi, \pi^0_A; \mathbf{h})$ corresponds also to a special case of $\mathcal{L}(\pi, \pi'; \mathbf{h}, \mathbf{h'})$, where $\mathbf{h}' = 1$ and $\pi' = \pi^0_A = C_0$, which is the $N$-cycle that for any integer $k$ from 1 to $N$, $C_0(k) = [k]_N + 1$

$$\mathcal{L}(\pi, \pi^0_A; \mathbf{h}) \equiv \mathcal{L}(\pi, C_0; \mathbf{h}, 1) = \text{Tr} \left[ P_h \left( C_0^{-1} \pi \right) \right] = \prod_i \delta \left( \prod_{k=1}^{r_i} h_{a_k} \right).$$  \hspace{1cm} (82)

Altogether, for a given network $\mathcal{N}(\mathbf{h}_n, \pi_n)$, defining the new variables $\varpi \equiv (\pi')^{-1} \pi$ and $\mathbf{H}$ given by (69) for each link, the corresponding link value is a product of $\chi(\varpi)$ delta function

$$\mathcal{L}(\pi, \pi'; \mathbf{h}, \mathbf{h'}) \equiv \mathcal{L}(\varpi; \mathbf{H}) = \text{Tr} \left[ P_{\mathbf{H}} (\varpi) \right] = \prod_i \delta \left( \prod_{k=1}^{r_i} H_{a_k} \right) \hspace{1cm} (83)$$

In particular, when $\pi = \pi'$, the link value $\mathcal{L}(\pi, \pi; \mathbf{h}, \mathbf{h'})$ is given by a product of $N$ delta functions as shown in (70) and we re-present it here

$$\mathcal{L}(\pi, \pi; \mathbf{h}, \mathbf{h'}) = \prod_a N \delta \left( (h_a')^i h_a \right) = \prod_a \delta \left( H_a \right) \hspace{1cm} (84)$$

which is non-zero only when $\mathbf{h} = \mathbf{h'}$.

So in the end the network is divided into several regions, in each of which $\pi_n$ and $\mathbf{h}_n$ are the same. The links which connect different regions identify boundaries between each pair of different regions, called again domain walls. Corresponding to different domain walls and different assignments of permutation groups to each region, we have different patterns for the given network. We introduce pattern functions $\mathcal{P}_A(\pi_n)$ and $\mathcal{P}_0(\pi_n)$ such that

$$\mathcal{P}_A(\pi_n) \equiv \int \prod_n \text{d} \mathbf{h}_n \mathcal{N}_A(\mathbf{h}_n, \pi_n)$$

$$\mathcal{P}_0(\pi_n) \equiv \int \prod_n \text{d} \mathbf{h}_n \mathcal{N}_0(\mathbf{h}_n, \pi_n).$$  \hspace{1cm} (85)

Given a set of $\{\pi_n\}$, $\mathcal{P}_A(\pi_n)$ and $\mathcal{P}_0(\pi_n)$ correspond to a certain network pattern with fixed boundary conditions, illustrated in the following figure.

More explicitly,

$$\mathcal{P}_A(\pi_n) = \int \prod_n \text{d} \mathbf{h}_n \prod_{\ell \in \Gamma} \left[ \chi(\varpi) \delta \left( \prod_{k=1}^{r_{\ell_i}} H_{a_{\ell_k}} \right) \right] \prod_{\ell \in A} \left[ \chi(C_0^{-1} \pi_n) \delta \left( \prod_{k=1}^{r_{\ell_i}} h_{a_{\ell_k}} \right) \right] \prod_{\ell \in A} \left[ \chi(\pi_n) \delta \left( \prod_{k=1}^{r_{\ell_i}} h_{a_{\ell_k}} \right) \right].$$  \hspace{1cm} (87)
\[ \mathcal{P}_0(\pi_n) = \int \prod_n \text{d}h_n \prod_{\ell \in \Gamma} \left[ \chi(\omega_{\ell}) \prod_i \delta \left( \prod_{k=1}^{r_\ell} h_{\ell a_i^k} \right) \right] \prod_{\ell \in \partial \Gamma} \left[ \chi(\pi_{n\ell}) \prod_i \delta \left( \prod_{k=1}^{r_\ell} h_{\ell a_i^k} \right) \right] \] (88)

They are exactly the amplitudes of a topological BF field theory, with given boundary condition, discretized on a specific 2-complex among the \( N \) replica of networks, with each different pattern \( \mathcal{P} \) corresponding to a different 2-complex. Each edge of the 2-complex is associated with a holonomy \( h_{na} \) that is on node \( n \) and the \( a \)th replica. The two ends of the holonomy are the vertices of the 2-complex. The \( h_{na} \) inside a delta function form a loop holonomy, the corresponding edges of which form the face of the 2-complex. Then \( Z_N \) and \( Z^N_0 \) are sum of BF amplitudes with different 2-complexes.

\[ Z_N \equiv \mathcal{C}^{\mathcal{Vr}} \sum_{\pi_n \in \mathcal{S}_N} \mathcal{P}_A(\pi_n), \quad Z^N_0 \equiv \mathcal{C}^{\mathcal{Vr}} \sum_{\pi_n \in \mathcal{S}_N} \mathcal{P}_0(\pi_n) \] (89)

It is important to notice that this simple form of the various functions entering the calculation of the entropy, with the emergence of BF-like amplitudes, is not generic. It follows from the choice of GFT kinetic term, from the approximation used in the calculation of expectation values (neglecting GFT interactions) and from the special type of tensor network, in GFT language, that we have chosen (with simple delta functions associated to the links of the network). More involved, and interesting, cases could be considered.

