Topological and Entanglement Properties of Resonating Valence Bond wavefunctions

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We examine in details the connections between topological and entanglement properties of short-range resonating valence bond (RVB) wave functions using Projected Entangled Pair States (PEPS) on kagome and square lattices on (quasi-)infinite cylinders with generalized boundary conditions (and perimeters with up to 20 lattice spacings). Making use of disconnected topological sectors in the space of dimer lattice coverings, we explicitly derive (orthogonal) “minimally entangled” PEPS RVB states. For the kagome lattice, using the quantum Heisenberg antiferromagnet as a reference model, we obtain the finite size scaling with increasing cylinder perimeter of the vanishing energy separations between these states. In particular, we extract two separate (vanishing) energy scales corresponding (i) to insert a vison line between the two ends of the cylinder and (ii) to pull out and freeze a spin at either end. We also investigate the relations between bulk and boundary properties and show that, for a bipartition of the cylinder, the boundary Hamiltonian defined on the edge can be written as a product of a highly non-local projector, which fundamentally depends upon boundary conditions, with an emergent (local) $su(2)$-invariant one-dimensional (superfluid) $t$–$J$ Hamiltonian, which arises due to the symmetry properties of the auxiliary spins at the edge. This multiplicative structure, a consequence of the disconnected topological sectors in the space of dimer lattice coverings, is characteristic of the topological nature of the states. For minimally entangled RVB states, it is shown that the entanglement spectrum, which reflects the properties of the (gapless or gapped) edge modes, is a subset of the spectrum of the local Hamiltonian, e.g. half of it for the kagome RVB state, providing a simple argument on the origin of the topological entanglement entropy $S_0 = -\ln 2$ of the $Z_2$ spin liquid. We propose to use these features to probe topological phases in microscopic Hamiltonians and some results are compared to existing DMRG data.

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I. INTRODUCTION

Conventional ordering in condensed matter systems is traditionally associated to symmetry-breaking and to the existence of a local order parameter (Landau theory). Topologically ordered phases of matter⁶ offer completely new classes of systems for which the ground state (GS) degeneracy depends on topology (disc, cylinder, torus, etc…). The (short-range singlet) Resonating Valence Bond (RVB) wavefunction proposed by Anderson⁷ as the parent Mott insulator of high-temperature superconductors is a celebrated example. Such topological phases carry emerging fractionalized excitations and raise growing attention due to their potential to realize fault-tolerant setups for quantum computing.⁸

Experimental and theoretical search for topological liquids in quantum antiferromagnets⁹ and in related microscopic models¹⁰ has been a long-standing quest. One major problem is the existence of many possible (non-magnetic) nearby competing states like valence bond crystals¹¹ (spontaneously breaking lattice symmetry), clearly evidenced e.g. in quantum dimer models.¹² Recent advances in the Density Matrix Renormalisation Group (DMRG) techniques has reinforced the strong belief that a gapped spin liquid might be stabilized in the nearest neighbor (NN) $S=1/2$ Heisenberg quantum antiferromagnet (HAF).¹³¹⁴ This has also triggered the search for novel theoretical tools capable of better detecting topological order, in particular entanglement measures used in quantum information. A common setup consists of dividing the system into two regions (named $A$ and $B$) and compute the reduced density matrix (RDM) in the GS of e.g. the $A$ subsystem. The entanglement entropy (EE), defined as the Von Neumann entropy of the RDM $S_{V\!N} = -\rho_A \ln \rho_A$, contains an extensive term — proportional to the length of the boundary (area law) — and a universal sub-leading constant, the topological EE. Specific disc-like setups or cylindrical geometries can be used to extract the topological EE.

In fact, $-\ln \rho_A$ can be seen as a (dimensionless) Hamiltonian $H_b$, a key conceptual object. First, its spectrum, the so-called entanglement spectrum (ES), has been conjectured to show a one-to-one correspondence with the spectrum of edge states. This remarkable property was first established in fractional quantum Hall states¹⁵ and, then, in quantum spin systems.¹⁶ Furthermore, Projected Enganged Pair States (PEPS)¹⁷ offer a natural formulation of the relation between bulk and boundary. In Ref. [17] an explicit isometry was constructed which maps the Hamiltonian $H_b$ onto another one $H_b'$ acting on the space of auxiliary spins living at the edge of region $A$, while keeping the spectrum. Furthermore, for various...
two-dimensional (2D) models displaying quantum phase transitions, like a deformed AKLT\cite{18} or an Ising-type\cite{19} model, it was found\cite{20} that a gapped bulk phase with local order corresponds to a boundary Hamiltonian with local interactions, whereas critical behavior in the bulk is reflected in a diverging interaction length of $H_b$.

Entanglement properties of 2D topological phases are less well understood. Rokhsar-Kivelson (RK) wave functions, defined as equal-weight superposition of fully packed dimer coverings, exhibit critical behavior on bipartite lattice,\cite{23} or realize the simplest topological phase, the so-called $\mathbb{Z}_2$ liquid, on frustrated lattices.\cite{23} The topological EE of critical and topological RK wavefunctions have been computed using various topologies\cite{23} and the boundary Hamiltonian corresponding to the GS of Kitaev’s toric code\cite{3} was shown to be non-local.\cite{17} Unfortunately, RK-like wavefunctions are not generic – their ES is completely dispersionless \cite{23} and do not describe real quantum $S = 1/2$ spin systems.\cite{23,29,22} In contrast, (short-range) RVB states, defined as linear superposition of hardcore coverings of non-orthogonal nearest-neighbor $SU(2)$ singlets (see Fig. 1(a)), appear to be closer to physical systems. Very recently, the (Renyi) EE between (finite) cylindrical regions has been computed\cite{23} numerically for the critical,\cite{23,29,22} RVB state on the square lattice. Similarly, the (Renyi) topological EE of SU(2)-symmetric gapped chiral and $\mathbb{Z}_2$ spin liquids was obtained\cite{22} using Kitaev-Preskill prescription. Nevertheless, ES and boundary Hamiltonians of such RVB/spin liquids wavefunctions are unknown.

In this work, we study topological and entanglement properties of both critical (square lattice) and gapped topologically-ordered (kagome lattice) RVB wavefunctions\cite{24} on infinite cylinders making use of simple PEPS representations. Let us describe here the organization of the paper: First, in Sec. \textbf{II} we introduce RVB wavefunctions defined in the space of dimer (hardcore) coverings of square and kagome lattices. On cylinders with generalized boundary conditions, we review the construction of four disconnected topological sectors of dimer coverings (on the kagome lattice). Next, in Sec. \textbf{III} we introduce the PEPS representation of the RVB wavefunctions and, making use of the disconnected topological sectors, explicitly construct four orthogonal RVB states. Using the quantum Heisenberg model as a reference Hamiltonian, we obtain the generic behavior of their energy splittings versus cylinder perimeter. In Sec. \textbf{IV} we introduce a partition of the cylinder and compute the corresponding Reduced Density Matrix (RDM). The (hermitian) operator defined as minus the logarithm of the RDM can be viewed as a boundary Hamiltonian: it is can be naturally expressed in the PEPS formalism as an operator acting on the virtual indices on the edges (up to an isometry). We show that the boundary Hamiltonian can be written as a product of a highly non-local projector, which depends fundamentally on the boundary conditions, by a local one-dimensional $t$–$J$ model, which arises due to the symmetry properties of the auxiliary spins at the boundary and characterizes the (gapless or gapped) edge modes. This multiplicative structure is a direct consequence of the disconnected topological sectors in the space of dimer coverings of the lattice and, therefore, reflects the topological nature of the states. For sake of conciseness, more technical issues such as finite size scalings, etc... are treated in Appendices.

