Physics-Inspired Quantum Simulation of Resonating Valence Bond States—A Prototypical Template for a Spin-Liquid Ground State

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ABSTRACT: Spin liquids—an emergent, exotic collective phase of matter—have garnered enormous attention in recent years. While experimentally many prospective candidates have been proposed and realized, theoretically modeling real materials that display such behavior may pose serious challenges due to the inherently high correlation content of such phases. Over the last few decades, the second-quantum revolution has been the harbinger of a novel computational paradigm capable of initiating a foundational evolution in computational physics. In this report, we strive to use the power of the latter to study a prototypical model, a spin-1/2-unit cell of a Kagome antiferromagnet. Extended lattices of such unit cells are known to possess a magnetically disordered spin-liquid ground state. We employ robust classical numerical techniques such as the density-matrix renormalization group (DMRG) to identify the nature of the ground state through a matrix-product state (MPS) formulation. We subsequently use the gained insight to construct an auxiliary Hamiltonian with reduced measurables and also design an ansatz that is modular and gate-efficient. With robust error-mitigation strategies, we are able to demonstrate that the said ansatz is capable of accurately representing the target ground state even on a real IBMQ backend within 1% accuracy in energy. Since the protocol is linearly scaling $O(n)$ in the number of unit cells, gate requirements, and the number of measurements, it is straightforwardly extendable to larger Kagome lattices that can pave the way for efficient construction of spin-liquid ground states on a quantum device.

INTRODUCTION

Quantum spin liquids are exotic phases of matter that arise in systems of interacting quantum spins. Unlike conventional magnets, where the spins align in a well-defined pattern at low temperatures, spin liquids do not exhibit long-range magnetic order, even at absolute zero. Instead, they are characterized by a unique combination of long-range quantum entanglement, fractionalized excitations, and emergent gauge fields. The strong entanglement between spins over long distances is responsible for the global topological order observed in spin liquids, which is distinct from the broken symmetries seen in conventional ordered states.

Spin liquids can have either a gapless or a gapped excitation spectrum. Gapped spin liquids are well-characterized by the global topological structure of their ground-state wave functions. They exhibit emergent quasi-particle excitations that have nontrivial statistical interactions, for instance, anionic statistics in two dimensions. Gapless spin liquids, on the other hand, are more challenging to characterize as their ground states can be highly degenerate, and even quasi-particle description completely breaks down in some cases. Gapless spin liquids exhibit power-law decay of correlation profiles of observable quantities.

Among various spin-liquid systems, the antiferromagnetic Heisenberg model on the Kagome lattice holds a particular significance. The two-dimensional lattice consisting of hexagons with corner-sharing triangles (see Figure 1) introduces strong geometric frustration that leads to intriguing physics. Numerical simulations have suggested the existence of various spin-liquid states on the Kagome lattice, including the resonating valence bond (RVB) state and the $Z_2$ spin liquid. Such RVB states were first investigated by Pauling for describing $\pi$-bonded organic molecules and then subsequently used by Anderson in connection to Mott insulators. For extended lattices, the presence of an RVB ground state usually implies that the valence bond pairs are itinerant and can delocalize throughout the lattice.

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Experimental studies have identified several materials as potential hosts of spin-liquid behavior on the Kagome lattice. For instance, Herbertsmithite (ZnCu$_3$(OH)$_2$Cl$_2$), a mineral compound with copper ions arranged in a Kagome lattice, has shown signatures of spin-liquid behavior.\textsuperscript{10–12} Another mineral is Barlowite (Cu$_6$(OH)$_3$FBr), which exhibits a Kagome lattice formed by copper ions.\textsuperscript{13} Furthermore, Na$_3$Ir$_2$O$_6$ is a material where the Ir ions, having a spin-$1/2$, reside on a Hyperkagome 3D lattice. Studies suggest that the ground state of the Heisenberg model on this lattice may exhibit a quantum spin liquid phase with a spinon Fermi surface.\textsuperscript{14} Such spinon excitations arise due to itinerant unpaired electrons not participating in the ground-state bonding framework. The list extends to other polycrystallites like (CsX)Cu$_3$O$_2$(PO$_4$)$_2$\textsuperscript{15} and even in Tb$_3$Sb$_3$Mg$_3$O$_{14}$, where each of Tb$^{3+}$ and Sb$^{5+}$ have alternating Kagome sublattices\textsuperscript{16} or in other pyrochlore derivatives like in La$_2$Sb$_3$Zn$_3$O$_{14}$\textsuperscript{17,18}. It is important to note that the formation of a perfect Kagome lattice in natural materials can be challenging due to factors like lattice distortions, impurities, and disorder. Despite the emergence of promising candidates, experimentally confirming the existence of a pristine spin-liquid material remains a significant challenge.