What we are interested in is the leading term of \( Z_N \) and \( Z^N_0 \), while the dimension \( D \) of Hilbert space \( \mathcal{H} \) is much larger than 1. This leads us to seek the most divergent term of \( \mathcal{P}_A(\pi_n) \) and \( \mathcal{P}_0(\pi_n) \). In other words, we need to know the degree of divergence of \( \mathcal{P}_A(\pi_n) \) and \( \mathcal{P}_0(\pi_n) \). The divergence degree of BF amplitudes discretized on a lattice has been the subject of a number of works, both in the spin foam an GFT literature (see for example
Let us first focus on a sub-region $R$ of the network such that $\pi_n = \pi$ for all nodes $n$ inside of $R$. Suppose that there are $L_i$ links inside $R$ and $L_e$ links connecting with other regions. Since we only consider 4-valent nodes, the number of nodes inside $R$ is

$$V \equiv \frac{1}{4} (2L_i + L_e) = \frac{L_i}{2} + \frac{L_e}{4}$$

A minimum spanning tree (MST) $T$, which contains $\#_T = V - 1$ links, can be found in $R$.

According to (84), since $\pi_n = \pi$, there are $N$ delta functions on each link. The integrals over $h_n$ would eliminate the $(V - 1)N$ deltas associated to the MST and leave only one set of $N$ integrals over $h = \{h_a\}$ and $(L_i/2 - L_e/4 + 1)N \delta(1)$’s. Here we keep indicating the divergent factor as the delta function evaluation originating it, but of course it should be understood more properly as a function of the cut-off used to regularize it. The pattern function of region $R$ is then

$$P_R(\pi) \equiv \int \prod_{n \in R} dh_n \prod_{\ell \in MST}^{L_i} \prod_a^{N} \delta(H_{\ell a}) \prod_{\ell \in MST}^{L_e} \prod_a^{N} \delta(H_{\ell a}) \prod_{\ell \notin MST}^{L_e} \prod_a^{N} \delta(H_{\ell a})$$

$$= \int \prod_{n \in R} dh_n \prod_{\ell \in MST}^{L_i} \prod_a^{N} \delta(H_{\ell a}) \prod_{\ell \notin MST}^{L_e} \prod_a^{N} \delta(H_{\ell a}) \prod_{\ell \in MST}^{L_e} \prod_a^{N} \delta(H_{\ell a})$$

$$= \int dh \prod_{\ell} \prod_i \delta \left( \prod_{k=1}^{L_e} H_{\ell a_k} \right) \left[ \frac{L_i}{2} - \frac{L_e}{4} + 1 \right] N \int dh \prod_{\ell} \prod_i \delta \left( \prod_{k=1}^{L_e} H_{\ell a_k} \right) \left[ \frac{L_i}{2} - \frac{L_e}{4} + 1 \right] N$$

In the calculation, we have used

$$\int \prod_{n \in R} dh_n \prod_{\ell \in MST}^{L_i} \prod_a^{N} \delta(H_{\ell a}) = \int dh \quad (93)$$

and ($h_n = h$)

$$\prod_{\ell \notin MST}^{L_e} \prod_a^{N} \delta(H_{\ell a}) = \left[ \delta(1) \right]^{\left( \frac{L_i}{2} - \frac{L_e}{4} + 1 \right)} N \quad (94)$$

The above calculation shows that we can coarse-grain the region $R$ into one single $L_e$-valent node which is colored by $\pi$ and $h$.

$$\int \prod_{n \in R} dh_n = \int \prod_{n \in R} dh_a = \int dh \quad [\delta(1)]^{\left( \frac{L_i}{2} - \frac{L_e}{4} + 1 \right)} N$$

(95)
So the degree of divergence in region $R$ is: the number of internal links $#_i = L_i$ subtracted the number of links in the MST $#_T = V - 1$, and then times the number of replica $N$,

$$
#_R = (#_i - #_T)N = (L_i - V + 1)N = \left(\frac{L_i}{2} - \frac{L_e}{4} + 1\right)N.
$$

(96)

Since the boundary condition of $N_0$ is $\pi = 1$ and $h = 1$, the boundary of $N_0$ can be coarse-grained into a single node with $\pi = 1$ and $h = 1$. The same consideration holds for $N_A$: its boundary can be coarse-grained into two nodes, one of which corresponds to $A$ with $\pi = C_0$, $h = 1$ and the other to $B$ with $\pi = 1$ and $h = 1$. The corresponding closed graphs are denoted as $\Gamma_0$ and $\Gamma_{AB}$. A certain pattern $\mathcal{P}(\pi_n)$ divides $\Gamma_0$ and $\Gamma_{AB}$ into $M$ regions that can be coarse-grained into $M$ nodes, each of which is colored with permutation group $\pi_m$ and $N$ integrals over $h_m$. Denote the graph with pattern $\mathcal{P}(\pi_n)$ as $\Gamma_0(\pi_m)$ and $\Gamma_{AB}(\pi_m)$, and denote the corresponding coarse-grained graphs as $\Gamma^c_0(\pi_m)$ and $\Gamma^c_{AB}(\pi_m)$.

One can show that, for $\Gamma_0$, the pattern in which all nodes have assigned the same permutation group $\pi = 1$ has the highest degree of divergence $#_0$.

$$
#_0 = (#_{\ell \in \Gamma_0} - #_{\mathcal{T}_0})N
$$

(97)

where $#_{\ell \in \Gamma_0}$ is the number of links in graph $\Gamma_0$. Let us consider a coarse-grained graph $\Gamma^c_0(\pi_m)$. Denote the number of links in region $m$, between regions $m$ and $m'$, and between region $m$ and boundary $\partial \Gamma$ are $L_m$, $L_{mm'}$, and $L_{m0}$, respectively. The proof goes as follows:

