Hexagon-singlet solid ansatz for the spin-1 kagome antiferromagnet

Wei Li\textsuperscript{1,2}, Andreas Weichselbaum\textsuperscript{1}, Jan von Delft\textsuperscript{1}, and Hong-Hao Tu\textsuperscript{3,*}

1. Physics Department, Arnold Sommerfeld Center for Theoretical Physics, and Center for NanoScience, Ludwig-Maximilians-Universität, 80333 Munich, Germany
2. Department of Physics, Beihang University, Beijing 100191, China
3. Max-Planck-Institut für Quantenoptik, Hans-Kopfermann-Str. 1, D-85748 Garching, Germany

(Dated: January 5, 2015)

We perform a systematic investigation on the hexagon-singlet solid (HSS) states, which are a class of spin liquid candidates for the spin-1 kagome antiferromagnet. With the Schwinger boson representation, we show that all HSS states have exponentially decaying correlations and can be interpreted as a (special) subset of the resonating Affleck-Kennedy-Lieb-Tasaki (AKLT) loop states. We provide a compact tensor network representation of the HSS states, with which we are able to calculate physical observables efficiently. We find that the HSS states have vanishing topological entanglement entropy, suggesting the absence of intrinsic topological order. We also employ the HSS states to perform a variational study of the spin-1 kagome Heisenberg antiferromagnetic model. When we use a restricted HSS ansatz preserving lattice symmetry, the best variational energy per site is found to be $e_0 = -1.3600$. In contrast, when allowing lattice symmetry breaking, we find a trimerized simplex valence bond crystal with a lower energy, $e_0 = -1.3871$.

PACS numbers: 75.10.Kt, 75.10.Jm

I. INTRODUCTION

Frustrated antiferromagnets on the kagome lattice have attracted great research interest recently. The ground state of a spin-1/2 kagome Heisenberg antiferromagnet (KHAF) has been disclosed to be a disordered state without any spontaneous symmetry breaking, i.e., a quantum spin liquid.\textsuperscript{1,2} However, less is known for the higher-spin KHAF models ($S > 1/2$). Numerical studies on the KHAF models with spin magnitude up to $S = 3$ showed that\textsuperscript{3} while long-range magnetic order appears for $S \geq 3/2$, the ground states for the $S = 1/2$ and $S = 1$ KHAF models remain non-magnetic. Experimentally, a number of spin-1 kagome compounds have been synthesized and analyzed, e.g., m-MPYNN-BF\textsubscript{4}\textsuperscript{4–7} and Ni\textsubscript{3}V\textsubscript{2}O\textsubscript{6}\textsuperscript{8}. The former has been found to be non-magnetic even at very low temperatures (30mK),\textsuperscript{5} and a spin gap has also been observed.\textsuperscript{9} This thus raises an interesting question: do spin-1 kagome antiferromagnets also support an intriguing spin liquid ground state?

Various spin liquid proposals have been put forward for the spin-1 frustrated antiferromagnet.\textsuperscript{9–17} For the spin-1 KHAF model, Hida proposed a hexagon-singlet solid (HSS) state as a candidate ground state.\textsuperscript{15} This state is constructed by projecting two virtual spin-1/2 particles around each vertex of the kagome lattice into physical spin-1 degrees of freedom. The name HSS refers to the fact that the six virtual spin-1/2’s in a hexagon of the kagome lattice form an entangled singlet [see Fig. 1(a)]. An alternative candidate is the spin-1 resonating Affleck-Kennedy-Lieb-Tasaki loop (RAL) state,\textsuperscript{16,17} which is an equal weight superposition of all possible AKLT loops\textsuperscript{18} on the kagome lattice and constitutes a $Z_2$ spin liquid. Compared to the HSS ansatz, the RAL has a lower variational energy (for the spin-1 KHAF model) on small clusters; while in the thermodynamic limit, the energy per site is clearly higher, $e_0 \approx -1.27.17$

Very recently, several extensive numerical studies have been devoted to the spin-1 KHAF model, exploring its ground state properties. Changlani and Läuchli (CL)\textsuperscript{19} employed the density matrix renormalization group (DMRG)\textsuperscript{20} to simulate the model with cylindrical geometries; at the same time, Liu et al.\textsuperscript{21} adopted tensor network methods\textsuperscript{22,23} to explore the same model on an infinitely large kagome lattice and also on infinitely long cylinders with various widths. These independent calculations, as well as another related tensor network simulation by Picot and Poilblanc,\textsuperscript{24} concluded that the ground state of the spin-1 KHAF is non-magnetic, but that it breaks lattice inversion symmetry and possesses a simplex valence-bond crystal (SVBC) order that favors trimerization,\textsuperscript{25} in that the energies (per triangle) differ between two neighboring triangles. The energy per site of

![FIG. 1. (Color online) Schematic plots of (a) the spin-1 HSS ansatz on the kagome lattice and (b) the spin-3/2 HSS ansatz on the honeycomb lattices. Six spin-1/2 virtual particles (red dots) form a spin singlet in each hexagon. In (a) the blue ovals denote projectors mapping two virtual spin-1/2’s to a physical spin-1, while in (b) the circles denote projectors mapping three spin-1/2’s onto a spin-3/2 space.](image-url)
this SVBC state was determined as $e_0 \approx -1.41$, in both DMRG and tensor network calculations.\cite{19,21,24}

Even more recently, Nishimoto and Nakamura (NN14)\cite{26} came to a different conclusion: Based on DMRG calculations for clusters of various types of shapes and boundary conditions, they argued that the ground state is a HSS state and not the SVBC state advocated in Refs. 19, 21, and 24. However, NN14 were not able to directly access states with HSS or SVBC structure; instead they sought to access them indirectly, using purposefully-designed boundary conditions that favor either HSS or SVBC structure. They then estimated the bulk values of $e_0$ by finite-size extrapolations to the thermodynamic limit. They reported $e_0 = -1.391(2)$ from SVBC-favoring clusters, $e_0 = -1.40988$ and $e_0 = -1.409(5)$ from clusters with cylindrical or periodic boundary conditions, respectively, and $e_0 = -1.41095$ from HSS-favoring clusters, thus concluding that HSS states win.

In our opinion, NN14’s strategy is intrinsically flawed on very general grounds: in the thermodynamic limit boundary effects should vanish, thus a tool (here ground-state DMRG on finite-sized clusters) that relies on boundary effects to distinguish two types of states (here HSS and SVBC), can not reliably estimate the difference in bulk $e_0$ values for these two types of states. If different boundary choices lead to different finite-size extrapolated $e_0$ values, it just means that the clusters are not yet large enough to reliably capture the true bulk $e_0$ value of the true ground state, whatever it is. (More detailed comments on NN14’s work are presented in Appendix C.)

To reliably access the bulk properties of HSS-type states for the spin-1 KHAF, tools are needed that directly implement HSS structure in the variational candidate ground state, without relying on boundary effects. In this work, we devise two such tools, one analytical, the other numerical, and use them to perform a systematic investigation of HSS states.

Our analytical tool is a formulation of the HSS ansatz in terms of SU(2) Schwinger bosons. In the Schwinger boson picture, we are able to prove that the HSS states have exponentially decaying correlation functions and thus describe gapped spin liquids. Furthermore, we also reveal that the HSS states have a hidden resonating AKLT-loop picture. However, we show that they have zero topological entanglement entropy,\cite{27,28} concluding that they are topologically trivial and thus do not belong to the same phase as the RAL state.

Our numerical tool is a compact tensor network representation of the HSS ansatz, with which we perform accurate tensor-network-based simulations. The weights of different hexagon-singlet configurations within a hexagon are treated as variational parameters to seek the lowest possible variational energy for the spin-1 KHAF model. It is found to be as low as $e_0 \approx -1.3600$, which is significantly higher than the reported ground state energy ($-1.41$) of the SVBC state.\cite{29} Moreover, we mimic a single step of imaginary-time evolution (on one of the two kinds of triangles), and thus add one additional parameter $\tau$. This gives a clear gain in energy ($e_0 = -1.3871$), for a state breaking the lattice symmetry between two neighboring triangles. This variational calculation indicates that the HSS state, which does not break any lattice symmetry, may not be energetically favorable for the spin-1 KHAF model.

The paper is organized as follows: in Sec. II we introduce the HSS ansatz in terms of the Schwinger boson representation. In Secs. III and IV we show a compact tensor network representation for the HSS states, and use it to calculate the physical quantities, including the variational energy for the spin-1 KHAF model and various correlation functions. Sec. V is devoted to the summary and discussions.