Despite the progress in experimental investigations, theoretical modeling of Kagome spin liquids remains challenging due to the strong correlations involved. In this paper, we focus on the theoretical investigation of a spin-1/2 antiferromagnetic Heisenberg model on the Kagome lattice. A potential initial step in understanding the Kagome lattice involves preparing its ground state and subsequently conducting investigations using the prepared ground state. This article focuses on achieving the ground state of Kagome lattices using a variational quantum eigensolver (VQE). Such quantum simulations, variational and otherwise, have become quite popular in recent years and been performed for a variety of other systems using diverse methodologies.\textsuperscript{19–29} However, even constructing the ground state of a single plaquette within the Kagome lattice requires 12 qubits. Various studies\textsuperscript{30,31} have demonstrated that when attempting to attain the ground state of Hamiltonians with a significant number of qubits (>8), any Hamiltonian-agnostic ansatz, beyond a certain depth, inevitably leads to a barren plateau. This barren plateau phenomenon is attributed to the excessive expressibility of the ansatz.\textsuperscript{32–34} To circumvent the issue, it is necessary to employ an ansatz that possesses just enough expressibility to capture the system’s ground state, without being overly expressive for other arbitrary states.

In this study, we develop a physics-inspired analogue that is capable of capturing the ground state of a single plaquette system. In order to achieve this, we generate two-point correlation profiles between various pairs of lattice sites using the density-matrix renormalization group (DMRG), which is computationally cheaper than exact diagonalization. We choose an ansatz that also conforms to the same correlation profile based on which pairs of lattice sites exhibit a substantial correlation. One can even go one step further to pick ansatz that has the same sign of the correlations as the ones obtained via DMRG. Furthermore, we also define two auxiliary Hamiltonians, which can be treated as reduced versions of the original Hamiltonian. Through appropriate classical preprocessing, we show that the ground state of the auxiliary Hamiltonians is identical to the ground state of the target Hamiltonian. By utilizing the reduced Hamiltonian, we can achieve certain benefits when simulating the real device compared to using the original Hamiltonian. Specifically, the reduced Hamiltonian significantly diminishes the number of gates required for computing expectations of the Hamiltonian multiple times during the optimization process. This reduction in the number of measurements enhances the performance of the VQE within the present era of noisy intermediate-scale quantum (NISQ) technologies. Naturally, all our results will be validated by performing calculations resulting in high-fidelity ground states on actual quantum hardware.

Even though there exists another work that attempts to perform a digital quantum simulation of the Kagome unit cell using Hamiltonian variational ansatz,\textsuperscript{35} however unlike us, the said report does not perform computations on an actual hardware but merely provides results using noise models on a simulator. Also being inspired from a classical preprocessing based on a thorough DMRG analysis, the approach we follow and the ansatz we use is quite different from them. The modular two-qubit ansatz that ref 35 employs requires the use of two single-qubit gates and one $\sqrt{\text{Sim}}$ gate that in turn requires a SWAP gate to be implemented (see Section IV in ref 35). Each such SWAP gate would need on an average three CNOT gates for implementation (sequential arrangement of such CNOT gates will lead to some fortunate cancellation, which motivates the discussion of an average case). As we shall see later, our ansatz is also modular with two-qubit connectivity requiring just a single CNOT gate. This ensures crisp convergence to the ground state even on the real hardware with a statistical error being $\leq$1%. In addition, as we shall discuss, our approach would also linearly scale in the number of such two-qubit gates even for an extended Kagome lattice with multiple plaquettes, making it efficient and less susceptible to noise injected from NISQ platforms. We shall also present a systematic analysis for the choice of optimizers, initial parametrization, and error mitigation strategies to use that is not explored elsewhere. Analogue simulation of Kagome ground state has also been undertaken using the 219-atom programmable quantum simulator.\textsuperscript{36} In this approach, arrays of $^{87}$Rb atoms were placed...
on the edges of a Kagome lattice. Such atoms are excitable to a high-energy Rydberg state, thereby turning on the dipolar interactions with neighboring vertices and effectively mimicking the dimer bonds in a RVB. Rydberg blockade naturally limits the density of such dimeric bonds on the atomic assembly at an appropriate filling fraction. The assembly was adiabatically time-evolved to create quantum states that inhabit the span of several paired valence bond configurations with no local order. The onset of a quantum spin-liquid phase was detected by using both diagonal and off-diagonal string operators. Since the present work is a digital simulation, it is naturally apparent that our approach is entirely different from the aforesaid attempt but shall act as a complementary recipe.

In the following section, we investigate the Kagome unit cell classically using DMRG as a precursor for choosing a suitable ansatz. Thereafter, we discuss the choice of ansatz and methods to define an auxiliary Hamiltonian to afford measurement reduction. We then present a discussion on the choice of optimizers, initial parametrization, and error-mitigation strategies that shall be used and present the results on the actual hardware. We conclude thereafter discussing the extendability of our ansatz for larger systems and using recent experimental efforts to study myriad applications of RVB in physics and chemistry even beyond the precincets of spin liquids. This highlights the importance of our investigation.