1. The permutation group on links between coarse-grained nodes $m$ and $m'$ is $\varpi_{mm'} \equiv \pi_m^{-1} \pi_{m'}$. As given by (83), the number of the delta functions on one of the links is the number of the disjoint cycles in $\varpi$, which is $\chi(\varpi_{mm'}) < N$. Since all links between $m$ and $m'$ are identical, having the same link value, which is given by (83), when one integrate over $h_m$ and $h_{m'}$, only $\chi(\varpi_{mm'})$ deltas will be eliminated and left with $\delta(1)$
to the order of $\chi(\varpi_{mm'})(L_{mm'} - 1)$ and $2N - \chi(\varpi_{mm'})$ $h$ integrals. In fact

$$
\int \mathrm{d}h \mathrm{d}h' (\mathrm{Tr} [P_H (\varpi)])^L = \int \mathrm{d}h \mathrm{d}h' \prod_i \chi(\varpi) \left[ \delta \left( \prod_{k=1}^{r_i} H_{a_k}^{r_i} \right) \right]^{L-1}
$$

$$
= \int \mathrm{d}h \prod_i \chi(\varpi) \left[ \delta \left( \prod_{k=1}^{r_i} H_{a_k} \prod_{k=1}^{r_i} H_{a_k}^{r_i} \right) \right]^{L-1}
$$

$$
= [\delta (1)] \chi(\varpi)^{(L-1)} \int \mathrm{d}h
$$

(98)

2. MST can be chosen for $\Gamma_0(\pi_m)$, $\Gamma_0^c(\pi_m)$ and $M$ regions. It is obvious that, given a MST $T_m$ for each of the $M$ regions and a MST $T_{\Gamma_0^c(\pi_m)}$ for $\Gamma_0^c(\pi_m)$, rooting from the coarse-grained boundary node $\partial \Gamma$, a MST $T_{\Gamma_0(\pi_m)}$ of $\Gamma_0(\pi_m)$ can be constructed.

$$
T_{\Gamma_0(\pi_m)} = \bigcup_m T_m \cup T_{\Gamma_0^c(\pi_m)}
$$

(99)

The number of branches of the trees is

$$
\# T_{\Gamma_0(\pi_m)} = \sum_m \# T_m + \# T_{\Gamma_0^c(\pi_m)}
$$

(100)

3. The degree of divergence of region $m$ is given by (96)

$$
\#_m = (L_m - \# T_m)N
$$

(101)

Similarly, for the divergence degree $\#_{\Gamma_0^c(1)}$ of the pattern where all coarse-grained nodes have the same permutation $\pi_m = 1$ is

$$
\#_{\Gamma_0^c(1)} = \left( \sum_{0 \leq m < m' \leq M} L_{mm'} - \# T_{\Gamma_0^c(\pi_m)} \right)N
$$

(102)

The degree of divergence of $\Gamma_0^c(\pi_m)$ is smaller than $\#_{\Gamma_0^c(1)}$

$$
\#_{\Gamma_0^c(\pi_m)} < \#_{\Gamma_0^c(1)}
$$

(103)

This is because, after evaluating the delta functions on the MST $\Gamma_0^c(\pi_m)$ in accordance with (98), there are still $MN - \sum_{(mm') \in T_{\Gamma_0^c(\pi_m)}} \chi(\varpi_{mm'})$ integrals over $h$. Performing
these integrals makes the degree of divergence of $\Gamma_0^c(\pi_m)$ not bigger than the following quantity

$$\#\Gamma_0^c(\pi_m) \leq \sum_{0 \leq m < m' \leq M} L_{mm'} \chi(\varpi_{mm'}) - \sum_{(mm') \in T_{\Gamma_0^c}(\pi_m)} \chi(\varpi_{mm'})$$  \hfill (104)

$$= \sum_{(mm') \notin T_{\Gamma_0^c}(\pi_m)} L_{mm'} \chi(\varpi_{mm'})$$  \hfill (105)

$$+ \sum_{(mm') \in T_{\Gamma_0^c}(\pi_m)} (L_{mm'} - 1) \chi(\varpi_{mm'})$$  \hfill (106)

which is definitely smaller than $\#\Gamma_0(1)$ because $\chi(\varpi_{mm'}) < N$.

4. So the divergence degree of $\Gamma_0(\pi_m)$ is smaller than the divergence degree $\#_0$ for the pattern where all nodes have the same permutation.

$$\#\Gamma_0(\pi_m) = \#\Gamma_0^c(\pi_m) + \sum_m M \#_m < \#\Gamma_0^c(1) + \sum_m \#_m = (\#_\ell \in \Gamma_0 - \#T_{\Gamma_0}) N = \#_0$$  \hfill (107)

The leading term of $Z_0^N$ is $P_0(1)$, whose divergence degree is $\#_0$.

$$Z_0^N = C^{V_{\Gamma}}[\delta(1)]^\#_0 [1 + O(\delta^{-1}(1)) + O(\lambda)]$$  \hfill (108)

For $Z_N$, since the boundary is separated into two parts, the most divergent pattern $P_A(\pi_n)$ is the one such that its corresponding coarse-grained graph has only two coarse-grained nodes $A$ and $B$, which are connected by the minimum number of links $\min(\#_\ell \in \partial_{AB})$, whose divergence degree is

$$\#_{AB} = \#_A + \#_B + \min(\#_\ell \in \partial_{AB})$$

$$= (\#_\ell \in \Gamma_{AB} - \min(\#_\ell \in \partial_{AB}) - \#T_A - \#T_B) N + \min(\#_\ell \in \partial_{AB})$$

$$= (\#_\ell \in \Gamma_{AB} - \#T_A - \#T_B) N + (1 - N) \min(\#_\ell \in \partial_{AB})$$

$$= \#_0 + (1 - N) \min(\#_\ell \in \partial_{AB})$$  \hfill (109)

where the second equality is in terms of (96) and the forth equality is because $\#_\ell \in \Gamma_{AB} = \#_\ell \in \Gamma_0$ and $\#T_A + \#T_B = \#T_{\Gamma_0}$.  \hfill (96)

11 Since the boundary is coarse-grained into two nodes in $\Gamma_{AB}$, there are one more node in $\Gamma_{AB}$ than in $\Gamma_0$,

$$V_{\Gamma_{AB}} = V_{\Gamma_0} + 1$$  \hfill (110)