II. HEXAGON-SINGLET SOLID ANSATZ

In this section, we briefly review the hexagon-singlet solid ansatz.\cite{15} We start in Sec. II A by introducing the construction of the HSS ansatz in terms of Schwinger bosons, and then provide a physical picture of resonating AKLT loops for these states in Sec. II B.

A. Schwinger boson formulation

The hexagon-singlet solid ansatz for the spin-1 kagome antiferromagnet bears similarity to the construction of the 1D spin-1 AKLT state.\cite{18} For each physical spin-1 site, one associates two virtual spin-1/2 particles. Since the kagome lattice can be viewed as a lattice with corner-sharing hexagons, each hexagon contains six virtual spin-1/2 particles [see Fig. 1(a)]. On each hexagon, the six virtual spin-1/2 particles are combined into a SU(2) spin singlet. The final step is to recover a physical spin-1 wave function by symmetrizing the two virtual spin-1/2’s in the same lattice site into the spin-1 subspace. In contrast to ordinary AKLT states, there exist inequivalent singlet configurations per hexagon (depicted schematically in Fig. 2). Therefore, this construction provides a class of trial wave functions for the spin-1 kagome antiferromagnet.

Formally, it is convenient to formulate the HSS ansatz in terms of Schwinger bosons.\cite{29} In Schwinger boson language, the SU(2) spin operators for each site $i$ are represented as

$$S^a_i = \frac{1}{2} \sum_{\alpha \beta = \uparrow, \downarrow} b_i^\dagger \sigma^a_{\alpha \beta} b_i \quad (a = x, y, z),$$  \hspace{1cm} (1)

where $\sigma^a$ are Pauli matrices. On each site, a boson number constraint, $\sum_{\alpha = \uparrow, \downarrow} b_i^\dagger b_i = 2$, has to be imposed to guarantee the physical spin-1 Hilbert space. Here the bosonic statistics takes care of the symmetrization of two virtual spin-1/2’s into a spin-1 and, in terms of Schwinger
Schwinger bosons within the same hexagon and can be thus contains some configurations which are prohibited in the hexagonal inversion symmetry: \( |N\rangle \) with longer-range valence bonds in the hexagon. The arrow originates from the first to the second spins in a singlet, which is anti-symmetric towards permutation of the two constituting spin-1/2’s. Notice that these three hexagon singlets are all eigenstates of hexagon-inversion symmetry: \( |D_\pm\rangle \) and \( |N\rangle \) are odd (eigenvalue -1), while \( |D_+\rangle \) is even (+1).

FIG. 2. (Color online) Graphical representation of the hexagon singlets. (a) All dimers are between nearest neighbors, like a resonating benzene ring, \( |D_\pm\rangle \) is with + (−) sign convention in the superposition. (b) An allowed dimer pattern \( |N\rangle \) with longer-range valence bonds in the hexagon. The arrow originates from the first to the second spins in a singlet, which is anti-symmetric towards permutation of the two constituting spin-1/2’s. Notice that these three hexagon singlets are all eigenstates of hexagon-inversion symmetry: \( |D_\pm\rangle \) and \( |N\rangle \) are odd (eigenvalue -1), while \( |D_+\rangle \) is even (+1).

FIG. 3. (Color online) (a) Typical resonating AKLT loop configuration arising from the benzene construction. The red arrows denote (oriented) valence bonds formed by virtual spin-1/2 particles. When cutting the loop configurations either horizontally or vertically (denoted by dashed lines), it is only possible to intersect an even number of valence bonds. As a comparison, we also show a RAL configuration in (b). The RAL state has four virtual particles 0 ⊕ 1/2 on each vertex (see green circles in (b), and more details in Ref. 17), and is a fully packed equal weight superposition of all possible loops, thus contains some configurations which are prohibited in the HSS state.

bosons, the three spin-1 states are represented as

\[
|1\rangle = \frac{(b_1^\dagger)^2}{\sqrt{2}}|\text{vac}\rangle, \quad |0\rangle = b_1^\dagger b_4^\dagger|\text{vac}\rangle, \quad |-1\rangle = \frac{(b_4^\dagger)^2}{\sqrt{2}}|\text{vac}\rangle,
\]

where \( |\text{vac}\rangle \) is the vacuum of the Schwinger bosons.

By using the Schwinger bosons, the HSS state is written as

\[
|\Psi\rangle = \prod_{\circ} \mathcal{P}_{\circ}^+ |\text{vac}\rangle,
\]

generally written as

\[
\mathcal{P}_{\circ}^+ = \sum_{\alpha_1,\alpha_2,\ldots\alpha_6 = \uparrow,\downarrow} T_{\alpha_1,\alpha_2,\ldots,\alpha_6}^\dagger b_{\alpha_1,\alpha_2}^\dagger \cdots b_{\alpha_5,\alpha_6}^\dagger.
\]

As there are several inequivalent ways of combining six spin-1/2’s into a singlet, \( T_{\alpha_1,\alpha_2,\ldots,\alpha_6} \) in (4) reflects this freedom of choice. For the wave function in (3), there are exactly two Schwinger bosons for every site and no extra projector is needed to remove unphysical configurations.

To gain further insight into the HSS ansatz, we exploit the fact that the hexagonal singlet \( \mathcal{P}_{\circ}^+ |\text{vac}\rangle \) in (3) can always be decomposed into a superposition of valence-bond singlets (over-complete bases) as

\[
\mathcal{P}_{\circ}^+ = \sum_{\{ij,kl, mn\}} w_{ij}w_{kl}w_{mn} C_{\circ,ij}^+ C_{\circ,kl}^+ C_{\circ,mn}^+ \cdot
\]

where \( \{ij,kl, mn\} \) denotes all allowed singlet-pair configurations (e.g., \{12,34,56\}, \{13,25,46\}), the valence-bond singlet creation operator \( C \) is defined by

\[
C_{ij}^+ = b_{ij}^\dagger b_{ij}^\dagger - b_{ij}^\dagger b_{ij}^\dagger,
\]

and \( w_{ij} \) are coefficients controlling the weights of the valence bonds. Generically, \( w_{ij} \) can be viewed as a set of free parameters. Comparing to (4), an obvious advantage of the parametrization (5) is that the \( C_{ij} \) lattice symmetry can be easily imposed in these ansatz. For instance, one may consider a simple choice with \( w_{ij} \) only depending on the distance between sites \( i \) and \( j \).

With the help of the Schwinger boson representation, we are able to show rigorously that the HSS class of states have \textit{exponentially} decaying correlation functions, indicating that they describe a class of \textit{gapped} spin liquids. The technical details on the proof of this statement are given in Appendix A. In short, the proof utilizes the spin-coherent state representation of the Schwinger boson states to write the norm of the HSS ansatz \( |\Psi\langle \Psi|\rangle \) as the partition function of a two dimensional (2D) classical statistical model describing interacting O(3) vectors on the kagome lattice. Additionally, the two-point correlation functions, say, the spin-spin correlation \( \langle \Psi| S_i \cdot S_j |\Psi\rangle \), can be expressed as the correlation function between O(3) vectors in the 2D statistical model. As the statistical model is at \textit{finite} temperature and has only short-range interactions, long-range order spontaneously breaking O(3) symmetry is not allowed, according to the Mermin-Wagner theorem, and the correlations between O(3) vectors (equivalently, spin-spin correlations in the HSS ansatz) decay exponentially. This is a direct generalization of the results in Ref. 31 proving that 2D AKLT states have exponentially decaying correlations.

In addition, the HSS construction is not restricted to the kagome lattice, but applies as well to any other lattices possessing hexagon motifs. For example, we show a spin-3/2 HSS ansatz on the honeycomb lattice in Fig. 1(b), where the three spin-1/2 virtual particles surrounding a vertex are symmetrized to constitute the physical
spin-3/2 degree of freedom. It is not difficult to see that
our proof on the gapped spin liquid nature of the kagome
HSS states (in Appendix A) also applies to all such HSS
wave functions in 2D, including the spin-3/2 honeycomb
HSS state in Fig. 1(b).

B. Resonating Affleck-Kennedy-Lieb-Tasaki loop
picture

Let us now introduce a simple example belonging to
the HSS class on the kagome lattice, which we call the
Benzene Ring State (BRS). Based on the BRS example,
we uncover a resonating AKLT loop picture for the HSS
ansatz.