\section{RVB wavefunctions on cylinders}

\subsection{Set-up and boundary conditions}

Let us first start with a square lattice on a cylinder of length $N_b$ and circumference $N_v$ with Open Boundary Conditions (OBC) as depicted in Fig. \textbf{1(a)}. We consider the space of all nearest-neighbor (NN) $|\uparrow\downarrow\rangle - |\downarrow\uparrow\rangle$ singlet coverings of the lattice in such a way that each site belongs to one and only one dimer (so called “hard-core” coverings). Note that all singlets are oriented from one sublattice to the other. The Resonating Valence Bond state is then defined as the equal weight superposition of all such dimer (singlet) coverings. Besides OBC we also consider Generalized Boundary Conditions (GBC) as in Fig. \textbf{1(b)} by freezing some spins at the two boundaries $B_L$ and $B_R$ of the cylinder: in that case, dimers cannot involve these “frozen” sites any more. Because of the local hard-core constraints, the choice of the boundary conditions will affect the physics in the center of the cylinder, even in the limit of an infinitely long one. Similar dimer coverings and RVB wavefunctions can be considered on cylinders with a kagome lattice (see e.g. Fig. \textbf{2}). In that case, singlets are all oriented clockwise in both left and right triangles. It is known that RVB wavefunctions always exhibit short-range spin-spin correlations in two-dimensions (2D) although dimer-dimer correlations can be either short-range (kagome) or critical (square lattice) as mentioned above.

\subsection{Topological sectors}

Here we briefly review the crucial concept of topological sectors in the space of (hard-core) dimer coverings (focusing on the kagome lattice) and show that four RVB wavefunctions belonging to different topological sectors can be constructed on $N_v \times N_b$ cylinders with periodic (open and generalized) boundary conditions in the vertical (horizontal) direction when $N_v$ is even. The case of odd perimeter will also be discussed. For illustration, small $4 \times 2$ and $3 \times 2$ cylinders are drawn for simplicity in Figs. \textbf{2}, \textbf{3} and \textbf{4} but our arguments are valid for any system size.

Let us first consider the case of a cylinder with $N_v$ even. Topological sectors can be defined by considering (i) a closed loop in the vertical direction winding around the cylinder (see Fig. \textbf{2}) and (ii) two open lines along the
FIG. 1: (Color online) Typical valence bond configurations on a $N_v \times N_h$ cylinder with periodic boundary conditions along the vertical ($v$) direction. Ellipses represent singlets of two spins 1/2. Open (a) or generalized (b) boundary conditions on the $B_L$ and $B_R$ ends of the cylinder are considered [GBC can be obtained physically by freezing some spins at the boundaries, e.g. with local magnetic fields]. The RVB wavefunction is defined as the equal-weight superposition of all such configurations (for a fixed realization of $B_L$ and $B_R$).

Crystal directions $h_1$ and $h_2$ at 30° angles w.r.t. the horizontal axis (see Fig. 3), joining the two open ends $B_L$ and $B_R$ of the cylinder. As shown in Figs. 2, 3 for a given configuration, the parities of the numbers of dimers cut by these loops are conserved quantities under translation of the vertical loop (horizontal lines) along the horizontal direction (vertical direction). Since the product of the three parities is constrained to be either even or odd (depending on the choice of $N_h$ and $N_v$), NN dimer configurations can be grouped into four disconnected sectors. Their are “topological” in nature since any local Hamiltonian acting on the space of dimer configurations preserves the sectors.

It is interesting to note that one can go from one topological sector to the other by non-local moves of dimers. For example, let us consider the left configuration of dimers around a closed loop winding around the cylinder. As shown in Fig. 3, by translating any staggered arrangement of dimers along a closed loop winding around the cylinder by one lattice spacing, one permutes (changes) the parities $G_h$ measured along $h_1$ and $h_2$ for $N_h = 4p + 2$ ($N_h = 4p$). Because the space of NN dimer coverings is divided into two disconnected sectors (fixing OBC), two RVB states can first be constructed separately in each sector. Such states should have the same energy density in the middle of the cylinder (for a generic local su(2)-Hamiltonian) since nothing can distinguish the two states locally. However, on a finite cylinder, such RVB states don’t have the lowest variational energy since they break the mirror symmetry w.r.t. the horizontal direction (a symmetry assumed for the Hamiltonian) when $N_h = 4p + 2$. However, by taking their superpositions both with relative plus or minus signs (see Fig. 2), two appropriate variational GS $\Psi^+_{\text{RVB}}$ and $\Psi^-_{\text{RVB}}$ can be defined (strictly orthogonal for $N_h = 4p + 2$). Interestingly, starting from $\Psi^+_{\text{RVB}}$, one can pictorially obtain $\Psi^-_{\text{RVB}}$ by inserting a “vison” line going all the way from the left to the right boundaries of the cylinder e.g. along the $h_1$ direction: the vision operator counts the number of dimers cut by the line and adds a minus sign to the wave function for an odd number of cuts. In other words, the $\Psi^+_{\text{RVB}}$ (no-vison) and $\Psi^-_{\text{RVB}}$ (vison) are states with a definite $Z_2$ flux through the cylinder.

The two states $\Psi^+_{\text{RVB}}$ and $\Psi^-_{\text{RVB}}$ have been constructed for specific OBC for $B_L$ and $B_R$. Shifting by one lattice spacing a line of staggered dimers joining the two ends of the cylinder, will change the parity $G_v$ of the numbers of dimers cut by loops winding around the cylinder, hence providing a change from, let say, the “even” to the “odd” topological sector, as seen in Fig. 3. By applying this second type of non-local move to the two previous $\Psi^+_{\text{RVB}}$ and $\Psi^-_{\text{RVB}}$ wavefunctions, one can then construct four orthogonal variational RVB wavefunctions denominated $\Psi^+_{\text{RVB}}$, $\Psi^-_{\text{odd}}$, $\Psi^+_{\text{even}}$ and $\Psi^-_{\text{odd}}$.

Let us now briefly discuss the case of “odd” cylinders i.e. cylinders with an odd number $N_v$ of unit cells. As shown in Fig. 4, the parity of the number of dimers cut by closed loops encircling the cylinder along the vertical direction alternates along the cylinder. This indicates that two consecutive columns become non-equivalent and the system spontaneously dimerizes in the cylinder direction. By shifting a horizontal line of staggered dimers as before, one switches the parity of the “even” and “odd” columns. This defines two disconnected classes of configurations from which two related RVB states $\Psi^+_{\text{RVB}}$ and
FIG. 3: Two valence bond configurations on a $4 \times 2$ cylinder ($N_v = 4$ even). The two configurations are obtained from each other by translating all dimers (in purple) along a (single) open loop joining the two $B_L$ and $B_R$ ends of the cylinder (and adding extra spins). Such configurations can be distinguished from the parity $G_v = \pm 1$ of the number of dimers cut by any closed loop winding around the cylinder along the vertical direction and, hence, define two different “even” and “odd” topological sectors (and the corresponding RVB states).

FIG. 4: Same as Fig. 3 for a $3 \times 2$ cylinder ($N_v = 3$ odd). $\Psi_{RVB}^{\pm, 1}$ can be constructed as equal weight superposition of all dimer configurations of each class. In the center of long (enough) cylinders, these two RVB states are simply related by a unit translation along the cylinder. Of course, as before, a vision line can be inserted between the two ends of the cylinder to derive two new $\Psi_{RVB}^{\pm, 2}$ wavefunctions.