Thus the number of the branches of the MST in $A$ and $B$ is equal to the number of the MST branches in $\Gamma_0$

$$\#_A + \#_B = (V_A - 1) - (V_B - 1) = V_{\Gamma_{AB}} - 2 = V_{\Gamma_0} - 1 = \#T_{\Gamma_0}$$  \hfill (111)
Let us consider a graph $\Gamma_{AB}(\pi_m)$ and its corresponding coarse-grained graph $\Gamma_{c,AB}(\pi_m)$. The divergence degree of $\Gamma_{AB}(\pi_m)$ is given as

$$\# \Gamma_{AB}(\pi_m) = \# \Gamma_{c,AB}(\pi_m) + \sum_{m=1, \ldots, M, A, B} \# m$$

(112)

where $\# m$ is given by (96)

$$\# m = (L_m - \# T_m) N$$

(113)

Adapting the same argument as for $Z_N^0$, because of the integral over $h_n$, $\# \Gamma_{c,AB}(\pi_m)$ should not be bigger than the following quantity

$$\# \Gamma_{c,AB}(\pi_m) \leq \sum_{(mm') \notin T^A_{\Gamma_{c,AB}(\pi_m)} \cup T^B_{\Gamma_{c,AB}(\pi_m)}} L_{mm'} \chi(\varpi_{mm'})$$

$$+ \sum_{(mm') \in T^A_{\Gamma_{c,AB}(\pi_m)} \text{ or } T^B_{\Gamma_{c,AB}(\pi_m)}} (L_{mm'} - 1) \chi(\varpi_{mm'})$$

(114)

where we assume $m < m'$ in order to avoid double counting, and $T^A_{\Gamma_{c,AB}(\pi_m)}$ and $T^B_{\Gamma_{c,AB}(\pi_m)}$ are the MST rooting from coarse-grained nodes $A$ and $B$, respectively. The right hand side of the above formula corresponds to the divergence degree of pattern $P_A(\pi_m)$ on a graph $\Gamma_{c,AB}(\pi_m)$ with all $h_n = 1$, which differs from $\Gamma_{AB}(\pi_m)$ by $T^A_{\Gamma_{c,AB}(\pi_m)}$ and $T^B_{\Gamma_{c,AB}(\pi_m)}$, i.e.

$$\Gamma_{c,AB}(\pi_m) \equiv \Gamma_{AB}(\pi_m) \setminus \{T^A_{\Gamma_{c,AB}(\pi_m)}, T^B_{\Gamma_{c,AB}(\pi_m)}\}$$

(115)

As presented in section 2, the major difference between [23, 27] and our paper is that we are considering the gauge transformation $h_n$ on each node $n$. When all $h_n$ are set to be the identity, our $Z_N$ and $Z_N^0$ simplify to the ones in [23, 27] up to overall normalization. In this case, as shown in [23, 27], the patterns which gives only one domain wall for $\Gamma_{AB}$ have higher divergence degree than the divergent degree of multi-domain walls, which in our language means that the patterns whose corresponding coarse-grained graph contains only two coarse-grained nodes are more divergent than the patterns $P_A(\pi_m)$, which give more than two coarse-grained nodes. So the divergence degree of the pattern $P_A(\pi_m)$ on the graph $\Gamma_{c,AB}(\pi_m)$ is not bigger than the pattern $P_A(1, C_0)$. So we have

$$\# \Gamma_{AB}(\pi_m) = \# \Gamma_{c,AB}(\pi_m) + \sum_{m=1, \ldots, M, A, B} \# m$$

$$\leq \# \Gamma_{c,AB}(\pi_m) + \sum_{m=1, \ldots, M, A, B} \# m$$

$$\leq \# A + \# B + \# \ell \in \partial_{AB} = \# 0 + (1 - N) \# \ell \in \partial_{AB}$$

$$\leq \# AB = \# 0 + (1 - N) \min(\# \ell \in \partial_{AB})$$

(116)

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It follows that the amplitude $Z_N$ is

$$Z_N = C^{11 \gamma}[\delta(1)]^\#_a + (1-N)^{\min(\#_{\ell \in \partial AB})}[1 + \mathcal{O}(\delta^{-1}(1)) + \mathcal{O}(\lambda)] .$$  \hspace{1cm} (117)

Finally, the $N$th order Rényi entropy $S_N$ is then:

$$e^{(1-N)S_N} = \frac{Z_N}{Z_0} = [\delta(1)]^{(1-N)^{\min(\#_{\ell \in \partial AB})}}[1 + \mathcal{O}(\delta^{-1}(1)) + \mathcal{O}(\lambda)] .$$  \hspace{1cm} (118)

When $N$ goes to 1, $S_N$ becomes the entanglement entropy $S_{EE}$. The leading term of the entanglement entropy $S_{EE}$ is therefore

$$S_{EE} = \min(\#_{\ell \in \partial AB}) \ln \delta(1) \quad ,$$  \hspace{1cm} (119)

which can be understood as the Ryu-Takayanagi formula in a GFT context. The minimal number of links $\min(\#_{\ell \in \partial AB})$ represents the minimal surface area which separates the bulk.

Before moving on to a different derivation of the same result, we want to clarify the interpretation of this calculation.

The definition of the expectation value (54) in the GFT language shows that the exponential of $S_N$ can be interpreted as a GFT $2N$-point function, at least within the limits of the approximation made, focusing on the average over group field functions at each node, without recasting the whole generalized tensor network as a GFT correlation function. As shown in previous sections, the GFT amplitudes can in turn be written, by standard perturbative expansion, as a sum of Feynman amplitudes associated to Feynman diagrams, each of which corresponds to a different discretized “space-time” with fixed boundary, with the Feynman amplitude defining (for quantum gravity models) a lattice path integral for gravity discretised on the corresponding cellular complex. This allows a tentative (and partial) interpretation of the entropy formula we have derived, in geometric spatiotemporal terms. It implies, in fact, that, in the calculation of the entropy, not only the information of a time-slice of a space-time is considered, as encoded in a given network, but also its full quantum dynamics. This, at least, is true when the complete GFT partition function (for quantum gravity models) is employed in the computation of the entropy. The leading term, the free GFT amplitude, captures only a sector of that full quantum dynamics. With the specific (trivial) choice of kinetic term we have used, the quantum dynamics can at best correspond to (summing over) static space-times. When $N$ goes to 1, in particular, the amplitude becomes the trivial propagation of GFT states, with any given network propagating to itself.
This corresponds exactly to the context (static space-time) in which the Ryu-Takayanagi formula is usually derived. In other words, our calculation provides a realization of the Ryu-Takayanagi formula, at least in one extremely simple case, within the full dynamics of a non-perturbative approach to quantum gravity, the group field theory formalism, which can also be seen as a different definition of loop quantum gravity. Our result also shows that the same formalism allows to compute non-perturbative quantum gravity corrections to the Ryu-Takayanagi formula, by including the contributions from the GFT interaction term into the amplitude (as well as considering different choices for the GFT kinetic term).