## METHODS

**Model.** An antiferromagnetic Kagome lattice consists of several tessellated unit cells, each of which can be described as being composed of qubits interacting according to the following Heisenberg Hamiltonian with homogeneous interactions (XXX)

$$H(J) = \sum_{\langle i,j \rangle} J(X_i Y_j + Y_i Z_j + Z_i X_j)$$

(1)

where \((X, Y, Z)\) are usual Pauli operators and the indices \((i, j)\) refer to respective sites/qubits. The set \(\eta\) is defined according to the adjacency matrix (connectivity pattern) of the interaction graph illustrated in Figure 1. A pair of qubits (say \((k, m)\)) sharing a specified edge in the unit cell shown in Figure 1 contributes a \(J(X_k X_m + Y_k Y_m + Z_k Z_m)\). The connectivity graph we followed in this work defines the set \(\eta\) as

\[
\eta = \{(0, 1), (1, 2), (2, 3), (3, 4), (4, 5), (5, 0), (0, 6), \\
(1, 6), (1, 7), (2, 7), (2, 8), (3, 8), (3, 9), (4, 9), \\
(4, 10), (5, 10), (5, 11), (0, 11)\}
\]

(2)

The model can be interpreted as consisting of a flip–flop term \((X_i X_j + Y_i Y_j)\) that contributes to off-diagonal entries in the configuration basis (eigenbasis of \(\sum Z_i\)) by coupling configurations that have the same number of overall excitations but only differ in the orientation of spins at the \(i\)th and \(j\)th sites only. The remaining \(Z_i Z_j\) term contributes to diagonal entries and is responsible for primarily making the model antiferromagnetic given that the overall scale of the interaction strength \(J \geq 0\). The model possesses a set of discrete \(Z_i\) symmetries over reflection across the planes \(xy, yz,\) and \(xz\) that keeps the Hamiltonian defined in eq 1 invariant under transformation of \(X_i \rightarrow -X_i, Y_i \rightarrow -Y_i,\) and \(Z_i \rightarrow -Z_i \forall i\). In addition, the model also has a continuous \(U(1)\) that leads to particle number conservation making \(\sum Z_i\) commute with the Hamiltonian in eq 1. In other words, the Hamiltonian matrix in the configuration basis would be blocked wherein within each block one shall only have configurations that belong to a particular eigenspace of \(\sum Z_i\) tagged by a given eigenvalue. All such configurations have fixed number of excitations and the dimension of such a subspace having \(k\) excitations would be \(\chi^k\) where \(N\) is the total number of spins. We shall see that the ground state that we are interested in would belong to \(\langle \sum Z_i \rangle = 0\) sector.

**Classical Analysis: Motivation for the Choice of Ansatz.** To motivate the choice of a reasonable ansatz that can reduce the computational time of actual quantum hardware without compromising accuracy, we shall first study the unit cell classically. The technique of choice is DMRG using a matrix-product state ansatz. DMRG was first introduced in ref 37 as a method for studying one-dimensional quantum systems. The method has been routinely used ever since for analyzing strongly correlated electronic as well as spin systems with high accuracy. The algorithm is based on the idea of truncating the Hilbert space of the system while preserving its essential properties. The specific version of the algorithm we use represents the target state of the system as a matrix-product state (MPS). In such an MPS ansatz, each component of the state of the system is written as a tensor network comprising the product of individual tensors defined at a local site in the system. The bond indices among the tensors are contracted while keeping the physical indices free. Mathematically, an MPS can be described as a sum of the product of matrices

\[
|\psi\rangle = \sum_{\{\tau\}} \sum_{\alpha} \Lambda_{\tau_1}^{0} \Lambda_{\tau_2}^{0} ... \Lambda_{\tau_{\chi-1}}^{0} \Lambda_{\tau_{\chi-1}}^{0} |s_1, s_2 ... s_\chi\rangle
\]

where \(\Lambda_{\tau}\) are complex square matrices of order \(\chi\) known as the bond dimension. \(\alpha\) are the bond indices and \(s\) are the physical indices. The bond dimension of the local tensor quantifies the entanglement between the sites of the system. MPS with fixed bond dimension \(\chi\) can approximate a quantum state residing in a \(N\)-qubit Hilbert space using only \(O(N\chi^2)\) parameters. One can then use this ansatz and minimize the energy of the system by iteratively optimizing the expectation value of the Hamiltonian until a desired level of accuracy is reached.

Unlike in 1D, even though DMRG in 2D has an unfavorable scaling over the width of the lattice, a number of techniques have been developed over recent years to ameliorate the issues for practical usage as described in refs 38,39. This has made the algorithm more favorable than many other popularly used methods in condensed matter physics like analytical treatments using Bethe ansatz, conformal field theory (both of which are not extendable beyond 1D), and quantum Monte Carlo-based methods that often suffer from nonnegativity of numerically obtained probability values due to the infamous sign problem. Indeed, some of the benchmark studies on potential suspects of 2D spin liquids stabilized by anisotropy,40 multispin interactions16 were initiated using DMRG. In fact, DMRG has been used to study antiferromagnet Kagome lattices (an extended version of the exact system we are interested in)40,41 quite fruitfully.