We finish this Section with the case of the square lattice. Because of the much more constrained nature of dimer configurations on the square lattice, one can construct an extensive number $\propto N_v$ of topological sectors. This will be discussed in more details in Sec. IV.

III. PEPS REPRESENTATION OF RVB STATES

A. Mathematical construction

We start with the square lattice RVB wavefunction (NN $|\uparrow\downarrow\rangle - |\downarrow\uparrow\rangle$ singlets are all oriented from one sublattice to the other) on a cylinder of length $N_h$ and circumference $N_v$, depicted in Fig. 5(a,c), corresponding to an equal weight (and equal sign) summation of all (singlet) dimer coverings. The RVB wavefunction can be expanded in the local $S_z$-basis, $|\Psi_{RVB}\rangle = \sum_S c_S |s_1, s_2, \ldots, s_M\rangle$, where $s_n = 0, 1$ are qubits (representing the two $S_z = \pm 1/2$ spin components) on the $M = N_h N_v$ sites and $S = \{s_n\}$. Such a state can in fact be represented by a $D = 3$ PEPS [Ref.] (up to local unitaries) where each lattice site is replaced by a rank-5 tensor $A^s_{n, \alpha\alpha'; \beta\beta'}$ labeled by one physical index, $s = 0$ or 1, and by four virtual bond indices (varying from 0 to 2) along the horizontal ($\alpha, \alpha'$) and vertical ($\beta, \beta'$) directions. Physically, the absence of singlet on a bond is encoded by the virtual index being "2" on that bond. To enforce
the hardcore dimer constraint, one takes $A_{\alpha,\alpha',\beta,\beta'} = 1$ whenever three virtual indices equal 2 and the fourth one equals $s$, and $A_{\alpha,\alpha',\beta,\beta'} = 0$ otherwise. The amplitudes $c_s$ are then obtained by contracting all virtual indices, except the ones at the ends of the cylinder fixed by boundary conditions, as depicted in Fig. 5(b,d). For the kagome lattice, as shown in Fig. 6, the RVB state can be represented in terms of rank-3 tensors, (i) $A^s$ on the sites – $A_{2,2}^s = A_{4,2}^s = 1$ and zero otherwise – and (ii) on the center of each triangle, $R_{2,2,2} = 1$, and $R_{a,\beta,\gamma} = \epsilon_{a,\beta,\gamma}$ otherwise, with $\epsilon_{a,\beta,\gamma}$ the antisymmetric tensor. One can then group the 3 sites on each unit cell to obtain a rank-5 tensor (the physical dimension is now $2^3 = 8$) connected on an effective square lattice (Fig. 6(b,c)). Note that for the Kagome PEPS, one can find a local parent Hamiltonian for which the degeneracy is equal to 4 on the torus.

In the PEPS formulation the boundary conditions $B_L$ and $B_R$ can be simply set by fixing the virtual states on the bonds “sticking out” at each cylinder end. E.g. open boundary conditions as in Fig. 5(a) are obtained by setting the boundary virtual indices to “2” as shown in Fig. 5(b). Generalized boundary conditions can be realized as in Fig. 5(c,d) by setting some of the virtual indices on the ends to 0 or 1.

B. Topological energy splittings of kagome RVB wavefunctions

More and more numerical data from DMRG simulations support the claim that the NN quantum HAF on the kagome lattice is a topological $\mathbb{Z}_2$ spin liquid. It is therefore interesting (and relevant) to consider the previous topological NN-RVB wavefunctions as variational ground-state ansätze for the NN HAF Hamiltonian on the kagome lattice,

$$H = J \sum_{\langle ij \rangle} S_i \cdot S_j,$$

where $S_i$ is the spin-1/2 operator at site $i$ and $\langle ij \rangle$ stands for all NN bonds of the kagome lattice and the exchange constant is been set to $J = 1$ from now on. Although (i) the (local) parent Hamiltonian of the NN-RVB wavefunction contains much more complicated interaction and, conversely, (ii) the ground-state of the NN HAF is far more involved that a simple NN-RVB (e.g. containing singlets bonds beyond NN), we believe generic features on the finite size energy splitting between the different topological sectors (topological gap) can be obtained by using simple NN RVB wave functions. A schematic picture in Fig. 7 illustrates the expected GS multiplet structure for increasing system size. In the 2D thermodynamic limit, when both cylinder length and perimeter are infinite, one expects all energy splittings to vanish and the GS to become four-fold degenerate.

The PEPS formalism allows to compute exactly the variational energy of the NN RVB wavefunctions on cylinders of perimeter $N_v$ up to $N_v = 10$ and length $N_h \to \infty$. $\Psi_{\text{RVB}}$ is obtained using the local rank-5 tensors described above and OBC. To get $\Psi_{\text{even}}$ one inserts a “vison” line joining the two boundaries of the cylinder by putting a string of $Z = \text{diag}(1,1,-1)$ operators on the bonds. Finally, $\Psi_{\text{odd}}$ are obtained by using GBC for $B_L$ and $B_R$. The energy is computed at the center of the cylinder after full convergence with increasing cylinder length $N_h$ is reached (typically $N_h \sim 10 N_v$ is enough). We have checked numerically that all states possess mirror symmetry of the energy density w.r.t. the horizontal

FIG. 6: (Color online) On the kagome lattice, an effective rank-5 tensor is constructed on each 3-site unit cell. Three site tensors (red dots) carrying the physical indices and two 120-degree tensors (in the center of the shaded triangles) are grouped together (a,b) to construct the basic tensor (c). The kagome lattice is then mapped onto an effective square lattice. A partition of the cylinder in the vertical direction generates $L$ and $R$ edges (thick dotted line).

FIG. 7: Illustration of the energy splitting between the four (variational) RVB wavefunctions for the kagome HAF. From left to right, the cylinder length (at fixed perimeter $N_v$ even) and then, its perimeter are increased to infinity.
axis (as expected from their symmetry) and are uniform (staggered) for $N_v$ even (odd) as illustrated in Fig. 8. Interestingly, for $N_v = 8$ and $N_v = 10$ the lattice $C_{6v}$ symmetry around an hexagon center is almost fully recovered (i.e. the vertical and 30-degree bonds become equivalent). The energy (per site) of the four orthogonal RVB wave functions are plotted in Fig. 8(a) vs $1/N_v$. After averaging the energies of the even and odd states, one obtains very accurate fits of the exponentially fast convergence of the energies of the RVB wave functions in the “+” and “−” topological sectors, with a very short characteristic length-scale $\xi_E \sim 1.0$. The extrapolated energy agrees very well with a recent estimate based on a Gutzwiller-projected superconducting wavefunction$^{31}$. Although this variational energy is much higher than most recent variational estimates$^{29,30,124}$ (between $-0.437$ and $-0.439$), we believe the observed finite size behaviors and energy splittings (topological gaps) should be generic of $\mathbb{Z}_2$ spin liquids. For example, we find that the average over the variational energies of the four RVB wavefunctions exhibits surprisingly small size dependence, in striking correspondence with DMRG results.$^{30,12}$