IV. RYU-TAKAYANAKI FORMULA FOR SPIN-NETWORK STATES

We want now to perform a similar calculation of the Ryu-Takanayagi entropy using a different truncation of a generic GFT state, reformulated as a tensor network. We use a given linear combination of spin networks, corresponding to a specific assignment of spins to the links of the network, and thus to the tensors associated to its nodes.

As presented in Section 2, the spin representation of a GFT network is spin-network, in which each node is colored by a tensor $\phi_{jm}^{ijkl}$

$$\phi_{jm}^{ijkl} = \sum_{\mathbf{p}} i_p \psi_{pm}, \quad \{ \phi_{jm}^{ijkl} \} = \sum_{\mathbf{m}} \phi_{jm}^{ijkl} \langle j, m \rangle \in \otimes_{\ell} H_{j_{\ell}}$$

and each link is colored by matrix $M_{j_{mn}}^{ijkl}$

$$\{ M_{j_{mn}}^{ijkl} \} = \sum_{\mathbf{m}} M_{j_{mn}}^{ijkl} \langle j, m \rangle \otimes \langle j, m' \rangle \in \otimes_{j} H_{j}^{\otimes 2}$$

where $H_{j}$ is the spin-$j$ irreducible representation of SU(2).

A spin-network has a clear geometric interpretation. The graph $\Gamma$ is the dual of a 3d cellular complex. When all nodes are 4-valent, the graph is dual to a 3d simplicial complex. Each node is dual to a tetrahedron and each link is dual to a triangle. The area of the triangle is given by the spin-$j$ irreducible representation associated with the dual link of the triangle. More precisely, the area $A_{\ell}$ is

$$A_{\ell} = 8\pi \gamma \sqrt{j_\ell(j_\ell + 1)} \ell_{p}^{2} \quad ,$$

where $\Gamma$ is the Barbero-Immirzi parameter and $\ell_{p}$ is the Planck length (while this results follows both from a canonical quantization of General Relativity in the continuum, and from
the geometric quantization of simplicial geometries, the identification of the length scale with the Planck length is, of course, natural from the first perspective only).

A detailed analysis (see e.g. \[83–85\]) shows that the semi-classical regime of loop quantum gravity states, in which the Regge-Einstein gravity can be recovered, at least at the kinematical level, in the sense of approximating smooth geometries with simplicial ones, is at a scale intermediate between the Planck scale $\ell_p$ and the average background curvature scale $L_\Lambda$, which means that if we are working on this regime, area $A_\ell$ of the triangle should be

$$\ell_p^2 \ll A_\ell \ll L_\Lambda^2$$

(123)

Together with the relation $A_\ell/L_\Lambda^2 \sim \gamma^{-1}j^{1/2} \ll 1$ uncovered in \[83\], the above regime is equivalent to

$$\frac{1}{j} \ll \gamma \ll \frac{1}{j^{1/2}}$$

(124)

In a semi-classical regime, then, one has $A_\ell \approx \gamma j \ell_p^2$.

In \[51\], a special choice of $M^j_{mm'}$

$$M^j_{mm'} = \langle j, m | n^\dagger e^{-\pi \gamma L_z - \exp(1-2\pi \gamma L_z)} n' | j, m' \rangle$$

(125)

has been considered, with the property that the leading order of the entanglement entropy between the two $H_j$ on a link is proportional to the same area $A_\ell \approx \gamma j \ell_p^2$ in the semi-classical regime. In (125), $n$ and $n'$ are SU(2) elements; $L_z$ is the $SU(2)$ generator in $z$-axis. We use the same choice for $M^j_{mm'}$ in our calculation to obtain the Ryu-Takayanaki formula.

Considering the same graph $\Gamma$ as in the previous subsection, the spin-network state $|\Psi_\Gamma^j\rangle$ and its corresponding density matrix $\rho$ are given as

$$|\Psi_\Gamma^j\rangle \equiv \bigotimes_\ell \langle M^j_{\ell} \bigotimes_n \phi^{J_{\ell}^{a_{\ell} n}}_n \rangle, \quad \rho \equiv |\Psi_\Gamma^j\rangle \langle \Psi_\Gamma^j|.$$  

(126)

Just as in the previous calculation, we divide boundary $\partial \Gamma$ into two parts $A$ and $B$. The $N$th Rényi entropy is

$$e^{(1-\gamma)S_N} = \frac{Z_N}{Z_0^N} = \frac{\text{Tr}[\rho^N \mathcal{P}(\pi_0; N, d)]}{\text{Tr}(\rho)^N} = \frac{\text{Tr} \left[ \bigotimes_\ell \rho^N_\ell \bigotimes_n \mathcal{E}(\rho^N_n) \mathcal{P}(\pi_0; N, d) \right]}{\text{Tr} \left[ \bigotimes_\ell \rho^N_\ell \bigotimes_n \mathcal{E}(\rho^N_n) \right]}.$$  

(127)

The first key step is to calculate $\mathcal{E}(\rho^N_n)$. Because the gauge symmetry is already encoded in the intertwiner $\tilde{\ell}$ for $\phi^{\tilde{\ell}}_m$, $\phi^{\tilde{\ell}}_m$ is not a gauge symmetric tensor, which is in the invariant space
of $H^\otimes 4$ as introduced in Section 2, but rather an ordinary tensor in $\otimes H_j$. So the average
over $\rho_n^N$ can be performed in the same way as the one shown in [51]:

$$
E(\rho_n^N) \equiv \int d\phi^j f(\phi^j) (|\phi^j\rangle \langle \phi^j|)^{\otimes N} \equiv \int_{U(D)} dU f(\phi^j) \left(U|\phi_0^j\rangle \langle \phi_0^j| U^\dagger\right)^{\otimes N},
$$

(128)

where $f(\phi^j)$ is a distribution of $\phi^j$ and $U$ is the group element in the unitary group $U(D)$, in which $D = \prod_{\ell \in n} d_{j^\ell}$. $f(\phi^j)$ is invariant under the transformation of $U(D)$ and in our following calculation we focus on either the uniform or the Gaussian distribution, which keep the main calculation unchanged up to an overall normalization that will be canceled in the final result.