Motivated by such developments, we employ the algorithm to classically study the unit cell as described above. We use the Julia version of the ITensor library for all computations and employ an MPS with a maximum bond dimension of 200 and a maximum number of sweeps kept at 40 during the execution of the protocol. Arnold’s method is used for diagonalizing the
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Figure 2. (a) Energy obtained from DMRG and exact diagonalization (ED) as described in the text for the four lowest-energy eigenstates of Hamiltonian in eq 1. There are two degenerate ground states at $-18.0 J$ that are replicated by DMRG with near exact accuracy (states are marked as ground state 1 and ground state 2). (b) Energy error obtained from DMRG and exact diagonalization (ED) as described in the text for the four lowest-energy eigenstates of Hamiltonian in eq 1. There are two degenerate ground states at $-18.0 J$ that are replicated by DMRG with near-exact accuracy (states are marked as ground state 1 and ground state 2). (c) $(\sum Z_i)$ from the two ground states (see panel (a) or (b)) obtained from DMRG and ED. Both the states belong to the $(\sum Z_i) = 0$ eigenvalue sector.

reduced density matrices on the local Hilbert spaces. The spectrum of the latter was subsequently used to truncate the wave function MPS. To assess the quality of the numerical computation, we compare all results with exact diagonalization (note that exact diagonalization is not required for our protocol, and we just use it as a benchmark to show that our DMRG results are in great agreement. For larger systems, such a comparison need not be done).

We compute the four lowest-energy eigenstates of each from both methods and display the resulting energies in Figure 2a,b. We see for all four states the energy error is less than 2% of the exact values, with the error being near zero for the two ground states in which we are interested. In addition, in Figure 2c, we plot the $(\sum Z_i)$ value for the two ground states as obtained from DMRG and ED and show that within numerical precision, both the ground states inhabit the $(\sum Z_i) = 0$ eigenvalue sector.

Next, to assess the structural attributes of the MPS wave function and the exactly diagonalized one, we compute the Neel correlation function defined as follows

$$\langle S_\alpha^\alpha J_i S_\alpha^\alpha J_i \rangle \forall \alpha \in \{x, y, z\}, (k, i) \in \eta$$

where $S_\alpha$ are the spin matrices defined in terms of Pauli operators $(X, Y, Z)$ as

$$S_x = \frac{X}{2}, S_y = \frac{Y}{2}, S_z = \frac{Z}{2}$$

We display the results in Figure 3a,b for $\alpha = z$, i.e., with $S_z$ operators for each of the two ground states seen in Figure 2. We see astonishing anticorrelation between specific pair of spins that are $(0,6)/(6,0), (1,7)/(7,1), (2,8)/(8,2), (3,9)/(9,3), (4,10)/(10,4),$ and $(5,11)/(11,5)$ in the first ground state from DMRG and between pairs $(0,11)/(11,0), (1,6)/(6,1), (2,7)/(7,2), (3,8)/(8,3), (4,9)/(9,4),$ and $(5,10)/(10,5)$ in the second ground state. It must be emphasized that for each reference spin $k$, we see a spike in the Neel correlation for $i = k$ too, which is exactly 0.25. This is expected as $S_z^2 = \frac{1}{4} I$. The orthogonality of the two obtained ground states dictates the correlation profiles of each peak on different sites, leading to two sets of pairs as mentioned above. The states from ED for both the ground states seem to be such a superposition of the two degenerate ground states from the ones obtained from DMRG and hence show anticorrelation for both sets of pairs above. The profiles for $\langle S_x^\alpha J_i S_x^\alpha J_i \rangle$ and $\langle S_y^\alpha J_i S_y^\alpha J_i \rangle$ are similar to the ones for $S_z$ due to the directional symmetry in the Hamiltonian in eq 1. Based on this correlation profile, we are inclined to conclude that a valid representation of the ground state of the target Hamiltonian will be one wherein specific pair of spins have highly nontrivial interaction leading to anticorrelation.