The splittings between the variational RVB GS defined in Fig. 8(b) are plotted using a logarithmic scale in Fig. 9(b) as a function of the perimeter $N_v$ of the infinite cylinder. Exponential decay of the topological splittings vs $N_v$ are seen revealing two typical length-scales $\xi_{\text{topo}}^1 \sim 0.65$ and $\xi_{\text{topo}}^2 \sim 1.01$, associated to the even-odd and $+−$ gaps, respectively. Note that the dimerization energy of the $\Psi_{\text{RVB}}^{+,1}$ (or $\Psi_{\text{RVB}}^{+,2}$) states follows the same exponential decay as the even-odd topological gaps. For a very long cylinder with fixed boundary conditions, we therefore predict the following finite size scaling of the largest topological splitting (cost of inserting a horizontal vison line),

$$\Delta E_{+\text{−}} \sim 1.06 N_v N_h \exp (-0.99 N_v).$$

The cost of freezing an odd number of spins at the boundary is given by,

$$\Delta E_{\text{eo}} \sim 1.95 N_v N_h \exp (-1.54 N_v),$$

for the case where the state has a definite parity ($G_h = \pm 1$). For a definite $\mathbb{Z}_2$ flux in the cylinder (e.g. a state with or without a vison), moderate corrections occur for perimeter $N_v \leq 8$ as seen in Fig. 9(b). In DMRG, the two different even and odd sectors can be fixed$^{29}$ by moving a site from one end of the cylinder to the other, which would be the same as pinning sites with strong fields on either end. However, it is not clear whether the DMRG algorithm chooses a definite $G_h$ parity or a definite $\mathbb{Z}_2$ flux or none of the two.
IV. BOUNDARY HAMILTONIAN ON INFINITE CYLINDERS

A. Bipartition and reduced density matrix

To define the boundary Hamiltonian of the RVB wavefunctions, we partition the $N_v \times N_h$ cylinder into two half-cylinders of lengths $N_h/2$, as depicted in Fig. 5. Partitioning the cylinder into two half-cylinders (playing the role of two A and B subsystems as defined in the Introduction) reveals two edges L and R along the cut. Ultimately, we aim to take the limit of infinite cylinders, i.e. $N_h \to \infty$ as before.

For a topological state, the boundary Hamiltonian depends on the choice of the wavefunction within the (variational) GS degenerate manifold. In other words, it depends upon (i) the choice of the $B_L$ and $B_R$ cylinder boundaries that impose the parity of $G_v$ ($B_L$ and $B_R$ have to “match”) and (ii) the possible insertion of a horizontal vison line (or equivalently a $Z_2$ flux through the cylinder). For simplicity, we restrict ourselves to the + combination of Fig. 2 (no vison) but still consider arbitrary choices of the boundary conditions at the ends of the cylinder.

The boundary Hamiltonian $\tilde{H}_b$ can be derived from the reduced density operator $\sigma^2 = \exp(-\tilde{H}_b)$ acting on the edge indices, following the procedure given in Ref. [17]. For the kagome lattice, there is no reflection symmetry w.r.t. the cut so the RDM for the left (right) side takes the form

$$\sigma^2_{BL} = \sqrt{\sigma^2_L}\sqrt{\sigma^2_R} \quad (\sigma^2_{BR} = \sqrt{\sigma^2_L}\sqrt{\sigma^2_R})$$

where $\sigma_L$ and $\sigma_R$ are obtained by contracting the tensors of the left and right half-cylinders, respectively, as shown in Fig. 10 (see Ref. [17] for details). Note that $\sigma^2_{BL}$ and $\sigma^2_{BR}$ give identical ES. For clarity, we restrict ourself to $\sigma^2_{BL}$. Ultimately, we are interested in RVB cylinders with infinite lengths in both directions. First, we fix the cylinder perimeter ($N_v = 4, 6, 8$) and take the limit $N_h \to \infty$ as shown in Appendix A (in practice, the RDM for $N_h \sim 10N_v$ is fully converged). The behaviors of the boundary Hamiltonian and the ES as a function of cylinder perimeter is then analyzed (see Appendix B for explicit finite size scalings).

B. Boundary Hamiltonian

1. Disconnected topological sectors in the PEPS representation

The concept of boundary Hamiltonians is described in details in Ref. [17] and, for topological states, in Ref. [32]. Here, we provide the details of their numerical computation for the RVB states. A crucial feature of the topological states is that the RDM depends intrinsically on the choice of the boundary conditions ($B_L = B_R$ for simplicity), even when $N_h \to \infty$. Indeed, the configurations of virtual indices (on the horizontal bonds) of any vertical column is split in two disjointed sectors which, due to local constraints, are conserved from column to column (and hence can be addressed independently from proper choices of $B_L = B_R$). This is directly connected to the partition of the space of NN dimer coverings into disconnected (even and odd for the kagome lattice) topological sectors discussed earlier. Indeed, the number of $2^2$ states on a column of virtual bonds corresponds to the number of bonds with no dimers. Therefore, for the kagome lattice, in a given topological sector, the parity of the number of virtual $|2^2)$ states on the columns is conserved from column to column. Consequently, the boundary Hamiltonian of the Kagome RVB wave function conserves the parity of the number of $|2^2)$ states, as in Kitaev’s toric code. On the square lattice, extra constraints impose the conservation of the difference between the number of $|2^2)$ states between two alternating sublattices on the edge. Hence, although the RDM ($\tilde{H}_b$) acts on all $3^{N_v}$ degrees of freedom of the $L$ (or $R$) edge, in each sector it contains a finite fraction of zero-weight eigenvalues, i.e. a finite fraction of eigenstates of $\tilde{H}_b$ have infinite-energy. Calling $P$ the projector on the finite-energy subspace, we split $\tilde{H}_b$ as $\tilde{H}_b = H_1 + \beta_{\text{topo}}(\text{Id} - P)$, where Id is the $3^{N_v}$ identity operator, $\beta_{\text{topo}} \to \infty$, and $H_1$ is supported by the non-zero eigenvalues sector of the RDM. More precisely, $H_1$ can be factorized as $H_1 = H_{\text{local}}P$ where $H_{\text{local}}$ is a Hamiltonian (shown later to be local) acting on the whole boundary space commuting with $P$ and independent on BC. The Kagome cylinder has only two sectors defined by $P = P_{\text{even}}$ and $P = P_{\text{odd}} = \text{Id} - P_{\text{even}}$, which can be obtained by choosing an even or odd number of “2” external (virtual) indices for $B_L = B_R$, respectively.

On the square lattice cylinder, there are $N_v + 1$ disconnected sectors defined by projectors $P_{\Delta}$ enforcing a fixed difference $N_{2,x} - N_{2,y} = \Delta$ of the numbers of “2” on two X and Y alternating sublattices on the edge, $\Delta = -N_v/2, \cdots, N_v/2$. The fact that $H_1$ is known for all sectors implies that $H_{\text{local}}$ is uniquely determined as will be shown in the next Subsection.
2. Practical derivation of $H_{\text{local}}$

In practice, each numerical calculation is done for a specific choice of the boundary conditions $B_L$ and $B_R$ (for simplicity, we assume here $B_L = B_R$) on the cylinder ends which determines a given conserved sector (mathematically characterized by some projector $\mathcal{P}$), support of the corresponding boundary Hamiltonian. Reversely, all sectors (associated to different projectors $\mathcal{P}$) can be obtained from proper choices of the boundary conditions $B_L = B_R$ (like the sectors defined by the projectors $\mathcal{P}_{\text{even}}$ and $\mathcal{P}_0$ which can be addressed by choosing OBC).