The BRS states are defined by restricting hexagonal
singlets in (5) to short-range dimers between neighboring
sites. Two (inequivalent) such choices for \( w_{ij} \) in (5) are
given by

\[
\begin{align*}
  w_{12} &= w_{34} = w_{56} = 1, \\
  w_{23} &= w_{45} = w_{61} = 1,
\end{align*}
\]

and

\[
\begin{align*}
  w_{12} &= w_{34} = w_{56} = 1, \\
  w_{23} &= w_{45} = w_{61} = -1,
\end{align*}
\]

respectively. The graphical representations for the above
two hexagonal singlet choices, resembling resonating benzene
rings, are shown in Fig. 2(a).

When building the wave function (3) using (7) or (8),
exanding the product \( \prod_{ij} \) in (3) leads to a number of nearest-neighbor
valence-bond configurations. A typical configuration is shown in Fig. 3(a). An interesting ob-
ervation is that this configuration can be viewed as the
covering of spin-1 AKLT loops on the kagome lattice.
This is due to the fact that each site shares two spin-1/2
valence bonds (forming spin-1 physical sites), that is to
say, the valence bonds have to enter and leave every site,
which inevitably form fully packed loop structure. Based
on this observation, we conclude that the BRS state can
be viewed as an equal weight superposition of resonating
AKLT loops.

What about the HSS ansatz with longer-range valence
bonds within each hexagon? It is not difficult to see that,
when expanding the product \( \prod_{ij} \) in (5), the AKLT
loop structure in each configuration is still preserved,
though the loops can connect sites beyond nearest neigh-
bors (NN). Then, the role of the weights \( w_{ij} \) in (5) is to
control the loop tension. This shows that all HSS ansatz
(3) belong to the broader family of resonating AKLT loop
states.

Coming back to the BRS states formed by an equal
weight superposition of AKLT loops, it is rather interesting
to make a comparison between them with the RAL
state considered in Ref. 17. In the latter, the RAL state
is an equal weight superposition of all possible AKLT
loops [see Fig. 3(b)] and has \( \mathbb{Z}_2 \) topological order on
the kagome lattice. However, there is an additional con-
straint in the BRS due to the benzene ring construction
which requires that no loop can be formed by two suc-
cessive valence bonds within the same hexagon. This,
in further, means that the allowed loop configurations in
the BRS states are strictly less than those of the RAL
state. Actually, this leads to an important observation
which reveals the difference between them: When the
RAL state is defined on a torus, cutting the torus in ei-
ther horizontal or vertical direction always intersects even
number of valence bonds [see Fig. 3(a)]. However, this is
not the case for the BRS state, where such a cut can in-
tersect both even and odd number of valence bonds [Fig.
3(b)], which enables the existence of different topological
sectors.17 While the existence of such “parity” sectors is
essential for \( \mathbb{Z}_2 \) topological order, this already gives a hint
that there is no topological order (at least not \( \mathbb{Z}_2 \) type) in
the BRS state and they are distinct from the RAL
state, even though they share very similar loop structure
with the latter. In Sec. III, we will provide numerical
evidence that the HSS ansatz, including the BRS state,
has vanishing topological entanglement entropy and thus
no topological order.

III. TENSOR NETWORK REPRESENTATION
AND SIMULATION OF THE
HEXAGON-SINGLET SOLID STATES

In this section, we provide a compact tensor network
representation for the HSS states (see Fig. 4) and cal-
culate the physical quantities by using tensor-network-
based simulations. In Sec. III A we start with analyzing
the BRS, a special HSS state introduced in Sec. II B. In
Sec. III B we move on to the study of another special
HSS state introduced in Ref. 15.

According to Sec. II, the HSS state (3), instead of its

\[
\begin{align*}
  M &= \left( \begin{array}{ccc}
              1 & 0 & 0 \\
              0 & 1 & 0 \\
              0 & 0 & 1
            \end{array} \right), \\
  R &= \left( \begin{array}{ccc}
              1 & 0 & 0 \\
              0 & 1 & 0 \\
              0 & 0 & 1
            \end{array} \right)
\end{align*}
\]
particles at site six virtual spin-1/2 particles, and the hexagon. Where \( σ \) denotes the hexagon singlet formed by six virtual spin-1/2 particles, and \( P_i \) projects two virtual particles at site \( i \) onto the triplet subspace

\[
|Ψ⟩ = \bigotimes_i P_i \prod_0 |ψ⟩,
\]

and

\[
|ψ⟩ = \sum_{\{σ \in O\}} T_{σ_1,σ_2,σ_3,σ_4,σ_5,σ_6}|σ_1,σ_2,σ_3,σ_4,σ_5,σ_6⟩,
\]

where \( σ_i \) is a virtual spin-1/2 located at site \( i \) belonging to the hexagon. \( |ψ⟩ \) denotes the hexagon singlet formed by six virtual spin-1/2 particles, and \( P_i \) projects two virtual particles at site \( i \) onto the triplet subspace

\[
P = \sum_{σ_1,σ_2} \sum_m C_{mσ_1,σ_2}|m⟩⟨σ_1,σ_2|,
\]

where \( m \in \{±1,0\} \) denotes the physical, \( σ_1,σ_2 \in \{±1/2\} \) the virtual space, and \( C_{mσ_1,σ_2} \) is the Clebsch-Gordan coefficient symmetrizing two spin-1/2’s into a physical spin-1, with nonvanishing elements as \( C_{1/2,1/2} = C_{-1/2,-1/2} = 1 \) and \( C_{0,1/2,1/2} = C_{0,1/2,-1/2} = 1/\sqrt{2} \). In Fig. 4(a), the HSS structure is depicted: the sixth-order tensor \( T \) is represented by the ring within each hexagon and the dashed oval on each site indicates the projector \( P \) [Fig. 4(b)].

However, from a numerical point of view, this representation is not practical for calculations, owing to the high coordination number \( z = 6 \) of the \( T \)-tensors. To overcome this difficulty, we here introduce a scheme shown in Figs. 4(d,e): two neighboring spin-1/2 particles (in the hexagon) are fused into a composite virtual particle \( 0 \otimes 1 \) of dimension four, and the coordination number of the hexagon tensor is lowered down to \( z = 3 \). After this transformation, we get a tetragonal tensor network consisting of \( M \) and \( R \) tensors, which can be more easily treated with the tensor network techniques. Notice that one has the freedom to block the virtual particles in two different ways [odd-even, or even-odd pairs, as in Fig. 2(a)]. Although the tensor network representation would be different (the \( M \) tensor would be associated to different triangles), they represent essentially the same state.

A. The Benzene Ring State

We start with the BRS depicted in Fig. 3, which only contains NN valence bonds. There are two hexagon-singlet configurations in the BRS for two sign choices, i.e., \( |ψ⟩ \) in Eq. (9) can be chosen as \( |D_±⟩ \) in Fig. 2(a), and we thus construct \( |{\text{BRS}}⟩ = \bigotimes R_i P_0 \prod_0 |D_±⟩ \). The corresponding \( R \)-tensors are shown in Tab. 1, where \( R \) is described in the QSpace language, a practical framework for implementing non-abelian symmetries in tensor networks. In the QSpace framework, the tensor \( R \) can be decomposed into the reduced multiplet data \(|R⟩\) and the Clebsch-Gordan coefficients (CGCs) \( C \), i.e.,

\[
R = \sum_{\{S_i\}} \sum_{\{m_i\}} \|R⟩|S_1,S_2,S_3⟩(C^{S_{\text{hex}}=0}_{S_1,S_2,S_3})m_1,m_2,m_3, \quad (12)
\]

where \( \{S_i\} \) are the spin quantum numbers and \( m_i \in \{-S_i,+S_i\} \) are the \( z \) component of magnetic quantum number of paired-up virtual spins. Notice that \( R \) represents a tensor of order four where the fourth dimension specifies the symmetry of the total tensor, which here for the HSS state by definition always a singlet state \( (S_{\text{hex}} = 0) \). The last dimension therefore represents a singular dimension which can be safely ignored. Overall then, what represents a \( 4 \times 4 \times 4 \) tensor in full state space becomes a \( 2 \times 2 \times 2 \) tensor in the reduced multiplet space.

To be more specific, in Tab. 1, the first column enumerates different records or fusion channels, i.e., the different ways three virtual particles can be coupled into a hexagon spin singlet. The second column shows the spin quantum numbers (\( S \)-labels) of these virtual particles: Respectively, there are three spin-0 (in record No. 1), two spin-1’s and a spin-0 (No. 2-4), and three spin-1 particles (No. 5). The third column contains the dimensions of the CGC tensor \( (C^{S_{\text{hex}}=0}_{S_1,S_2,S_3})m_1,m_2,m_3, \) which vary in different records because of different spin quantum numbers \( \{S_1,S_2,S_3\} \). The last column gives the value of the reduced tensor element obtained when coupling \( S_1,S_2 \) and \( S_3 \) together to obtain a hexagon singlet. The norm of \(|R⟩\) is chosen such that the weight in the first record is 1, the other weights are denoted as \( \alpha \) in records 2-4, and \( \beta \) in the last record, respectively. Note that in Tab. 1 and other tables for QSpace tensors hereafter, the actual number of the CGC coefficients are not shown. In practice, the CGC tensor in each channel are normalized in such a way that the largest tensor element equals \( ±1 \), and the first nonzero coefficient being positive.