We then move on to analyze the mutual information $I(i,j)$ that is defined as follows

$$I(i,j) = S(\rho(i)) + S(\rho(j)) - S(\rho(i,j))$$

where $(i,j)$ denotes a pair in the set $\eta$ (in eq 2) and the entropy is defined as $S(X) = -\text{Tr}(X \ln(X))$ where X is the reduced density matrix of the corresponding qubits. We designate the pairs in eq 2 using the letters $A$–$R$ for convenience of representation. This encoding is defined as

$$A = (0, 1), B = (1, 2), C = (2, 3), D = (3, 4), E = (4, 5), F = (5, 0), G = (0, 6), H = (1, 6), I = (1, 7), J = (2, 7), K = (2, 8), L = (3, 8), M = (3, 9), N = (4, 9), O = (4, 10), P = (5, 10), Q = (5, 11), R = (0, 11)$$

The results from the two ground-state MPS of DMRG calculations are displayed in Figure 3c,d as a function of a list of qubit pairs (see eq 5). We see that for the first ground state, pairs like $G = (0,6), I = (1,7), K = (2,8), M = (3,9), O = (4,10),$ and $Q = (5,11)$ show unusually high mutual information but all other pairs have mutual information on zero. This is only possible if in this ground state (Gr-state-1) all other pairs except those above have factorizable reduced states but the two qubits in each of the pairs $G, I, K, M, O,$ and $Q$ have inherently quantum-correlated joint state $\rho(i,j)$ that is not factorizable (i.e., the correlation is such that the entropies of the one-particle reduced state of each...
A qubit in the pair cannot reproduce the total joint entropy. A similar conclusion can be drawn for the other ground state (Gr-state-2) too but for pairs $H$, $J$, $L$, $N$, $P$, and $R$ (see eq 5) for a definition of the pairs. Lastly, in Figure 3e,f, we plot $\sum_{i \in D} Z_i$ where the set $D = G, I, K, M, O$, and $Q$ is for the first ground state (see Figure 3e) and $D = H, J, L, N, P$, and $R$ is for the second.
isoenergetic ground states even though their excited states differ. (b) Energy error of the ground-state energy values of $H_{VBS-0}(J)$ and $H_{VBS-1}(J)$ from the reduced density matrix ground state (see Figure 3f). This is done by computing the Figure 4.

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(a) Energy values of the four lowest-energy eigenstates of $H_{VBS-0}(J)$, $H_{VBS-1}(J)$, and the full $H(J)$ in eq 1. All three Hamiltonians have isoenergetic ground states even though their excited states differ. But this alone does not prove that they share the exact same ground state (it only establishes all three ground states are isoenergetic). (c) To prove this, we use the MPS of the ground state of $H_{VBS-1}(J)$ and $H_{VBS-0}(J)$ and compute its energy with respect to the full Hamiltonian in eq 1, i.e., compute $\langle \psi_{VBS-0}(H(f)) | VBS \rangle$ and $\langle \psi_{VBS-1}(H(f)) | VBS \rangle$ where $H(f)$ is the full Hamiltonian defined in eq 1 and $\psi$ is the ground state of $H_{VBS-1}(J)$ and $H_{VBS-0}(J)$. We obtain the same energy value as before. This establishes that the ground states of $H_{VBS-1}(J)$ and $H_{VBS-0}(J)$ act as a valid representation of the ground state of $H(J)$ in eq 1. (d) We resolve each of the ground state of $H_{VBS-0}(J)$ and $H_{VBS-1}(J)$ in the basis of the eigenstates (both DMRG/ED) of the Hamiltonian in eq 1. We see for each the probability adds up to 1, indicating that $\psi_{VBS-0}$ and $\psi_{VBS-1}$ live in the span of the ground state of $H(J)$ in eq 1.

$H_{VBS-0}(J) = \sum_{(i,j) \in \xi} J(X_iX_j + Y_iY_j + Z_iZ_j)$

$H_{VBS-1}(J) = \sum_{(i,j) \in \xi} J(X_iX_j + Y_iY_j + Z_iZ_j)$

where the two sets $\kappa_1$ and $\kappa_2$ consist of pair of spins defined as follows

$\kappa_1 = \{G: (0, 6), I: (1, 7), K: (2, 8), M: (3, 9), O : (4, 10), Q: (5, 11)\}$

$\kappa_2 = \{R: (0, 11), H: (1, 6), J: (2, 7), L: (3, 8), N : (4, 9), P: (5, 10)\}$

We compute the four lowest-energy eigenstates of both $H_{VBS-0}(J)$ and $H_{VBS-1}(J)$ and compare the results with ED and DMRG calculations of the full system in eq 1 in Figure 4a. We see that even though the two auxiliary Hamiltonians $H_{VBS-0}(J)$ and $H_{VBS-1}(J)$ differ from the full system in terms of excited-state energy values, the ground-state energy values of $H_{VBS-0}(J)$ and $H_{VBS-1}(J)$ surprisingly match with the ground-state energy value of the Hamiltonian in eq 1. This does not yet prove that all of these Hamiltonians have the same ground state; it only shows that $H_{VBS-0}(J)$ and $H_{VBS-1}(J)$ have an isoenergetic ground state with the Hamiltonian in eq 1. We verify this again: in Figure 4b, we display the energy error of $H_{VBS-1}(J)$ and $H_{VBS-0}(J)$ from the energy of Hamiltonian in eq 1, indicating that the errors are actually quite low. In Figure 4c, we use the MPS of the ground state of $H_{VBS-1}(J)$ and $H_{VBS-0}(J)$ and compute its energy with respect to the full Hamiltonian in eq 1, i.e., compute $\langle \psi_{VBS-0}(H(f)) | VBS \rangle$ and $\langle \psi_{VBS-1}(H(f)) | VBS \rangle$ where $H(J)$ is the full Hamiltonian defined in eq 1 and $\psi$ is the ground state of $H_{VBS-1}(J)$ and $H_{VBS-0}(J)$. We see that $\psi_{VBS-0}$ and $\psi_{VBS-1}$ also act