On the Kagome cylinder, the two sectors defined by $\mathcal{P} = \mathcal{P}_{\text{even}}$ and $\mathcal{P} = \mathcal{P}_{\text{odd}} = 1^\otimes N_v - \mathcal{P}_{\text{even}}$ can be obtained by choosing even (e.g. OBC) or odd number of "2" external (virtual) indices on both the left and right boundaries of the cylinder, respectively. We can then construct a “mixed” RDM for the right part,

$$\sigma^2 = \sqrt{\sigma^t L \sigma^t R \sqrt{\sigma^t L}},$$  \hspace{1cm} (4)

by considering the linear superpositions

$$\sigma^t_L = \sigma_{L,\text{even}}^t + \sigma_{L,\text{odd}}^t,$$
$$\sigma^t_R = \sigma_{R,\text{even}}^t + \sigma_{R,\text{odd}}^t,$$  \hspace{1cm} (5)

where $\sigma_{T,p}$ are obtained by contracting the left ($T = L$) and right ($T = R$) half-cylinders (see Ref.17) with appropriate $p$="even" or $p$="odd" parity boundary conditions and the "equal weight" normalization condition,

$$\text{Tr}\{\sigma_{L,\text{even}}^t \sigma_{R,\text{even}}^t\} = \text{Tr}\{\sigma_{L,\text{odd}}^t \sigma_{R,\text{odd}}^t\} = 1.$$  \hspace{1cm} (6)

Since $\sigma_{T,\text{even}}$ and $\sigma_{T,\text{odd}}$ are supported on disconnected subspaces, the mixed RDM splits into orthogonal contributions, $\rho_p = \rho_{\text{even}} + \rho_{\text{odd}}$, where

$$\rho_{\text{even}} = \sqrt{\sigma_{L,\text{even}}^t \sigma_{R,\text{even}}^t \sqrt{\sigma_{L,\text{even}}^t}},$$
$$\rho_{\text{odd}} = \sqrt{\sigma_{L,\text{odd}}^t \sigma_{R,\text{odd}}^t \sqrt{\sigma_{L,\text{odd}}^t}}.$$  \hspace{1cm} (7)

Since $\sigma_p^2$ is supported by the whole Hilbert space, $H_{\text{local}}$ can be uniquely defined by setting $\sigma_p^2 = \exp(-H_{\text{local}})$, enabling a direct computation of $H_{\text{local}} = -\ln \sigma_p^2$ from Eqs. (4) and (5). Reversely, the generic form of the boundary Hamiltonian associated to the (normalized) RDM $\rho_p$ is given by,

$$\tilde{H}_b = H_{\text{local}} \mathcal{P}_p + \beta^\infty \mathcal{P}_{\bar{p}},$$  \hspace{1cm} (8)

with $\beta^\infty \to \infty$ and where $p$ ($\bar{p}$) refers to the "even" ("odd") or "odd" ("even") parity sector.

For the RVB wavefunction on the square lattice, there are $N_v/2 + 1$ orthogonal sectors defined by the projectors $\mathcal{P}_\Delta$ enforcing a fixed difference $|N_{2,A} - N_{2,B}| = \Delta$ on the two A and B alternating sublattices on the one-dimensional edge, $\Delta = 0, 1, \cdots, N_v/2$. We show (for $N_v = 6$) the expansion of these projectors in terms of N-body operators in Fig. 11(a), highlighting clearly their highly non-local character. We construct the “mixed” RDM $\sigma^2 b$ from the linear superposition

$$\sigma_b = \sum_{\Delta = 0}^{N_v/2} \sigma^2 \Delta,$$  \hspace{1cm} (9)

where $\sigma^2 \Delta$ is obtained by contracting the infinite (left or right) half-cylinder (see Ref.17) with appropriate boundary conditions and normalized according to $\text{Tr}(\sigma^2 \Delta) = 1$. The operators $\sigma^2 \Delta$ are supported on orthogonal subspaces which span the whole space of virtual indices on the edge i.e. $\sum \mathcal{P}_\Delta = 1^\otimes N_v$. Therefore, since $\sigma^2 b$ lives on the whole Hilbert space of the edge, one can uniquely define $H_{\text{local}}$ as $H_{\text{local}} = -\ln \sigma^2 b$. Consequently, $H_{\text{local}}$ can be computed numerically using Eq. 9. It also follows that, for each sector, the corresponding boundary Hamiltonian is,

$$\tilde{H}_b = H_{\text{local}} \mathcal{P}_\Delta + \beta^\infty \mathcal{P}_{\bar{\Delta}},$$  \hspace{1cm} (10)

with $\beta^\infty \to \infty$ and $\mathcal{P}_{\bar{\Delta}} = 1^\otimes N_v - \mathcal{P}_\Delta$ is the projector on the complementary subspace.

3. Expansion in terms of N-body operators: numerical results

Next, we wish to explore the non-local/local characters of the $H_1/H_{\text{local}}$ edge operators. Any operator $\mathcal{O}_{\text{edge}}$ acting on the edge can be expanded in terms of $32^N_v$ orthogonal operators. For this purpose, we use a local basis of $9$ (normalized) operators $\{\hat{x}_0, \cdots, \hat{x}_8\}$ which...
act on the local basis of configuration (at some site \(i\), \((\{0\}, \{1\}, \{2\})\)), e.g. \(\hat{x}_0 = 1\), \(\hat{x}_1 = \sqrt{\frac{3}{2}}(\langle 0| - |1\rangle \langle 1|)\) and \(\hat{x}_2 = \frac{1}{\sqrt{3}}(|0\rangle \langle 0| + |1\rangle \langle 1| - 2|2\rangle \langle 2|)\), for the diagonal matrices, complemented by \(\hat{x}_3 = \hat{x}_1^\dagger = \sqrt{3}|0\rangle \langle 1|\) acting as “spin” operators, and \(\hat{x}_4 = \hat{x}_1^\dagger = \sqrt{3}|2\rangle \langle 0|\) and \(\hat{x}_5 = \hat{x}_1^\dagger = \sqrt{3}|2\rangle \langle 1|\) acting as annihilation and creation (hardcore) bosonic operators. The expansion in terms of \(N\)-body operators reads (see Appendix C for more details),

\[
O_{\text{edge}} = c_0 N_v + \sum_{\lambda,i} c_{\lambda,i} \hat{x}_\lambda + \sum_{\lambda,\mu,r,i} d_{\lambda\mu}(r) \hat{x}_\lambda \hat{x}_\mu^i + r + \sum_{\lambda,\mu,\nu,r,r',i} e_{\lambda\mu\nu}(r,r') \hat{x}_\lambda \hat{x}_\mu^i \hat{x}_\nu^{i+r} + \cdots, \tag{11}
\]

where each group of terms involves products of \(N\) (\(1 \leq N \leq N_v\)) on-site \(x_\lambda\) (\(\lambda \neq 0\)) operators. Here the sums are restricted to non-equivalent relative distances and only translations giving distinct sets of sites are performed. The (real) coefficients appearing in (11) have been computed for \(O_{\text{edge}} = \mathcal{P}_{\text{even}}\) (Kagome lattice), \(O_{\text{edge}} = \mathcal{P}_{\Delta}\) (square lattice) and \(O_{\text{edge}} = \mathcal{H}_{\text{local}}\) (Kagome and square lattices) on infinitely-long cylinders of perimeters \(N_v = 6\) and \(N_v = 8\) up to order \(N = 6\).

As seen from the distribution of their weights in Figs. 11(a,b), projectors are highly non-local, conferring a fundamentally non-local character to the boundary Hamiltonian \(H_1\). This is also to be expected for realistic topological GS of microscopic Hamiltonians on geometries involving open or fixed BC in some directions.