Similar to the hexagon tensor \( R \), the triangle tensor \( M \) can also be determined by performing a few steps of contractions [as indicated in Fig. 4(d)], with the three physical \( S \) = 1 spins combined into a single state space \( (3^3 = 27 \text{ states reduced to 7 multiplets}) \), and this results in a tensor of order four. Given the hexagonal tensor-network state consisting of tensors \( M \) and \( R \), we take it as a variational ground-state wave function of the spin-1 KHAF model (with the Hamiltonian \( H_{\text{KHAF}} = \sum_{(i,j)} S_i \cdot S_j \)) and first calculate the energy.
per site \( e_0 \) using the infinite projected entangled-pair state (iPEPS) contraction algorithm [via the boundary matrix product state (MPS) scheme].\(^{22,23}\) The resulting energy per site is \( e_0 = \frac{1}{3.1670602} \) for \(|\text{BRS}\rangle\), while \( e_0 = \frac{1}{0.83127138} \) for \(|\text{BRS}\rangle\).

In Fig. 6(a), we show the numerical results of various correlation functions of \(|\text{BRS}\rangle\), which are also obtained by iPEPS contractions (thus measured on an infinite kagome lattice). The correlation functions include the spin-spin \( C_{SS}(j-i) = \langle S_i \cdot S_j \rangle \), the quadrupole-quadrupole \( C_{QQ}(j-i) = \langle Q_i \cdot Q_j \rangle \), and the dimer-dimer \( C_{DD}(j-i) = \langle (S_i + S_{i+1}) (S_j + S_{j+1}) \rangle - \langle S_i \rangle \cdot \langle S_j \rangle \) correlations. All the correlation functions are calculated in an SU(2)-invariant manner, i.e., the \( C_{SS} \) and \( C_{QQ} \) correlations are computed using irreducible tensor operators \( S^{(1)} = \{ S^+, S^z, S^- \} \) and \( Q^{(2)} = \{ (S^+)^2, -(S^+ S^- + S^- S^+), \sqrt{\frac{3}{2}} (S^z)^2 - 2, (S^- S^z + S^z S^-), (S^-)^2 \} \), respectively. The correlations are measured along the vertical line marked as a red dashed line in Fig. 5(b), with length unit \( a_x = 1 \) being specified there, and are found to decay exponentially, as expected from the rigorous proof in terms of Schwinger bosons (Sec. II and Appendix A). The correlation lengths \( \xi \), extracted by linear fittings from the semi-log plot, are found to be rather short.

Besides the infinite kagome lattice, we are also interested in evaluating the properties of BRS on the cylindrical geometries (see the XC and YC geometries in Fig. 5). In Fig. 6(b), we show the entanglement entropy results \( S(L) \) of XC and YC geometries, versus various cylinder circumferences (up to \( L = 16 \) for both geometries), which measures the quantum entanglement between the two half-infinite cylinders. As shown in Fig. 6(b), we extrapolate \( S(L) \) using the formula \( S = \alpha L - \gamma \) and get \( \gamma \approx 0 \) as \( L \to 0 \), in both XC and YC cases. This observation shows unambiguously that the BRS possesses no long-range entanglement and thus no intrinsic topological order, this is due to the local constraint arising from the benzene construction (related discussions in Sec. II B).

### Table I. Hexagon tensor \( R \) of even and odd BRS, as a Q-space object. \( \{ S_i \} \) are the spin quantum numbers, and the third column after it demonstrates the dimensions of corresponding CGC tensor in each channel (simply also determined from the symmetry labels \( S_i \)), the fourth column stores the reduced multiplet elements \(| \langle R \rangle |\).

| B(R)even | \( S_1, S_2, S_3 \) | dimensions | \(| \langle R \rangle |\) |
|----------|-----------------|-------------|-----------------|
| 1. 0 , 0 , 0 | 1 \times 1 | 1 |
| 2. 0 , 1 , 1 | 1 \times 3 | 1/3 |
| 3. 1 , 0 , 1 | 3 \times 1 | 1/3 |
| 4. 1 , 1 , 0 | 3 \times 3 | 1/3 |
| 5. 1 , 1 , 1 | 3 \times 3 | 1/3 |

FIG. 6. (Color online) (a) The spin-spin, quadrupole-quadrupole, and dimer-dimer correlation functions of the \(|\text{BRS}\rangle\), obtained by SU(2) iPEPS contractions. All of the correlation functions \( C(x) \) are found to decay exponentially, as \( C(x) \sim \exp(-x/\xi) \). The correlation lengths \( \xi \) are obtained from fitting the data. (b) Entanglement entropies of the odd BRS on XC and YC geometries with various circumferences (up to \( L = 16 \)), the entropy data extrapolate to \( \gamma = 0 \) in the \( L = 0 \) limit.

### FIG. 7. (Color online) (a) Energy expectation values (per site) \( e_0 \) of Hida’s HSS state on cylinders with various circumferences \( L \) [see Fig. 5 for the illustration of the XC and YC geometries, the length units \( a_x, a_y \) are also shown in Fig. 5(b)]. The energy results converge very fast to the thermodynamic limit obtained by iPEPS contractions. (b) Entanglement entropy results extrapolate to \( \gamma \approx 0 \) as \( L \to 0 \), for both XC and YC geometries.

### B. Hida’s Hexagon-Singlet Solid State

Next, we turn to the original HSS state proposed in Ref. 15 (henceforth to be referred to as Hida’s HSS state, or more briefly the Hida point), where the hexagon singlet \(|\psi_0\rangle \) in Eq. (9) is chosen to be the ground state of a six-site (hexagonal) spin-1/2 Heisenberg ring,

\[
H_0 = J_1 \sum_{i,j} S_i \cdot S_j + J_2 \sum_{\langle i,j \rangle} S_i \cdot S_j + J_3 \sum_{\langle \langle i,j \rangle \rangle} S_i \cdot S_j,
\]

where \( J_1 \) is the NN coupling, \( J_2, J_3 \) are the second- and third-NN interactions, and \( S \)'s are the \( S = 1/2 \) spin operators. Following Ref. 15, we diagonalize the Heisenberg ring with only NN couplings (couplings \( J_1 = 1, J_2 = J_3 = 0 \)) in a hexagon \( h \), and find five orthonormal singlet eigenstates, with energies \(-2.8028, -1.5000, -0.5000, -0.5000, \) and \( 0.8028 \). Notice that there is a considerable gap (~1.3) between the resulting ground and the first
excited states. We select the lowest singlet ground state, and determine the corresponding SU(2)-invariant tensor $R$ as shown in Tab. II. Since there are only two multiplets (a $S = 0$ singlet and a $S = 1$ triplet) on each geometric bond, the reduced bond dimension is thus $D^* = 2$, corresponding to four individual states ($D = 4$).

Table II. Hexagon tensor $R$ of Hida’s HSS state, where $\alpha$ and $\beta$ are used as variational parameters below. Notice that to keep the $C_3$ discrete rotational lattice symmetry, the coefficients of records 2 to 4 should be equal (denoted by $\alpha$).

| No. | $[S_1, S_2, S_3]$ dimensions | $||R||$ |
|-----|-------------------------------|--------|
| 1   | [0, 0, 0] 1x1x1              | 1      |
| 2   | [0, 1, 1] 1x3x3              | $\alpha = -0.2457$ |
| 3   | [1, 0, 1] 3x1x3              | $\alpha = -0.2457$ |
| 4   | [1, 1, 1] 3x3x1              | $\alpha = -0.2457$ |
| 5   | [1, 1, 1] 3x3x3              | $\beta = 0.1315$ |

Table III. Energy expectation values of the Hida’s HSS state for the spin-1 KHAF model, obtained by SU(2) iPEPS contractions.

| $D^*$ | $D_c$ | $e_0$ | truncation error |
|-------|-------|-------|-----------------|
| 4     | 8     | $-1.359944730698$ | $2 \times 10^{-6}$ |
| 5     | 11    | $-1.359910140148$ | $4 \times 10^{-8}$ |
| 6     | 16    | $-1.359909517302$ | $2 \times 10^{-12}$ |
| 10    | 26    | $-1.359909517316$ | $2 \times 10^{-13}$ |
| 16    | 44    | $-1.359909517316$ | $2 \times 10^{-15}$ |

Given the local tensors, we again consider two kinds of geometries for the evaluation of observables: the infinite kagome lattice and cylinders (including XC and YC geometries). Firstly, the energy per site $e_0$ is calculated through iPEPS contractions and the results are shown in Tab. III. The small truncation error suggests that the data are very well converged when more than $D^*_c = 10$ multiplets (corresponding to $D_c = 26$ states) are retained in the geometric bond of the boundary MPS.