$\psi_{VBS-1}(H(f))$ and $\psi_{VBS-0}(H(f))$.
interacting pairs like \( (i, j), (k, l), (m, n) \in \kappa_i \). The gadget implements a Givens unitary rotation capable of generating a state in the single-excitation subspace, i.e., in the span of \(|10\rangle\) and \(|01\rangle\). The elemental gate decomposition of the gadget is given alongside. The number of unique parameters \((\theta, \phi)\) is 2 due to symmetry in the desired state as discussed before. (b) Complete 12-qubit ansatz as used for the VQE calculations in this work. The two-qubit gadget \( U(\theta, \phi) \) in this case again acts only between interacting pairs like \((i, j), (k, l), (m, n) \in \kappa_i \). The gadget comprises a single-qubit U_{1} gate and a CNOT gate controlled on state \( |0\rangle \) of the control qubit. The unique parameter count is 2 and these parameters are denoted as \((\theta, \phi)\). They collectively define the operations of the \( U_{1} \) gate in each couplet. The CNOT in each pair creates a correlated two-qubit state in the span of \(|10\rangle\) and \(|01\rangle\). We discuss about such interacting couplets in the next section specifically. Note that this ansatz consists of 1 CNOT gate per interacting pair that makes it more economical and less susceptible to gate infidelities compared to panel (a), thereby justifying its choice.

Figure 5. (a) Prospective choice of an ansatz that consists of a two-qubit gadget \( U(\theta, \phi) \) between pairs like \((i, j), (k, l), (m, n) \in \kappa_i \). The gadget safely prepare the ground states of \( H_{\text{VBS-1}}(J) \) and \( H_{\text{VBS-0}}(J) \) and claim that those are ground states of the target Hamiltonian defined in eq 1. As established from the DMRG calculations, the ground states of \( H_{\text{VBS-1}}(J) \) and \( H_{\text{VBS-0}}(J) \) are known to be singlet pairs (also, the ground state of eq 1 possesses properties satisfying this as seen in the correlation profile before) with even a subspace \( U(1) \) symmetry of a single excitation within each interacting pair, and we propose to use either of two prospective ansatze as described in Figure 5a,b. Each of the two ansatze has a modular structure with exclusive connectivity between the following interacting pairs of spins

\[
\kappa_i = \{ G: (0, 6), I: (1, 7), K: (2, 8), M: (3, 9), O: (4, 10), Q: (5, 11) \}
\]

and each of which can prepare the ground state of \( H_{\text{VBS-0}}(J) \).

Both the prospective candidates prepare a state \( \psi(\theta, \phi) \) between each such interacting pair described as \( \psi(\theta, \phi) = \cos(\theta/2)01 + \sin(\theta/2) \exp(i\phi)|10\rangle \) in the above list. The joint many-body ground state will be Kronecker product of the singlet pairs represented by two-qubit states \( \psi(\theta, \phi) \). The ansatze in Figure 5a does this by starting from the initial configuration of \( |01\rangle \) that is generated by the use of the first X gate (little endian ordering is followed) in the inset of Figure 5a. Once such a configuration is created in the single-excitation subspace, the two-qubit gadget \( U(\theta, \phi) \) for this ansatze thereafter implements a Givens unitary \( (G(\theta, \phi)) \) defined as follows

\[
\begin{align*}
U(\theta, \phi) &= X \tau(\phi) \tau(\theta) X, \\
\tau(\theta) &= e^{i\theta/2}, \\
\tau(\phi) &= e^{i\phi/2}. 
\end{align*}
\]
Thus, the overall unitary is $U(\theta, \phi) = (X \otimes I)G(\theta, \phi)$ that creates the desired superposition. However, as is seen in the inset of Figure 5a wherein the full elemental decomposition of $G(\theta, \phi)$ is explicated, the ansatz requires 3 CNOT gates per interacting pair that enhances the likelihood of the implementation to be error-prone on a near-term device. Contrary to this popular choice, we present in this manuscript another ansatz that is far more economical. Like the one above, this ansatz also comprises of a pairwise interacting gadget $U(\theta, \phi)$ that only couples pair of spins $(p, q) \in \mathcal{N}$. The unique parameter count as before will be 2 and denoted $(\theta, \phi)$. It is kept the same for all pairs due to similar correlation profiles, mutual information values, and symmetry of each pair (see Figure 3). However, unlike before, the ansatz in Figure 5b (see inset) implements the target unitary as $U(\theta, \phi) = (U_z(\theta, \phi, 0) \otimes I)(X \otimes I)\text{CNOT}(X \otimes I)$, thereby requiring just 1 two-qubit gate ($\text{CNOT}$) for each pair. The single-qubit $U_z(\theta, \phi, 0)$ gate creates the superposition of the control qubit, thereby leading to the formation of a state in the span of $|00\rangle$ and $|10\rangle$. The $\text{CNOT}$ gate being controlled on the state $|10\rangle$ thereby exclusively transforms $|00\rangle \rightarrow |10\rangle$. To use this scheme, one must emphasize that it is mandatory to initialize the qubits in $|00\rangle^{\otimes N}$. Henceforth, we shall use this ansatz for all computations in the manuscript.