The total weight corresponding to each order of the expansion of \(H_{\text{local}}\) in terms of \(N\)-body operators are shown in Fig. 11(a,b) as a function of the order \(N\). Finite size effects are remarkably small and we believe the results for \(N_v = 8\) are converged. The data reveal clearly an exponential decay of the weight with the order \(N\). In other words, \(H_{\text{local}}\) contains primarily one- and two-body contributions (in addition to the normalization constant). This is the first part of the proof that \(H_{\text{local}}\) is indeed local. However, one still needs to go beyond the analysis and investigate further the \(r\)-dependence of the leading two-body contributions. In the next Subsection, we show that \(H_{\text{local}}\) of the \(\mathbb{Z}_2\) topological RVB is basically a short-range two-body Hamiltonian. In contrast, the RVB wave function on the square lattice exhibits a long-range two-body potential term.

4. Local boundary Hamiltonian: an effective one-dimensional t–J model

Next, we investigate the exact connection between the boundary Hamiltonian and the bulk properties of the system. We look for its explicit form, trying to make the connection with \(D = 3\) models, with \(\text{su}(2)\) symmetry corresponding to the \(1/2 \oplus 0\) representation.

The boundary Hamiltonian belongs to the \(1/2 \oplus 0\) representation of \(\text{su}(2)\) and its Hilbert space is the same as the one of a bosonic t–J model. Therefore, \(H_{\text{local}}\) is formally equivalent to a one-dimensional (1D) “t–J model” describing motion of (bosonic) “holes” (the “0”) in a spin fluctuating background (the “1” and “2”) in a spin fluctuating background (the “0”) in a spin fluctuating background (the “1” and “2”) in a spin fluctuating background (the “0”) in a spin fluctuating background (the “1” and “2”).

The form of the Hamiltonian components are dictated by the spin symmetry of the boundary Hamiltonian. We restrict here to the (dominant) 1- and 2-body terms of \(H_{\text{local}}\). The unique one-body (diagonal) term can be written as a chemical potential term \(H_2(r)\):

\[
\sum_i \hat{x}_i^i = \frac{3}{\sqrt{2}} \sum_i (n_i - 2/3) = \frac{3}{\sqrt{2}} H_2, \tag{12}
\]

where \(n_i = n_{i,0} + n_{i,1}\) counts the number of 0 or 1 on site \(i\). The diagonal 2-body density-density operators takes the form of a density-density (repulsive) interaction \(H_V\):

\[
\sum_i \hat{x}_i^i \hat{x}_{i+r}^i = \frac{9}{2} \sum_i (n_i - 2/3)(n_i-r - 2/3) = \frac{9}{2} H_V(r), \tag{13}
\]
Defining the pseudo-spin $S = \frac{1}{2} \sum_{s' \in \{0,1\}} \sigma_{ss'} |s\rangle \langle s'|$ involving a combination of $\hat{x}_1$, $\hat{x}_3 = \sqrt{3} |0\rangle \langle 1|$ and $\hat{x}_4 = \sqrt{3} |1\rangle \langle 0|$, and combining three 2-body terms (that appears in $H_1$ and $H_{local}$ with the same weights) we obtain an effective Heisenberg-like couplings $H_J(r)$:

$$
\sum_i \left( \hat{x}_1^r \hat{x}_1^{i+r} + \hat{x}_3^r \hat{x}_3^{i+r} + \hat{x}_4^r \hat{x}_4^{i+r} \right) = 6 \sum_i S_i \cdot S_{i+r} = 6 H_J(r).
$$

(14)

By symmetry, one also get (short-range) hopping terms $H_h(r)$ by combining:

$$
\sum_i \left( \hat{x}_1^r \hat{x}_5^{i+r} + \hat{x}_5^r \hat{x}_1^{i+r} + \hat{x}_6^r \hat{x}_6^{i+r} + \hat{x}_8^r \hat{x}_8^{i+r} \right)
$$

$$
= 3 \sum_i \left( b_{i,s}^r b_{i+1,s} + b_{i,s}^r b_{i+1,s}^r \right)
$$

$$
= 3 H_h(r),
$$

(15)

and Josephson couplings $H_J(\Delta)$ by combining:

$$
\sum_i \left( \hat{x}_6^r \hat{x}_6^{i+r} - \hat{x}_8^r \hat{x}_8^{i+r} + \hat{x}_7^r \hat{x}_7^{i+r} - \hat{x}_5^r \hat{x}_5^{i+r} \right)
$$

$$
= 3 \sum_i \left( b_{i,0} b_{i+1,1} - b_{i,1} b_{i+1,0} \right) + h.c.
$$

$$
= 3 H_J(\Delta),
$$

(16)

which describe fluctuations of (s-wave) short-range singlet pairs. The local Hamiltonian takes then the final form:

$$
H_{local} = c_0 N_v + \frac{3V_2}{\sqrt{2}} H_2 + \sum_r V_r H_V(r)
$$

$$
+ \sum_r t_r H_t(r) + \sum_r J_r H_J(r)
$$

$$
+ \sum_r \Delta_r H_{\Delta}(r) + H_{rest},
$$

(17)

where $H_{rest}$ contains all negligible $N > 3$ contributions. The new physical parameters are simply related to the amplitudes appearing in the expansion of $H_{local}$: $t_r = 3 d_{57}(r) = 3 d_{68}(r)$, $J_r = 6 d_{11}(r) = 6 d_{34}(r)$, $V_r = \frac{3}{2} d_{22}$ and $\Delta_r = 3 d_{33}(r) = 3 d_{37}(r)$.

For the $Z_2$ RVB liquid on the kagome lattice, as seen on Fig. 12(b), all weights $d_{\lambda \mu}^r(r)$ (and hence all the physical parameters $t_r$, $J_r$, $V_r$ and $\Delta_r$) decay exponentially fast with $r$ so that $H_{local}$ is a truly local operator. The dominant 2-body contribution to $H_{local}$ is the (negative) hopping term. The density-density interaction is attractive between nearest-neighbor sites ($V_1 < 0$) while it becomes repulsive (and very small) at longer distance ($V_r > 0$ for $r \geq 2$). Finally, we note that the small Heisenberg spin interaction is ferromagnetic at all distances ($J_r < 0$). For the critical RVB wave function on the square lattice, as seen on Fig. 12(a), all weights $d_{\lambda \mu}^r(r)$ also decay exponentially fast with $r$ except the (diagonal) density-density interaction $H_V(r)$ which remains long-range. These remarkable features are to be connected to the bulk correlations of the RVB wavefunctions: short-range (critical) bulk correlations translate into short-range (long-range) boundary Hamiltonians. We have therefore established a one-to-one correspondence between the long-range behavior of the bulk correlations and the range of the boundary Hamiltonian of RVB wavefunctions. This extends the previous findings to the case of topological order.

### C. Entanglement spectra and edge modes

We now move to the investigation of the full bipartite ES which is given by the spectrum of $H_{local}$. Our results are summarized in Figs. 13(a,b) and 14(a,b), for infinitely-long cylinders with kagome and square lattices. For convenience, the GS energy of $H_{local}$ (corresponding to the largest weight in the RDM) is subtracted from the spectra. The (excitation) ES are shown as a function of momentum around the cylinder and the eigenstates are labelled according to their spin-multiplet structure inherited from the $su(2)$ symmetry of the RVB state, although with the $1/2 \oplus 0$ representation. A careful finite size scaling (see Appendix B) suggests that the kagome (square) lattice cylinder ES is gapless (gapped) in the limit $N_v \to \infty$. Since these features are opposite to what is expected for the energy excitation spectra of the corresponding bulk systems (according to their long wavelength properties), we deduce that the ES characterizes specifically the nature of the L and R edge modes (Fig. 3). Note that for given choice of BC, the actual ES is the spectrum of a projected $\hat{H}_1$ Hamiltonian and, hence, is a subset of the full ES. For example, in a kagome lattice (square lattice) cylinder with OBC, a common set-up in numerical simulations, only (a sub-set of) the integer spin eigenstates are obtained.