Because of Schur’s lemma, $E(\rho_n^N)$ is the invariant tensor in $(\otimes H_j)^{\otimes N}$, which can be written as a sum of permutations

$$
E(\rho_n^N) = \mathcal{C} \sum_{\pi_n \in \mathcal{S}_N} \mathcal{P}(\pi_n; N, D) = \mathcal{C} \sum_{\pi_n \in \mathcal{S}_N} \prod_{\ell \in n} \mathcal{P}(\pi_n; N, d_{j^\ell})
$$

(129)

where $\mathcal{C}$ is an normalization factor which depends on the distribution. Then $Z_N$ and $Z_0^N$ can be written as a sum of different patterns $\mathcal{P}(\pi_n)$

$$
Z_N = \mathcal{C}^V \sum_{\pi_n \in \mathcal{S}_N} \mathcal{P}_A(\pi_n), \quad Z_0^N = \mathcal{C}^V \sum_{\pi_n \in \mathcal{S}_N} \mathcal{P}_0(\pi_n)
$$

(130)

where $\#$ is the number of nodes in $\Gamma$. $\mathcal{P}_A(\pi_n)$ and $\mathcal{P}_0(\pi_n)$ can be written as products of link values $\mathcal{L}(\pi_n, \pi'_n)$

$$
\mathcal{P}_A(\pi_n) = \prod_{\ell \in \Gamma} \mathcal{L}_\ell(\pi_n, \pi'_n) \prod_{\ell \in A} \mathcal{L}_\ell(\pi_n, \pi_0\pi_A) \prod_{\ell \in B} \mathcal{L}_\ell(\pi_n, 1)
$$

(131)

$$
\mathcal{P}_0(\pi_n) = \prod_{\ell \in \Gamma} \mathcal{L}_\ell(\pi_n, \pi'_n) \prod_{\ell \in \partial \Gamma} \mathcal{L}_\ell(\pi_n, 1)
$$

(132)

where $\mathcal{L}_\ell(\pi, \pi')$ is defined as

$$
\mathcal{L}_\ell(\pi, \pi') \equiv \text{Tr}[\mathcal{P}(\pi; N, d_{j^\ell}) \rho_n^N \mathcal{P}(\pi'; N, d_{j^\ell})]
$$

(133)

Suppose $\varpi \equiv (\pi')^{-1} \pi = \prod_i C_i$, where $C_i$ is an $r_i$-cycle, and impose (125) into (133). $\mathcal{L}(\pi, \pi')$ becomes

$$
\mathcal{L}(\pi, \pi') = \prod_{i=1}^{\chi(\varpi)} \chi_j \left(e^{-r_{i}2\pi\gamma L_z-r_{i} \exp(1-2\pi\gamma L_z)\frac{1}{2\pi\gamma}}\right)
$$

(134)
In the semi-classical regime \(^{(124)}\), the leading contribution of \( \mathcal{L}(\pi, \pi') \) is obtained as

\[
\mathcal{L}(\pi, \pi') \approx \prod_{i=1}^{\chi(\varpi)} \frac{1}{r_i} e^{-1 + (1 - r_i)2\pi \gamma j - r_i \exp(1 - 2\pi \gamma j)}
\]

\[
= e^{-\chi(\varpi) + (\chi(\varpi) - N)2\pi \gamma j - N \exp(1 - 2\pi \gamma j)} \prod_{i=1}^{1} \frac{1}{r_i}
\]\n
\((135)\)

A detailed calculation from \((134)\) to \((135)\) can be found in the appendix. When \(\varpi = 1\), i.e. \(\pi = \pi'\) and \(\chi(\varpi) = N\), \(\mathcal{L}(\pi, \pi)\) is then

\[
\mathcal{L}(\pi, \pi) \approx e^{-N - N \exp(1 - 2\pi \gamma j)}
\]\n
\((136)\)

It is straightforward to check that \(\mathcal{L}(\pi, \pi) \geq \mathcal{L}(\pi_1, \pi_2)\). In fact, because the sum of \(r_i\) equals to \(N\), \(\mathcal{L}(\pi, \pi)\) can be rewritten as

\[
\mathcal{L}(\pi, \pi) = \prod_{i=1}^{\chi(\varpi)} e^{-r_i - \frac{\exp(1 - 2\pi \gamma j)}{2\pi \gamma}}
\]\n
\((137)\)

Then the ratio between \(\mathcal{L}(\pi_1, \pi_2)\) and \(\mathcal{L}(\pi, \pi)\) is

\[
\frac{\mathcal{L}(\pi_1, \pi_2)}{\mathcal{L}(\pi, \pi)} = \prod_{i=1}^{\chi(\varpi)} \frac{e^{-1 + (1 - r_i)2\pi \gamma j - r_i \exp(1 - 2\pi \gamma j)}}{r_i e^{-r_i - \frac{\exp(1 - 2\pi \gamma j)}{2\pi \gamma}}} = \prod_{i=1}^{\chi(\varpi)} \frac{e^{(1 - r_i)(2\pi \gamma j - 1)}}{r_i} \leq 1
\]\n
\((138)\)

The last inequality holds because \(r_i \geq 1\) and in the regime \(^{(124)}\) \(\gamma j \gg 1\). The equality holds if and only if \(\pi_1 = \pi_2\).