**Choice of the Optimizer.** To analyze the performance of various optimizers, we performed numerous calculations on the `ibmq-qasm` simulator by importing noise models (choice of the noise model is due to FakeHanoi backend) using several available optimizers. The code base to perform all quantum simulations is implemented using Qiskit.\textsuperscript{49} Figure 6 shows the convergence of the energy as a function of the epochs (iteration index), and it can be seen that most optimizers perform well on this ansatz. We see that while most optimizers perform reasonably well and eventually attain the desired accuracy, COBYLA provides the fastest convergence within 10–20 iterations. This has been corroborated multiple times for this specific case with different initial parameters. Since enhanced number of iterations on a quantum device would incur more error due to increased number of operations and decoherence of the qubit register, we stick to using COBYLA as the preferred optimizer for all cases henceforth.

### RESULTS AND DISCUSSION

#### Initial Parameters and Error-Mitigation Schemes Using a Single Interacting Component of the Ansatz: A Two-Qubit System.**

Our analysis in the Classical Analysis: Motivation for the Choice of Ansatz section revealed anticorrelation between two adjacent lattice sites inhabiting the triangular facets of the Kagome unit cell (see Figure 1). Also, the pinwheel structure of the Kagome lattice unit cell has $C_6$ symmetry. Therefore, starting with the anticorrelated two-qubit system can provide a good foundation for comprehending the ground state of the 12-qubit Kagome lattice. Based on this finding for a two-qubit system, we propose an ansatz given by $\psi(\theta, \phi) = \cos(\theta/2)|01\rangle + \sin(\theta/2)\exp(i\phi)|10\rangle$. The corresponding ansatz can be implemented onto a quantum circuit just by using a $U_z(\theta, \phi, 0)$ gate, and a CNOT gate as shown in Figure 7.

![Figure 7. Two-Qubit ansatz used within the VQE calculations.](Image)

Intuition about the performance of this two-qubit ansatz can be beneficial for achieving convergence in the overall optimization algorithm for all 12 qubits, particularly when operating on real-world quantum devices, which are often noisy. We shall therefore use this two-qubit ansatz to study what kind of initial parameters are to be used for warm starting on a real quantum device for the 12-qubit case. We will also study what should be the optimal choice of the error-mitigation schemes. Incorporating such effective error-mitigation techniques can significantly improve the accuracy of the final results. The two-qubit Hamiltonian for any one interacting pair (say $(0,6)$th qubit pair) is given by

$$\hat{H} = X_0X_1 + Y_0Y_1 + Z_0Z_1$$

(11)

where we have relabeled the dummy indices. The ground state of this Hamiltonian is variationally obtained by minimizing the parameters $(\theta, \phi)$ within the ansatz described in Figure 7. To do this, we also use several different choices of error-mitigation techniques by evaluating their effectiveness in our specific case against the imported noise model (choice of the noise model is due to FakeHanoi backend) in the `ibmq-qasm` simulator. These schemes include zero-noise extrapolation (ZNE),\textsuperscript{50} Pauli Twirling (popularly called T-Rex),\textsuperscript{51} and measurement error mitigation (MEM),\textsuperscript{52} which are the commonly used methods to mitigate errors in quantum computing. The code base to perform all such simulations with and without error-mitigation schemes is implemented using Qiskit.\textsuperscript{49} Various factors influence the performance of each of these techniques when it is employed to attenuate the effect of noise, including but not limited to the following. For instance, ZNE can become increasingly complex as the number of qubits grows as well as the

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**Figure 6. Performance of different optimizers when VQE computations are performed for the full 12-qubit ansatz by importing the noise model (FakeHanoi backend) on the `ibmq-qasm` simulator.**

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fidelity of quantum gates is high as it requires running the same circuit multiple times with different levels of noise. In contrast, Pauli twirling can be applied more efficiently as it involves preparing a more robust single noisy state that can be used for multiple operations. Furthermore, the MEM method builds a calibration matrix that is shown in Figure 8a. The size of this calibration matrix grows exponentially, which therefore can be depending on the choice of optimizer used in the optimization process that in turn drastically affects the quality of convergence.3