### D. Topological entropy

It is of great interest to investigate the entanglement entropy which can give access to the quantum dimension $D$ and hence provides clear fingerprints of topological order. We recall that we consider here the RVB wavefunction for which the same sign enters in the linear superposition of the dimer coverings (defined, on the kagome lattice, as $\Psi_{RVB}$ with no vison line, i.e. no $Z_2$ flux through the cylinder). We investigate infinite cylinders and study the behavior of the EE as a function of the perimeter. From specific choices of the cylinder boundary conditions we can select specific conserved sectors on the edge (e.g. OBC for the kagome cylinder selects the even sector, etc...). The EE is given by the Von Neumann entropy $S_{VN} = -Tr\{\sigma_3^2 \ln \sigma_3^2\}$.

As shown in Fig. 15(a), the EE of the square lattice cylinder with OBC ($\Delta = 0$ sector) shows strong
entanglement entropy

$2044$

Fig. 15(b). The existence of a finite $H_1$ as a direct consequence of the particular structure of $\Psi_{RVB}$ can be seen as a direct consequence of the particular structure of $H_1$ according to the following argument: The EE is given (crudely) by $-\ln \mathcal{N}$ where $\mathcal{N}$ is the number of eigenstates of $H_1$ below a fixed energy scale of order 1. For fixed cylinder boundaries, the support of $H_1 = H_{local}P \ (P = P_{even} \text{ or } P = P_{odd})$ contains $\mathcal{N} \approx \frac{1}{\pi} 3^{N_v}$ states and $S_0 = -\ln 2$, as expected for a topological $Z_2$ spin liquid with quantum dimension $D = 2$. Note that $\Psi_{RVB}^{+even}$ and $\Psi_{RVB}^{odd}$ can be seen as “minimally entangled states” naturally produced by the DMRG algorithm from amongst the quasi-degenerate ground states of the $Z_2$ topological phase. Linear combination of them (or e.g. of $\Psi_{RVB}^{+even}$ and $\Psi_{RVB}^{even}$) should give a larger topological entropy.

Summing over all sectors amounts to taking $H_1 = H_{local}$ so that all eigenstates of the ES contribute and $S_{VN} \propto N_v \ (\text{as can be shown rigorously})$ as seen in Fig. 15(b). For the square lattice, severe constraints leads to an extensive number (i.e. proportional to the perimeter $N_v$) of disconnected sectors on the edge of dimension $\mathcal{N} \approx \frac{1}{\pi} 3^{N_v}$, therefore introducing negative logarithmic corrections $-\ln N_v$ to the EE for any boundary conditions (see e.g. data for OBC on Fig. 15(a)). The long-range diagonal interaction in $H_{local}$ (Fig. 12(a)) may also be responsible for deviations from the area law, even when considering all sectors.

V. DISCUSSION AND OUTLOOK

Introducing PEPS representations and using Tensor Networks techniques, we have examined topological and entanglement properties associated to gapped and gapless RVB states using cylindrical geometries with arbitrary boundary conditions. The formalism allows to take the limit of infinite cylinders. Using the simple topological structure of the space of dimer coverings on the
kagome lattice, we construct four quasi-degenerate (for a generic quantum HAF) orthogonal RVB states and obtain the finite size scalings of the energy splittings amongst them (topological gaps) which could be compared to numerical simulations. Incidentally, our results identify two very different energy splittings decaying with two clearly different length scales. The largest energy scale corresponds to inserting a (horizontal) vison line (or a $\mathbb{Z}_2$ flux in the cylinder). The second energy scale corresponds to pinning a site with a strong field on either end of the cylinder, which would be the same as moving a site from one end to the other. Although it has been suggested that the DMRG algorithm (naturally) selects a minimally entangled state\textsuperscript{[11]} it is still not clear how to reconcile the fact that the finite size corrections of the groundstate energy are very small\textsuperscript{[11]} while our RVB computation predicts clear finite-size effects for the states with a definite $\mathbb{Z}_2$ flux. On the other hand, we find that the energy averaged over the four (minimally entangled) RVB states shows very small finite size effects.

In addition, we show that boundary Hamiltonians can be written as $H_{local}(\mathcal{P}) + \beta_{\text{topo}}(\text{Id} - \mathcal{P})$, $\beta_{\text{topo}} \rightarrow \infty$. In particular, we have established the existence of a projector $\mathcal{P}$ (which intrinsically depends on the boundary conditions) onto a restricted subspace at the edge (as for Kitaev toric code\textsuperscript{[3,17]}), a consequence of the disconnected topological sectors in the space of dimer coverings of the lattice. We argue that the non-local character of the resulting boundary Hamiltonian is the fingerprint of topology. The ES is a subset (associated to $\mathcal{P}$) of the spectrum of the emerging local Hamiltonian $H_{\text{local}}$ acting on the unrestricted edge space. In contrast to the toric code for which $H_{\text{local}}$ is trivial, here $H_{\text{local}}$ takes the form of a short-range (bosonic) t–J model (including a long-range diagonal interaction for the critical RVB state). We argue that the topological features (e.g. finite size scaling of topological gaps) and entanglement properties (e.g. structure of boundary Hamiltonians) of the NN RVB wave functions are characteristic of topological phases. We propose to use these features to detect topological order in microscopic models\textsuperscript{[9,12]}

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**APPENDIX A: TAKING THE LIMIT OF THE INFINITE CYLINDER**

The (excitation) ES are shown in Fig.\textsuperscript{[16]} as a function of momentum around the cylinder and the eigenstates are labelled according to their spin-multiplet structure inherited from the su(2) symmetry of the RVB state, although with the $1/2 \oplus 0$ representation. From the data shown in Fig.\textsuperscript{[16]} we see that, for a fixed perimeter, the ES converge rapidly when increasing the length towards the infinite-cylinder limit ($N_h = \infty$). The latter is reached as soon as $N_h > N_v$ ($N_h \gg N_v$) for the Kagome (square) lattice: in practice, the RDM for $N_h = 50$ is fully converged. This is clear from the finite size scaling analysis of some of the low-energy excitations of the ES shown in Fig.\textsuperscript{[17]}
FIG. 17: (Color online) Finite size scaling of the lowest energy levels marked by arrows in the ES of Fig. 16. Excellent convergence is found when $N_h \to \infty$ (at constant $N_v = 8$). Note the alternating behavior according to the parity of $N_h/2$ for the square lattice, in contrast to the kagome lattice showing a straight exponential convergence.

APPENDIX B: FINITE SIZE SCALING OF THE ENTANGLEMENT SPECTRUM VS CYLINDER PERIMETER

After taking the limit of the infinite cylinder ($N_h \to \infty$), we investigate the dependence of the ES as a function of the cylinder perimeter $N_v$. Note that when taking the $N_h \to \infty$ limit, one still has to specify the choice of the $B_L$ and $B_R$ boundaries that uniquely determine the projector (even or odd) involved at the edges along the cut. Here we only consider OBC which select the integer spin sector of the boundary Hamiltonian. Our results are summarized in Fig. 18 for infinitely-long cylinders with kagome and square lattices. A careful analysis of these spectra and some of their low-energy excitations (see Fig. 19(a)) as a function of cylinder perimeter suggests that the square (kagome) lattice cylinder ES is gapped (gapless) in the limit $N_v \to \infty$. Since these features are opposite to what is expected in the corresponding bulk systems, we deduce that the ES characterizes specifically the nature of the edges (L and R in Fig. 1(a-d)).