Besides the iPEPS calculations, we also performed exact contractions on various cylinders, the results are shown in Fig. 7(a). Notably, the energy expectation value is determined as $e_0 = -1.359910231678$ for XC12, in excellent agreement with the accurate iPEPS results in Tab. III. This value is also in accordance with that in Ref. 19, where the HSS energy is estimated as $-1.36$ based on the exact diagonalization results on several small clusters. This variational energy is lower than that of the RAL state ($\approx -1.27$) in Ref. 17, but still higher than the best estimate $e_0 \approx -1.410$ (of an SVBC state) in Refs. 19 and 21 for the actual ground state of the spin-1 KHAF model.

In Fig. 7(b), we show the entanglement entropies on (both XC and YC) cylinders of various circumferences $L$. They extrapolate to zero in the $L = 0$ limit, meaning that Hida’s HSS state possesses no intrinsic topological order, just as the [BRS]$_O$ investigated in Sec. IIIA. This is an expected and consistent observation, because the rigorous proof in Sec. II guarantees that all HSS states are gapped spin liquids and thus should belong to the same (non-topological) phase.

In addition, we also studied the entanglement spectra (ES) of the HSS states on various cylinders through exact contractions.$^{34,35}$ In the Appendix B, we show results at the Hida point, where a nonvanishing triplet gap has been observed in the ES [Fig. S1(a)]. This is in contrast to the $S = 2$ AKLT state on a square lattice, where the gaps in the ES decrease as the system size increases, and finally vanish in the thermodynamic limit [see Fig. S1(b)].$^{34,35}$ The absence of a gapless edge excitation in the HSS state indicates that it has no symmetry-protected topological (SPT) order, since gapless edge modes necessarily appear in the SPT phases.$^{37,38}$

IV. VARIATIONAL STUDY OF THE KAGOME HEISENBERG ANTFERROMAGNETIC MODEL

In this section, we discuss variational energies of the HSS states for the spin-1 KHAF model, and furthermore search for lower variational energy within the present $D^* = 2$ ($D = 4$) tensor network ansatz. Among various ways to perform the variational studies, we will discuss three cases below, which turn out to produce consistent results.

First, as discussed in Sec. II B, we can expand the hexagon singlet with the over-complete basis of valence bonds (as shown in Fig. 2). In addition, by comparing the tensor elements in Tabs. I and II, we can see that Hida’s HSS state has similar weights as [BRS]$_O$ in corresponding channels, which suggests that these two states may have a big overlap, and the dominating hexagon-singlet configuration in the former might consist of NN valence bonds. Thus we could add hexagon-singlet configuration containing longer-range bonds in addition to the configurations $|D_\alpha\rangle$ of the [BRS]$_O$ [Fig. 2(a)]. Here we consider the configuration in Fig. 2(b), which consists...
of two NN and one second NN bonds in each hexagon. Associating a weight ω with this hexagon-singlet configuration (denoted as |N⟩), i.e., the singlet state can be defined as

\[ |\psi_\omega(\omega)⟩ = |D_–⟩ + ω|N⟩ \] (14)

in each hexagon, and we thus construct a one-parameter HSS state family \( |\Psi(\omega)⟩ = 0, 1 \prod_\omega |\psi_\omega(\omega)⟩ \) according to Eqs. (9-11). As shown in Fig. 8(a), by tuning ω, we can indeed connect smoothly the |BRS⟩ of the Hida point (at ω ≃ 0.5826). Notably, the lowest energy state turns out to deviate slightly from the Hida point, although the energy difference between them is rather tiny. In addition, the exact expansion (14) at the Hida point (ω ≃ 0.5826) means that Hida’s HSS state, like the BRs in Fig. 3(a), also has a simple resonating AKLT-loop picture, but with both NN and third-NN valence bonds (instead of only NN bonds) across each hexagon.

At a first glance, the fact that the hexagon singlet |G⟩ in Hida’s state can be expressed exactly as a superposition of |D–⟩ [of odd BRs, Fig. 2(a)] and |N⟩ [Fig. 2(b)] as in Eq. (14) is surprising, because chances are these three normalized while non-orthogonal states (vectors) do not lie on the same plane. In fact, there are two independent components α and β for the C3 lattice rotational invariant states, as shown in Tabs. I and II, which can parametrize a sphere after properly reorganized. Nevertheless, it turns out that these three states satisfy the condition \( \langle G|D–|^2 + ⟨G|N|^2 + ⟨N|D–|^2 = 1 + 2⟨G|D–⟩⟨G|N⟩⟨N|D–⟩ \), meaning |G⟩, |N⟩, and |D–⟩ are on the same unit circle and thus are linearly dependent. As a more careful analysis shows, the reason for this is that these hexagon singlets (and the corresponding non-symmetry-breaking HSS states) are also eigenstates of the hexagon-inversion symmetry operator (with eigenvalues ±1). Therefore, it adds one more constraint and leaves only one independent parameter ω in Eq. (14) when one expands one hexagon singlet state (say, |G⟩) with two other states (say, |N⟩ and |D–⟩) of the same inversion parity (−1 here). Interestingly, the other state |D+⟩ in Fig. 2 not used in the expansion of |G⟩ above belongs to an +1 eigenstate of hexagon-inversion symmetry, which completes the set of three bases (|D+⟩ and |N⟩) for expanding any HSS states preserving C3 rotational symmetry.

Second, since α and β are the coefficients of the orthogonal normal-singlet channels (see Tabs. I and II), tuning them is the most efficient way to explore all states in the HSS family. In Fig. 8(b), e0 versus energy difference εA – εB, which are the energy expectation values on two kinds of triangles, are shown for each curve with fixed α and varying β parameters. From this we can see that the global minimum is located at e0 = −1.36000, which is again only slightly lower than the value e0 = −1.35991 of the Hida point.

Besides the above two simulations, the third approach we have adopted is to introduce second- (J2) and third-NN (J3) couplings in the hexagon [see Eq. (13)], so as to strengthen longer-range valence bonds. Through tuning J2 and J3 in (13), we find the same lowest variational energy for the spin-1 KHAF model, which, within the HSS states, is e0 = −1.36000, located at εA – εB = 0, i.e., without any trimerization order.

Beyond the HSS states, we are also interested in improving the variational energy of the spin-1 KHAF, within the present \( D^* = 2 \) tensor network ansatz shown in Fig. 4. Notice that the present M (obtained by combining three projectors P) itself leaves the three physical spins uncorrelated, the correlation only enters through the singlet construction in R. If we now instead introduce correlations directly between physical spins within M across hexagons (where R still allows singlets within the hexagons only), the HSS picture will generally be shuffled [see Fig. 9(a)]. In practice, we add one more parameter τ, in addition to α and β, to tune tensor M and thus explore the variational energies. This is inspired from the more sophisticated imaginary-time evolution approach, \( 39-41 \) where the operators O(τ) = e−τH,τ are applied iteratively to an initial state until it converges (to the ground state), where \( H_{\Delta(V)} = \sum_{(i,j)\in\Delta(V)} S_i \cdot S_j \)
are the local Hamiltonians (consisting of NN couplings, S’s are spin-1 operators) acting on local spins in Δ (\(\nabla\)) triangle [see Fig. 9(b)]\(^{21,24,42,43}\). Here, instead of diving into the heavy numerics, we instead mimic a single step of projection by applying \(O(\tau)\) on down (denoted by A) triangles,\(^{35}\) which can be expanded as \(1 − \tau H_0 + O(\tau^2)\).

We consider the first two terms, absorb it to the \(M\) tensor, and thus introduce a new variational parameter \(\tau\) in the tensor network state.