If we assume that all \(j_\ell\) are in the same order of magnitude, because of \((138)\), one can observe immediately that the leading term of \(Z_0^N\) is \(\mathcal{P}_0(1)\), i.e. the permutation group for all nodes is \(\pi_n = 1\). Suppose there are \(L_i\) internal links and \(L_e\) external links in \(\Gamma\), then

\[
Z_0^N \approx C_{V} \prod_{\ell} e^{-N - N \exp(1 - 2\pi \gamma j_\ell)}
\]\n
\((139)\)

The \(N\)th order Rényi entropy becomes

\[
e^{(1 - N)S_N} = \frac{Z_N}{Z_0^N} \approx \sum_{\pi_n} \prod_{\ell} \prod_{i} \frac{\chi(\varpi_\ell) e^{(1 - r_i)(2\pi \gamma j_\ell - 1)}}{r_i}
\]\n
\((140)\)

As shown in \([27]\), in order for the single domain wall pattern to contribute the most to the Rényi entropy, when three domain walls intersect, they should satisfy

\[
\prod_{i} \frac{\chi(\varpi_1)}{r_i} \frac{e^{(1 - r_i)(2\pi \gamma j_1 - 1)}}{r_i} \geq \prod_{i} \frac{\chi(\varpi_2)}{r_i} \frac{e^{(1 - r_i)(2\pi \gamma j_2 - 1)}}{r_i} \prod_{i} \frac{\chi(\varpi_3)}{r_i} \frac{e^{(1 - r_i)(2\pi \gamma j_3 - 1)}}{r_i}
\]\n
\((141)\)
where $\varpi_1 \varpi_2 \varpi_3 = 1$. The above inequality can be simplified to

$$e^{[C(\varpi_2) + C(\varpi_3) - C(\varpi_1)](2\pi \gamma j - 1) - \ln N} \geq 1,$$

where $C(\varpi)$ is the Cayley weight of a permutation $\varpi$ which satisfies the triangular inequality $C(\varpi_1 \varpi_2) \leq C(\varpi_1) + C(\varpi_2)$. In general, when $C(\varpi_1) < C(\varpi_2) + C(\varpi_3)$, the above inequality is satisfied because when $\gamma j \gg 1$ the exponential part of the inequality is dominant. When $C(\varpi_1) = C(\varpi_2) + C(\varpi_3)$, one can check that the inequality is satisfied at least for $N \leq 3$.

Since we are only interested in the entropy while taking the limit $N \to 1$, this inequality is well satisfied. The Rényi entropy $S_N$ for small $N$ is given as

$$e^{(1 - N)S_N} \approx \prod_{\ell \in \partial_{AB}} \exp \left((1 - N)(2\pi \gamma j\ell - 1) - \ln N\right).$$

When $N$ goes to zero, we have

$$S_{EE} \approx \sum_{\ell \in \partial_{AB}} \left[2\pi \gamma j\ell - 1 - \lim_{N \to 1} \frac{\ln N}{1 - N}\right] = \frac{A_{\partial_{AB}}}{4\ell_p^2},$$

which is exactly the Ryu-Takayanagi formula. Comparing with the calculation in [27], we both reproduce the Ryu-Takayanagi formula from the spin-network state in the semi-classical regime IV of loop quantum gravity and GFT states. This gives further support to the expectation that a classical gravitational theory can be recovered in this formalism. Differently from [27], however, our result directly relies on the fundamental degrees of freedom of the theory.

V. RANDOMNESS AND UNIVERSALITY

The dictionary we have established between GFT states and (generalized) random tensor networks suggest the potential for useful cross-over of results across these two research areas. In particular, one can already envisage a direct application of results concerning the quantum dynamics of GFT models and the statistical properties of random tensor models to problems

Using the geometric inequality, the left hand side of the above inequality becomes

$$\prod_{i}^{x(\varpi_2)} r_i \prod_{i}^{x(\varpi_3)} r_i \geq \left(\frac{C(\varpi_2) + 1)(C(\varpi_1) - C(\varpi_2) + 1)}{\left(\frac{N - C(\varpi_1)}{N - C(\varpi_1)}\right)^{N - C(\varpi_1)}}\right)^{N - C(\varpi_1)} \geq (C(\varpi_1) + 1) \left(\frac{N - C(\varpi_1)}{N}\right)^{N - C(\varpi_1)}$$

This simplification is very rough since one has to keep $\frac{N}{N - C(\varpi_1)}$ to be integer. Even in this approximate situation, we could find that it is bigger than 1 when $N$ is a bit smaller than 3.
in statistical mechanics and condensed matter that can be formulated in terms of random
tensor networks.

Indeed, our path integral analysis generalises the statistical derivation given in [23], where
the random character of the tensors allowed to map the computation of typical Rényi en-
tropies to the evaluation of partition functions of generalized Ising models with inverse
temperature $\beta \propto \log D$, $D$ being the dimension of each leg of each tensor in the network.
Interestingly, in the original work, the form of the averaged entropies was derived only in the
large $D$ limit, where the fluctuations of the partition functions are effectively suppressed.
In the large $D$ (low temperature) limit, corresponding to the long-range ordered phase for
the Ising models, the entropies of a boundary region can be directly related to the energy
of a domain wall between different domains of the order parameter: the Ising action can
be estimated by the lowest energy configuration and the minimal energy condition of the
domain wall naturally leads to the RT formula.

One set of results that appears immediately useful in this context concerns universality
properties of probability distributions over random tensors, in the limit of large $D$ [86]. They
represent a generalization to tensor distributions of the central limit theorem for ordinary
probability distributions.

Indeed, a recently proved universality theorem for random tensor fields [86] states that
a rank-d random tensor whose entries are $N^d$ independent, identically distributed, complex
random variables, and whose distribution is a trace invariant (of the type defining the
interactions of tensorial GFTs as well), converges in distribution in the large $D$ limit to
the distributional limit of a Gaussian tensor model, namely a Gaussian tensor field theory.
This is already quite remarkable. However, a second, stronger, universality result [86] states
that under only the assumption that the joint probability distribution of tensor entries is
invariant, assuming that the cumulants of this invariant distribution are uniformly bounded,
the large $D$ limit the tensor distribution again converges to the distributional limit of a
Gaussian tensor model.