APPENDIX C: EXPANSION IN TERMS OF MANY-BODY OPERATORS

Any operator $\mathcal{O}_{\text{edge}}$ like projectors $\mathcal{P}$ or boundary Hamiltonians acting on the edge can be expanded in terms of $3^{2N_v}$ orthogonal (real) operators $\hat{X}_\alpha$,

$$\mathcal{O}_{\text{edge}} = c_0 N_v + \sum_\alpha A_\alpha \hat{X}_\alpha .$$

(18)

The scalar product in the operator basis is defined as $\langle \hat{u} | \hat{v} \rangle$, where $\langle \cdots \rangle = \frac{1}{2} \text{Tr}(\cdots)$ and the trace is a priori

FIG. 18: (Color online) ES of infinite cylinders with increasing perimeter $N_v$. OBC are used for $B_L$ and $B_R$. Kagome (top) and square (bottom) lattices.

FIG. 19: (Color online) Finite size scaling of low excitation energies of infinite cylinders vs inverse perimeter, suggesting a vanishing (finite) gap in the thermodynamic limit for the kagome (square) lattice. For the square lattice, the average (divided by 3) between the lowest $K = \pi$ singlet and triplet excitations is shown.
ori performed over the full basis of $Z = 3^{N_v}$ states. For convenience, the constant term $c_0 = \frac{1}{2} \langle \mathcal{O}_{\text{edge}} \rangle$ has been separated so that we can assume all other operators satisfy $\text{Tr} \tilde{X}_\alpha = 0$. Simple algebra shows that the coefficients can be obtained by taking the trace of the corresponding operators with $\mathcal{O}_{\text{edge}}$ as,

$$A_\alpha = \langle \tilde{X}_\alpha \mathcal{O}_{\text{edge}} \rangle / \langle \tilde{X}_\alpha \tilde{X}_\alpha^\dagger \rangle,$$

where the trace in the numerator involves, in fact, the sum over the projected subspace. We also obtain some "sum-rule":

$$\langle \mathcal{O}_{\text{edge}}^2 \rangle = (c_0 N_v)^2 + \sum_\alpha A_\alpha^2 \langle \tilde{X}_\alpha \tilde{X}_\alpha^\dagger \rangle,$$

which enable to compute the weight associated to each operator.

To go further and expand $\mathcal{O}_{\text{edge}}$ in the full operator basis, it is convenient to use a local basis of 9 (normalized) operators $\{\hat{x}_0, \cdots, \hat{x}_8\}$ which act on the local site configuration, $\{\{0\}, \{1\}, \{2\}\}$, e.g. $\hat{x}_0 = 0$, $\hat{x}_1 = \sqrt{\frac{3}{2}} (|0\rangle \langle 0| - |1\rangle \langle 1|)$, and $\hat{x}_2 = \frac{1}{\sqrt{2}} (|0\rangle \langle 0| + |1\rangle \langle 1| - 2|2\rangle \langle 2|)$, for the diagonal matrices, complemented by $\hat{x}_3 = \hat{x}_4 = \sqrt{3}|1\rangle \langle 1|$ acting as "spin" operators, and $\hat{x}_5 = \hat{x}_7 = \sqrt{3}|2\rangle \langle 2|$ and $\hat{x}_6 = \hat{x}_8 = \sqrt{3}|2\rangle \langle 1|$ acting as annihilation and creation (hardcore) bosonic operators. These operators satisfy $\text{Tr}(\hat{x}_3) = 0$ (for $\lambda \neq 0$) and $\text{Tr}(\hat{x}_3 \hat{x}_3) = 3$, where "tr" is the trace over the local degrees of freedom (of some site $i$). From now on, we extend the action of these local operators to the whole edge, assuming a trivial (implicit) action on the $N_v - 1$ unspecified sites, i.e. $\hat{x}_\lambda \equiv \hat{x}_\lambda \otimes 1^{\otimes (N_v-1)}$, so that $\text{Tr}(\hat{x}_\lambda (\hat{x}_\lambda)^\dagger) = 3^{N_v}$ and $\langle \hat{x}_\lambda (\hat{x}_\lambda)^\dagger \rangle = 1$. Using the local basis of operators, one can then uniquely expand any edge operator like $H_1$ in terms of N-body operators as,

$$\mathcal{O}_{\text{edge}} = c_0 N_v + \sum_\lambda c_\lambda \sum_i \hat{x}_\lambda \sum_{\lambda,\mu, r} d_{\lambda \mu}(r) \sum_i \hat{x}_\lambda \hat{x}_\mu^{i+r},$$

$$+ \sum_{\lambda,\mu, r, r'} e_{\lambda \mu}(r, r') \sum_i \hat{x}_\lambda \hat{x}_\mu^{i+r} \hat{x}_\nu^{i+r'} + \cdots,$$

where each group of terms involves products of $N = 1, 2, \cdots, N_v$ on-site operators $\hat{x}_\lambda$, $i$ labeling the sites. Here the sums do not contain the identity, the sums over distances are restricted to non-equivalent relative distances. $\sum^m$ means that only translations giving distinct sets of sites are performed (no multiple counting). Hence, the N-body translationally invariant operators $\tilde{X}_\alpha$ in [21], where $\alpha$ combines all the labels of the coefficients of the expansion (e.g. $\alpha = (\lambda, \mu, r)$ and $\tilde{X}_\alpha = \sum_{\lambda, \mu} \hat{x}_\lambda \hat{x}_\mu^{i+r}$), are normalized as $\langle \tilde{X}_{\alpha} \tilde{X}_{\alpha}^\dagger \rangle = N_v/g_\alpha$, where $g_\alpha$ are “multiplicity” factors that count the number of times the operator maps onto itself under all $N_v$ translations. The (real) coefficients in [21] are obtained by taking the trace (in operator space) of the corresponding operators with the operator $\mathcal{O}_{\text{edge}}$,

$$c_\lambda = \frac{1}{N_v} \langle \sum_{i=1}^{N_v} \hat{x}_\lambda \rangle \mathcal{O}_{\text{edge}},$$

$$d_{\lambda \mu}(r) = \frac{1}{N_v} \langle \sum_{i=1}^{N_v} \hat{x}_\lambda \hat{x}_\mu^{i+r} \rangle \mathcal{O}_{\text{edge}},$$

$$e_{\lambda \mu}(r, r') = \frac{1}{N_v} \langle \sum_{i=1}^{N_v} \hat{x}_\lambda \hat{x}_\mu^{i+r} \hat{x}_\nu^{i+r'} \rangle \mathcal{O}_{\text{edge}},$$

where one can make advantage of translation symmetry to compute the r.h.s of these equations. The sum-rule for the weights takes then the form:

$$\frac{1}{N_v} \langle \mathcal{O}_{\text{edge}}^2 \rangle = c_0^2 N_v + \sum_\lambda \sum_{\lambda, \mu} \sum_r d_{\lambda \mu}(r)^2 + \cdots,$$

where the (second) sums are restricted to non-equivalent sets of distances and the multiplicity factors only depend on the latter.

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