In Fig. 10(a), we show that when fixing the parameter \(\tau \neq 0\), and scanning through various \(\alpha, \beta\) parameters, the lowest energy state turns out to be located at \(e_A − e_B \neq 0\); while the sign and magnitude of \(\Delta e = 2(e_A − e_B)/3\) depends on the values of the three parameters of \(\alpha, \beta\) and \(\tau\). We repeat the analysis of Fig. 10(a) for various \(\tau\)-values and for each collect that curve containing the lowest variational energies, like the \(\alpha = 0.5\) curve for \(\tau = 0.23\) in Fig. 10(a). We show them in Fig. 10(b), from which it is found that the best variational energy is \(e_0 = −1.3871\), significantly lower than the non-symmetry-breaking HSS state, and the energy difference \(\Delta e = 2(e_A − e_B)/3\approx 0.2083\), about 20% of \(e_0\). This implies that the SVBC has lower energy than any HSS states with \(D^* = 2\). This conclusion and even the magnitude of trimerization order parameter, obtained only by considering three parameters here, are in nice agreement with previous calculations of Refs. 19, 21, and 24, where the estimate \(e_0 \approx −1.41\) of a SVBC state was obtained with much larger bond dimensions and more sophisticated numerical algorithms. However, our conclusion that SVBC states yield a lower energy than HSS states disagrees with the main conclusion of NN14 in Ref. 26. Possible reasons for this disagreement are given in Appendix C.

V. SUMMARY AND DISCUSSION

To summarize, we have performed a systematic investigation of the hexagon-singlet solid states for the spin-1 kagome antiferromagnet. Through the Schwinger boson representation, we have shown that the HSS states are gapped paramagnets, which have a hidden resonating AKLT-loop picture when the hexagon singlet is decomposed within the over-complete valence-bond bases. However, in contrast to the RAL state (equal weight superposition of all possible loop configurations) which is a \(\mathbb{Z}_2\) spin liquid, the HSS states, owing to the local constraint of hexagon-singlet construction, turn out to have no intrinsic topological order. By performing numeric simulations using the tensor network representation, we have shown that the HSS states are indeed gapped spin liquid, with all correlation functions decaying exponentially, and the results of the entanglement entropy and spectra confirm that these states are topologically trivial. Furthermore, we find out that the best variational energy for the spin-1 KHAF model, among all the non-symmetry-breaking HSS states, is \(e_0 = −1.3600\). Moreover, by an enlightening variational study, we have shown that, within the present \(D^* = 2\) tensor network ansatz, the simplex valence bond crystal state \((e_0 = −1.3871)\) is more energetically favorable than the non-symmetry-breaking HSS state.

An interesting issue we leave for a future study is to find a realistic Hamiltonian which could stabilize the HSS phase: such a Hamiltonian might contain second- or even third-NN couplings in the hexagons, which can still be conveniently treated in the present tensor network ansatz in Fig. 4. Moreover, through the investigations of the RAL\(^{17}\) and HSS states, it has been shown that the resonating AKLT loops, which constitute a rather general representation of spin-1 many-body singlets, are able to describe a variety of states, ranging from the topological ordered RAL to the topologically trivial HSS states. Therefore, we expect that there could be more exotic quantum states by exploring the resonating AKLT loop family in future studies.

VI. ACKNOWLEDGMENT

HHT acknowledges Tai-Kai Ng for introducing him the hexagon-singlet ansatz and Yi Zhou for stimulating discussions. WL and HHT would like to thank Meng Cheng and Zheng-Xin Liu for fruitful discussions on spin-1 kagome antiferromagnets. WL is further indebted to Tao Liu and Gang Su for helpful discussions. This work has been supported by the DFG project SFB-TR12, the EU project SIQS, and the DFG Cluster of Excellence NIM. AW further acknowledges support by DFG grant WE-4819/1-1.

Appendix A: Proof of the exponentially decaying correlations in HSS ansatz

In this Appendix, we provide technical details on the proof that the HSS ansatz all have exponentially decaying spin correlations. This relies on the spin-coherent state representation of the SU(2) Schwinger boson states. Following Refs. 29 and 31, \(S \equiv 1\) spin coherent states are defined by

\[
|\Omega\rangle = \frac{1}{2}(ub_{\uparrow} + vb_{\downarrow})^2|\text{vac}\rangle, \quad \text{(S1)}
\]

where the \(O(3)\) vector \(\Omega\) is parametrized by the solid angle, \(\Omega = (\sin \theta \cos \phi, \sin \theta \sin \phi, \cos \theta)\), and \(u\) and \(v\) are given by \(u(\theta, \phi) = \cos \frac{\theta}{2} e^{-i\frac{\phi}{2}}\) and \(v(\theta, \phi) = \sin \frac{\theta}{2} e^{i\frac{\phi}{2}}\). The spin coherent states satisfy the following relations:

\[
\frac{3}{4\pi} \int d\Omega \langle \Omega | \Omega \rangle = I, \quad \text{(S2)}
\]

\[
\frac{3}{2\pi} \int d\Omega \Omega_i |\Omega\rangle \langle \Omega| = S, \quad \text{(S3)}
\]

where the integration \(\int d\Omega\) is over the solid angle, \(\int d\Omega = \int_0^\pi \sin \theta d\theta \int_0^{2\pi} d\phi\).
By using these relations, the norm of the HSS wave function (3) is expressed as

$$\langle \Psi | \Psi \rangle = \frac{3}{4\pi} \int d\Omega \langle \Psi | \Omega \rangle \langle \Omega | \Psi \rangle$$

$$= \frac{3}{4\pi} \int d\Omega |\Psi(\Omega)|^2,$$  \hspace{1cm} (S4)

where $\Psi(\Omega)$ is given by

$$\Psi(\Omega) = \langle \text{vac} | \prod_{\Omega} \left[ \sum_{ij,kl,mn} \tilde{w}_{ij} \tilde{w}_{kl} \tilde{w}_{mn} (u_{\Omega,i}v_{\Omega,j} - v_{\Omega,i}u_{\Omega,j})(u_{\Omega,k}v_{\Omega,l} - v_{\Omega,k}u_{\Omega,l})(u_{\Omega,m}v_{\Omega,n} - v_{\Omega,m}u_{\Omega,n}) \right] \prod_i |\Omega_i\rangle \right],$$  \hspace{1cm} (S5)

and $|\Psi(\Omega)|^2$ is written as

$$|\Psi(\Omega)|^2 \propto \prod_{\Omega} \left[ \sum_{ij,kl,mn} \tilde{w}_{ij} \tilde{w}_{kl} \tilde{w}_{mn} (u_{\Omega,i}v_{\Omega,j} - v_{\Omega,i}u_{\Omega,j})(u_{\Omega,k}v_{\Omega,l} - v_{\Omega,k}u_{\Omega,l})(u_{\Omega,m}v_{\Omega,n} - v_{\Omega,m}u_{\Omega,n}) \right]^2.$$  \hspace{1cm} (S6)

Note that the norm $\langle \Psi | \Psi \rangle$ in (S4) is proportional to the following partition function of a classical statistical model defined on the same kagome lattice:

$$Z = \int d\Omega \ |\Psi(\Omega)|^2$$

$$= \int d\Omega \prod_{\Omega} \left[ \sum_{ij,kl,mn} \tilde{w}_{ij} \tilde{w}_{kl} \tilde{w}_{mn} (u_{\Omega,i}v_{\Omega,j} - v_{\Omega,i}u_{\Omega,j})(u_{\Omega,k}v_{\Omega,l} - v_{\Omega,k}u_{\Omega,l})(u_{\Omega,m}v_{\Omega,n} - v_{\Omega,m}u_{\Omega,n}) \right]^2$$

$$= \int d\Omega \exp \left( \ln \prod_{\Omega} \left[ \sum_{ij,kl,mn} \tilde{w}_{ij} \tilde{w}_{kl} \tilde{w}_{mn} (u_{\Omega,i}v_{\Omega,j} - v_{\Omega,i}u_{\Omega,j})(u_{\Omega,k}v_{\Omega,l} - v_{\Omega,k}u_{\Omega,l})(u_{\Omega,m}v_{\Omega,n} - v_{\Omega,m}u_{\Omega,n}) \right]^2 \right)$$

$$= \int d\Omega \exp \left( 2 \ln \sum_{\Omega} \left[ \sum_{ij,kl,mn} \tilde{w}_{ij} \tilde{w}_{kl} \tilde{w}_{mn} (u_{\Omega,i}v_{\Omega,j} - v_{\Omega,i}u_{\Omega,j})(u_{\Omega,k}v_{\Omega,l} - v_{\Omega,k}u_{\Omega,l})(u_{\Omega,m}v_{\Omega,n} - v_{\Omega,m}u_{\Omega,n}) \right] \right)$$

$$= \int d\Omega \exp \left( - \sum_{\Omega} h_{\Omega} \right),$$  \hspace{1cm} (S7)

where $h_{\Omega}$ is the classical Hamiltonian describing local interactions among the O(3) vectors within each hexagon

$$h_{\Omega} = -2 \ln \left[ \sum_{ij,kl,mn} \tilde{w}_{ij} \tilde{w}_{kl} \tilde{w}_{mn} (u_{\Omega,i}v_{\Omega,j} - v_{\Omega,i}u_{\Omega,j})(u_{\Omega,k}v_{\Omega,l} - v_{\Omega,k}u_{\Omega,l})(u_{\Omega,m}v_{\Omega,n} - v_{\Omega,m}u_{\Omega,n}) \right].$$  \hspace{1cm} (S8)