We expect these theorems to have direct applicability to random tensor networks, and
even to the generalized class corresponding to the infinite dimensional group fields, where
the large $D$ limit refers to the regime in which any UV cut-off on group representations is
removed.

The key point to be careful about is that such theorems generally apply to distributions
of invariant tensor observables, constructed out of trace (bubble) invariants for bipartite
d-colored graphs \[10\]. Therefore, it does not directly apply to simple products of tensors as
we have dealt with in this paper. However, one may wonder how much of such universal
behavior survives for generic graphs when distributions of generic tensor observable are
considered, e.g. including polynomials made by contractions of tensors which leave some
indices free, as for the case of a contracted tensor network state associated to an open graph.

Intuitively, if one randomizes tensors at the nodes independently of contractions, one
can still rely on such results, to some extent, but the conclusions become much less solid,
because contractions do affect the scaling of the tensors. Much more solid would be to treat
the whole tensor network as an observable in a random tensor or GFT model; then, for
tensor networks associated to d-colored graphs (trace invariants), the universality theorems
would apply, thereby indicating a new direction for further characterizations of the tensor
network states. We postpone this type of evaluations to future work, alongside the complete
reformulation of tensor network states and their statistical average within the 2nd quantized
GFT framework.

VI. CONCLUSIONS

Let us summarize our results in this paper. We have established a precise dictionary
between GFT states and (generalized) random tensor networks. This dictionary also implies,
under different restrictions on the GFT states, a correspondence between LQG spin network
states and tensor networks, and a correspondence between random tensors models and tensor
networks. Next, we have computed the Rényi entropy and derived the RT entropy formula,
for GFT and spin network techniques, first using a simple approximation to a complete
definition of a random tensor network evaluation seen as a GFT correlation function, but still
using a truly generalized tensor network seen as a GFT state, and then considering directly a
spin network state as a random tensor network. This elucidates further the correspondence
and its potential. Finally, we have discussed how universality theorems for random tensor
models can be applied to tensor network states, as a first example of application of results
from the theory of random tensors and GFT to tensor networks. We are convinced that
these results can be just the beginning of many further developments, made possible by the
fertile meeting between tensor networks and fundamental quantum gravity, along the lines
we have established.

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Appendix A: From (134) to (135)

In this appendix we perform the calculation from (134) to (135). $\mathcal{L}(\pi, \pi')$ is given by (134). Let us denote $2\pi\gamma$ as $c$ for simplicity, then $\mathcal{L}(\pi, \pi')$ can be written as

$$ \mathcal{L}(\pi, \pi') = \prod_{i=1}^{\chi(\pi)} \chi_j \left( e^{-rc_Lz - r \exp(1 - cLz)} \right) \equiv \prod_{i=1}^{\chi(\pi)} I_i. \quad (A1) $$

$I_r$ can be written in terms of SU(2) coherent state as

$$ I_r = dj \int dn \langle j, j| n^\dagger e^{-rc_Lz - r \exp(1 - cLz)} n| j, j \rangle = dj \sum_k (-)^k \frac{r^k e^k}{k! c^k} \int dn \langle j, j| n^\dagger e^{-(r+k)cLz} n| j, j \rangle \equiv dj \int dn e^{S_r^{(k)}} \equiv dj \int dn e^{S_r} \quad (A2) $$

where $S_r$ is the total action and $S_r^{(k)} \equiv 2j \ln \langle \uparrow| n^\dagger e^{-c(r+k)Lz} n| \uparrow \rangle$ and $| \uparrow \rangle \equiv | \frac{1}{2}, \frac{1}{2} \rangle$. In the semi-classical regime of loop gravity, i.e. the large spin-$j$ regime, the leading contribution of $I_r$ is from the critical point of $S_r^{(k)}$, which is the solutions of the equations of motion

$$ \delta_n S_r^{(k)} = 0 \quad \Rightarrow \quad n^\dagger e^{-c(r+k)Lz} n = e^{-\alpha Lz} \quad (A4) $$

One can obtain the solutions

$$ n^\dagger L_z n = \pm L_z, \quad \alpha^\pm = \pm c(r+k) \quad (A5) $$

Bring the solutions back to $I_r$, we can get

$$ I_r \sim dj \sum_{\epsilon=\pm} \frac{e^{S_r^{(\epsilon)}}}{\sqrt{\det H_\epsilon}} \equiv \sum_{\epsilon=\pm} I_r^{(\epsilon)} \quad (A6) $$
where $S^r_{r0}$ is the total action $S_r$ on the critical point

$$S^r_{r0} \equiv -\epsilon rcj - r \frac{\exp (1 - \epsilon cj)}{\gamma} \quad (A7)$$

and $H^r_r$ is the Hessian matrix of $S_r$

$$H^r_r \equiv \frac{1}{2} \delta^2 S_r|_c \quad (A8)$$

After perform the second derivation on $S_r$, one can obtain

$$\text{det} \, H^r = 4j^2 r^2 (-c + \exp (1 - \epsilon cj))^2 \quad (A9)$$

In the semi-classical and low energy limit

$$\text{det} \, H^r \sim 4j^2 N^2 \exp 2 (1 - \epsilon cj) \quad (A10)$$

Then $I^r_r$ becomes

$$I^r_r \sim \exp \left( -1 + \epsilon cj (1 - r) - r \frac{\exp(1-\epsilon cj)}{c} \right) \quad (A11)$$

One can observe that $I^+_r \gg I^-_r$ since when $\epsilon = -$, in the large spin regime $I^-_r$ goes to zero. $I_r$ thus becomes $I^+_r$, which is one of the term in the product of (135).

$$I_r \approx I^+_r = \frac{\exp \left( -1 + cj (1 - r) - r \frac{\exp(1-cj)}{c} \right)}{r} \quad (A12)$$

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