Similarly, the spin-spin correlation function can be represented by using spin coherent states as

$$\frac{\langle \Psi | S_i \cdot S_j | \Psi \rangle}{\langle \Psi | \Psi \rangle} = \frac{4}{Z} \int d\Omega |\Psi(\Omega)|^2 \Omega_i \cdot \Omega_j \hspace{1cm} (i \neq j)$$

$$= \frac{4}{Z} \int d\Omega \Omega_i \cdot \Omega_j \exp \left( - \sum_{\Omega} h_{\Omega} \right).$$  \hspace{1cm} (S9)
FIG. S1. (Color online) The entanglement spectra (ES) on YC geometries, with various cylinder widths $L$. Each symbol in the figure labels a multiplet, instead of an individual plain state. The spectra of different circumferences have been offset so as the lowest eigenvalues are zero. (a) The ES of Hida’s HSS state on cylinders with $L = 4, 6, 8, 10, 12, 14$. There exists a non-vanishing singlet-triplet gap ($\delta \sim 1.6$) in the spectra. (b) The ES of the $S = 2$ square-lattice AKLT state on cylinders with $L = 4, 6, 8, 10, 12, 14$, where the singlet-triplet gaps $\delta \sim (1/L)$ extrapolate to zero in the large $L$ limit.

This shows that the spin-spin correlation function in the HSS ansatz is equivalent to the two-point correlation function between the O(3) vectors in a 2D classical statistical model. Since the classical statistical model is at finite temperature and has only short-range interactions, Mermin-Wagner theorem rules out the possibility of long-range order and the correlation function (S9) decays exponentially.

Other correlation functions, such as the quadrupole-quadrupole and dimer-dimer correlation functions, can be similarly represented as correlation functions in the same classical statistical model. Apart from a qualitative understanding of their behaviors using the Mermin-Wagner theorem, an additional benefit of this quantum-classical mapping is that Monte Carlo techniques can also be applied directly to compute the physical quantities accurately (see e.g. Ref. 44 for this application to the 3D AKLT states).

Appendix B: Entanglement Spectra

In this Appendix, we show the results of the entanglement spectra (ES) on cylinders with various widths $L$. By performing exact contraction for small $L$ and MPS-based approximate contractions for large $L$ ($\geq 12$), one can get converged left and right vectors $V_l(\alpha, \alpha')$ and $V_r(\alpha, \alpha')$ after a sufficient number (typically $10 \sim 20$) of iterations. The vectors $V_l, V_r$ are the dominating (left and right) eigenstates of the transfer-matrix of the double-layer cylinder tensor network: and $|\alpha\rangle, |\alpha'\rangle$ are the virtual bases on the geometric bond. Since the $|\alpha\rangle$ and $|\alpha'\rangle$ bases are not orthonormal to each other, it is important to notice that $V_l, V_r$ themselves are not the reduced density matrices of the half-cylinder. The reduced density matrix can be obtained through transformation $\rho = \sqrt{V_r} V_l \sqrt{V_r}$.

By diagonalizing $\rho = U \Lambda U'$, in Fig. S1 we show the entanglement spectrum, i.e., the minus logarithm of the density matrix spectrum $-\log(\Lambda)$, on cylinders of various widths. Notice that since SU(2) symmetry has been implemented in the tensor network, each symbol in Fig. S1 represents a multiplet with well-defined spin quantum number $S$, so we can tell the spin $S$ of each level. In Fig. S1(a), there is a distinct gap between lowest singlet and the first excited triplet state, which does not vanish as $L$ increases. This is in contrast to a typical $S = 2$ AKLT state on a square lattice: in Fig. S1(b) we show the ES of the $S = 2$ AKLT state on cylinders, for which the singlet-triplet gap $\delta$ is proportional to the inverse width $1/L$. $\delta$ vanishes in the infinite-$L$ limit, suggesting a gapless edge mode, in agreement with the results in Refs. 34 and 36. The different behaviors of the HSS and $S = 2$ square-lattice AKLT states implies that the former is a trivial insulator phase, and does not belong to a SPT phase.

Appendix C: Comments on NN14’s DMRG study on HSS states

In this Appendix, we address the relation between our present tensor network study of the HSS state and the work of NN14 in Ref. 26, who studied the same model using DMRG, but, in contrast to us and Refs. 19, 21, and 24, concluded that the model’s ground state is an HSS state. They performed DMRG simulations for the spin-1 KHAF on four types of clusters: (i) cylindrical clusters; (ii) periodic clusters; (iii) clusters with open boundary conditions (OBC) purposefully designed to favor a HSS ground state, by choosing cluster shapes that contain only hexagons along the edges and putting spin-1/2’s on the outermost sites (called HSS clusters); and (iv) clusters with OBC purposefully designed to favor SVBC order, by choosing cluster shapes that contain only triangles around the edges (called SVBC clusters). For (i) to (iii) they found ground states without clear signa-
tures of trimerization, while for (iv) they found a ground state that clearly trimerizes. Their conclusion of vanishing trimerization order on (i) is in direct contradiction to CL’s observation that stable trimerization order exists on cylinders with width $L = 8$.\textsuperscript{19} NN14 estimated the energy per site in the bulk, $\epsilon_0$, by finite-size extrapolations to the thermodynamic limit. Their SVBC clusters, (iv), yielded the highest extrapolated energy ($\epsilon_0 = -1.391(2)$); their cylindrical and periodic clusters, (i) and (ii), yielded extrapolated energies ($\epsilon_0 = -1.40988$ and $\epsilon_0 = -1.409(5)$, respectively) consistent with those of CL\textsuperscript{19} and Liu et al.\textsuperscript{21}; and their HSS clusters, (iii), yielded the lowest extrapolated energy ($\epsilon_0 = -1.41095$). Based on this evidence, NN14 argued that the ground state is a HSS state, and not the SVBC state advocated in Refs. 19, 21, and 24.

In our opinion, NN14’s conclusion is flawed because the tool that they use to distinguish the HSS and SVBC states, namely ground-state DMRG on finite-size clusters, relies on boundary effects to distinguish these states. However, such a tool can not reliably estimate the difference in bulk $\epsilon_0$ for these two types of state, since by definition, the bulk value of $\epsilon_0$ is the value obtained for clusters so large that boundary effects vanish. Not surprisingly, the data for their HSS and SVBC clusters shown in their Fig. 4 have a much stronger finite-size dependence (larger slope of data plotted versus inverse system size) than their cylinder and periodic clusters. The fact that their extrapolations from the HSS and SVBC clusters disagree with each other in the thermodynamic limit just means that these clusters are not yet large enough to give reliable bulk $\epsilon_0$ values. Thus the discrepancy between NN14’s $\epsilon_0$ values for HSS and SVBC clusters in our opinion does not reflect the true difference in the $\epsilon_0$ values of bulk SVBC and bulk HSS states; rather, it reflects the error bar, induced by boundary effects, in their determination of the bulk $\epsilon_0$ of the true ground state.

We would also like to make a comment regarding the trick of pinning spin-1/2’s on the open boundary of a spin-1 model. This trick was first developed for simulating a spin-1 Heisenberg chain, where two spin-1/2’s are put on the ends of the open chain to remove the edge states (and related degeneracies).\textsuperscript{45} Some of us (WL, AW, JvD) have actually used this trick ourselves in recent work on spin-1 Heisenberg chains.\textsuperscript{46} In that context, this trick leads to a better numerical convergence for the bulk properties, since it binds the edge modes localized on both open ends. Importantly, however, it does not change the bulk physics, i.e., the same value for $\epsilon_0$ is obtained with or without employing spin-1/2’s on the ends.\textsuperscript{45,46} In contrast, in NN14’s work, the presence or absence of spin-1/2 particles on the outermost sites of their HSS or SVBC cluster, respectively, is an attempt to use boundary effects to stabilize two states with different bulk properties. As argued above, this strategy is intrinsically flawed. Thus, in our opinion NN14 are mistaken in believing that their DMRG calculations on HSS clusters can be used to reliably predict the bulk properties of HSS states. Their DMRG calculations target purely ground state properties, and if a certain type of state (here the HSS state) happens not to be the actual ground state (as argued in the present paper and in Refs. 19, 21, and 24), then its bulk properties are simply inaccessible to NN14’s DMRG simulations. In contrast, through the present work we find an explicit tensor network state representation for the HSS state family, which makes Hida’s HSS state readily accessible with tensor network methods. We show unambiguously that the best HSS variational energy for the spin-1 KHAF model is $-1.3600$, much higher than NN’s “HSS” energy ($-1.41095$).

In contrast to NN14’s HSS and SVBC data, their data for cylinder and periodic clusters show much weaker finite-size effects (smaller slopes when plotted versus inverse system size). We believe that the $\epsilon_0$ values extrapolated from this data do reflect the bulk $\epsilon_0$ value of the true ground state rather accurately – they are certainly consistent with our own $\epsilon_0$ values and those of CL. We find it all the more surprising and perplexing that NN14 did not observe evidence for SVBC order in their cylinder clusters, whereas CL, who used essentially similar cylindrical clusters, did.\textsuperscript{19} We suspect that in NN14’s calculations the presence of a symmetry-breaking ground state was still hidden by boundary effects, which can cause a non-symmetry-breaking ground state to appear through superposition. Indeed, CL reported that in their calculations SVBC order emerged clearly only in their cylinders of largest circumference (width 8).

We conclude with a comment about the bond dimension used in the present tensor network study, namely $D^* = 2$ (corresponding to $0 \oplus 1$ in Tabs. I and II), or $D = 4$. One might ask: is our conclusion in Section IV that HSS states have higher energy than SVBC states robust with respect to increasing $D$? Perhaps HSS states would yield a lower energy than SVBS states if each hexagon were allowed to involve not only spin-1/2 virtual particles but also higher-spin virtual particles? In this regard, we note that the HSS states studied here are a subset of the much more general class of tensor network states studied by Liu et al. in Ref. 21, where tensor network variational calculations were performed with bond dimensions as large as $D = 20$, implying that many more virtual particles with higher spin were included. Those large-$D$ calculations showed unambiguously that the spin-1 KHAF model possesses a SVBC ground state with $\epsilon_0 = -1.41035$. Therefore, the conclusion of the present work, that HSS states lie significantly higher in energy than SVBC states, is not specific to using an HSS ansatz with $D = 4$, as done here; instead, the results of Ref. 21 demonstrate that it holds throughout as $D$ is increased up to $D = 20$, a value sufficiently large that the results reported in Ref. 21 were well converged.
1. S. Yan, D. A. Huse, and S. White, Science 332, 1173 (2011).
2. S. Depenbrock, I. P. McCulloch, and U. Schollwöck, Phys. Rev. Lett. 109, 067201 (2012).
3. O. Götze, D. J. J. Farnell, R. F. Bishop, P. H. Y. Li, and J. Richter, Phys. Rev. B 84, 224428 (2011).
4. K. Awaga, T. Okuno, A. Yamaguchi, M. Hasegawa, T. Inabe, Y. Maruyama, and N. Wada, Phys. Rev. B 49, 3975 (1994).
5. I. Watanabe, N. Wada, H. Yano, T. Okuno, K. Awaga, S. Ohira, K. Nishiyama, and K. Nagamine, Phys. Rev. B 58, 2438 (1998).
6. N. Wada, T. Kobayashi, H. Yano, T. Okuno, A. Yamaguchi, and K. Awaga, J. Phys. Soc. Jpn. 66, 961 (1997).
7. T. Matsushita, N. Hamaguchi, T. Okuno, A. Yamaguchi, and H. Ishimoto, J. Phys. Soc. Jpn. 69, 4003 (2000).
8. H. Yao, L. Fu, and X.-L. Qi, arXiv:1012.4470 (2010).
9. W. Li, S. Yang, M. Cheng, Z.-X. Liu, and H.-H. Tu, Phys. Rev. B 89, 245128 (2014).
10. S. Bieri, M. Serbyn, T. Senthil, and P. A. Lee, Phys. Rev. B 84, 180403(R) (2011).
11. F. Wang and C. Xu, arXiv:1110.4091.
12. C. Xu, F. Wang, Y. Qi, L. Balents, and M. P. A. Fisher, Phys. Rev. Lett. 108, 087204 (2012).
13. S. Bieri, M. Serbyn, T. Senthil, and P. A. Lee, Phys. Rev. B 86, 224409 (2012).
14. H.-H. Tu, Phys. Rev. B 87, 041103 (2013).
15. K. Hida, J. Phys. Soc. Jpn. 69, 4003 (2000).
16. H. Yao, L. Fu, and X.-L. Qi, arXiv:1012.4470 (2010).
17. W. Li, S. Yang, M. Cheng, Z.-X. Liu, and H.-H. Tu, Phys. Rev. B 89, 174411 (2014).
18. I. Affleck, T. Kennedy, E. H. Lieb, and H. Tasaki, Phys. Rev. Lett. 59, 799 (1987); Commun. Math. Phys. 115, 477, (1988).
19. H. J. Changlani and A. M. Läuchli, arXiv:1406.4767 (2014).
20. R. N. White, Phys. Rev. Lett. 69, 2863 (1992).
21. T. Liu, W. Li, A. Weichselbaum, J. von Delft, and G. Su, arXiv:1406.5905 (2014).
22. F. Verstraete and J. I. Cirac, arXiv:cond-mat/0407066v1 (2004); V. Murg, F. Verstraete, and J. I. Cirac, Phys. Rev. A 75, 033605 (2007).
23. J. Jordan, R. Orus, G. Vidal, F. Verstraete, and J. I. Cirac, Phys. Rev. Lett. 101, 250602 (2008).
24. T. Picot and D. Poilblanc, arXiv:1406.7205 (2014).
25. Z. Caï, S. Chen, and Y. P. Wang, J. Phys.: Condens. Matter, 21, 456009 (2009).
26. S. Nishimoto and M. Nakamura, arXiv:1409.5870 (2014).
27. A. Kitaev and J. Preskill, Phys. Rev. Lett. 96, 110404 (2006).
28. M. Levin and X.-G. Wen, Phys. Rev. Lett. 96, 110405 (2006).
29. A. Auerbach, Interacting Electrons and Quantum Magnetism (Springer-Verlag, New York, 1994).
30. N. D. Mermign and H. Wagner, Phys. Rev. Lett. 17, 1133 (1966).
31. D. P. Arovas, A. Auerbach, and F. D. M. Haldane, Phys. Rev. Lett. 60, 531 (1988).
32. A. Weichselbaum, Ann. Phys. 327, 2972 (2012).
33. H.-C. Jiang, Z. Wang, and L. Balents, Nature Phys. 8, 902 (2012).
34. J. I. Cirac, D. Poilblanc, N. Schuch, and F. Verstraete, Phys. Rev. B 83, 245134 (2011).
35. D. Poilblanc and N. Schuch, Phys. Rev. B 87, 140407(R) (2013).
36. J. Lou, S. Tanaka, H. Katsura, and N. Kawashima, Phys. Rev. B 84, 245128 (2011).
37. X. Chen, Z.-X. Liu, and X.-G. Wen, Phys. Rev. B 84, 235141 (2011).
38. Z.-X. Liu and X.-G. Wen, Phys. Rev. Lett. 110, 067205 (2013).
39. G. Vidal, Phys. Rev. Lett. 98, 070201 (2007).
40. R. Orús and G. Vidal, Phys. Rev. B 78, 155117 (2008).
41. H.-C. Jiang, Z.-Y. Weng, and T. Xiang, Phys. Rev. Lett. 101, 090603 (2008).
42. T. Liu, S.-J. Ran, W. Li, X. Yan, Y. Zhao, and G. Su, Phys. Rev. B 89, 054426 (2014).
43. Z.-Y. Xie, J. Chen, J.-F. Yu, X. Kong, B. Normand, and T. Xiang, Phys. Rev. X 4, 011025 (2014).
44. S. A. Parameswaran, S. L. Sondhi, and D. P. Arovas, Phys. Rev. B 79, 024408 (2009).
45. S. R. White and D. A. Huse, Phys. Rev. B 48, 3844 (1993).
46. W. Li, A. Weichselbaum, and J. von Delft, Phys. Rev. B 88, 245121 (2013).