We use the two-qubit Hamiltonian defined in eq 11 and optimize the ansatz in Figure 7 with respect to the variational parameters using a noise model imported from the FakeHanoi backend. The corresponding results for the energy function and the relative % energy error vs the epochs when using MEM are shown in Figure 8b,c. As can be seen in Figure 8c, the % error for the case of the FakeHanoi backend without considering any error-mitigation scheme is around 5–10%. When we use the calibration matrix from MEM, the error is mitigated and comes down to around 1–2% (Figure 8c) but not below that. On the other hand, if ZNE is used, from the results as illustrated in Figure 8d,e, we see that it is possible to reach even below 1% error in the obtained final energy. Thus, MEM was unable to reduce the error percentage as much as ZNE could do on the noisy model, and thus it will not be selected for executing VQE for the full 12-qubit system on an actual quantum device as we shall explore next. From the results of the computation with ZNE, the final converged parameters that minimized the energy for the two-qubit case were found to be \( \theta \sim \pi/2 \) and \( \phi \sim \pi \), which is essentially a Bell state of the kind \( \frac{1}{\sqrt{2}} (|01\rangle - |10\rangle) \), thereby further corroborating our previous findings of having anti-correlated pairs. We use this set of parameters as an initial warm start for the full 12-qubit case, which we investigate next.

Simulations on a Real Quantum Hardware. In this section, we show the results corresponding to the VQE calculation for finding the ground-state energy of the 12-qubit Kagome lattice on the ibm_hanoi that is one of the IBM Quantum Canary Processors. The code base to perform all such simulations on the real hardware with and without error-mitigation schemes is implemented using Qiskit.39 Based on the analysis of the aforementioned results on quantum devices, it is evident that for good accuracy, one needs to have an intuition of the starting step motivated by the physics of the problem. We use the ansatz as motivated in the Classical Analysis: Motivation for the Choice of Ansatz section and warm start the optimization by using initial parameters as discussed in the previous section. We fixed the choice of the optimizer as COBYLA since its performance is the best, as discussed. Also, we performed the optimization using the two error-mitigation schemes: Pauli–Twirling (T-Rex) and ZNE. The ground-state energy obtained upon optimization using the above specifications is well within the desired accuracy of relative energy error 1% on the ibm_hanoi device. ibm_hanoi is a 27-qubit device with the median CNOT error of 7.504 \( \times 10^{-3} \), median readout error of 1.150 \( \times 10^{-2} \), median CNOT error of 148.74 \( \mu s \), and median T2 error of 116.04 \( \mu s \) as per the latest calibration statistics.

We performed three runs using the T-Rex error-mitigation scheme and three runs using the ZNE scheme (Figure 9a–f). The results displayed in the figure show the convergence of energy as a function of the epochs. As can be seen in the figure, near the end of the optimization process, the converged energy is

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**Figure 8.** (a) Calibration matrix for all of the four configurations of the two-qubit example (see Figure 7). (b) Plots of energy, as a function of the number of epochs using a noise model coming from the FakeHanoi backend, with and without measurement error mitigation (MEM). (c) Same as in panel (b), except the dashed lines indicate % energy error = ±1%. (d) Plots of energy, as a function of the number of epochs using a noise model coming from the FakeHanoi backend, with and without zero-noise extrapolation (ZNE). (e) Same as in panel (d), except the y-axis is relative % energy error in energy with respect to the true energy of \( E_0 = -3J \), especially focusing on the last few iterations. The red dashed lines indicate % energy error = ±1%. Unlike in panel (c) where MEM could barely attain an accuracy within ±1% error range, we see ZNE results are below the same quite effortlessly. All results are obtained by importing the said noise model on the ibmq-qasm simulator with COBYLA as the optimizer.
very close to $-18J$ that is the desired value. Also, we show in each of the insets of Figure 9a–f the relative energy error computed using the difference between the true energy $E_0 = -18J$ and the obtained ground-state energy over the last few iterations, and the plots confirm that the accuracy of our optimization protocol is well within the relative error of 1% limit for most cases (the said limit is denoted using red dashed lines in insets of Figure 9a–f).

We summarize the results in Table 1 and also calculate the mean
energy and standard deviation using the last 15 iterations of the optimization process. The converged set of parameters \((\theta, \phi)\) correspond to the last iteration step, and the relative energy error \(\%\) using the mean energy in the third column is also presented.

### CONCLUSIONS

In this report, we have shown the construction of a physically inspired ansatz for the quantum simulation of a resonance valence bond state (RVB) that is the ground state of a Kagome unit cell, commonly prescribed as a candidate for a gapped spin liquid. The ansatz discussed in this report heavily relies on the fact that the DMRG calculations on the said unit cell showed significant two-point correlations only for spins present on the sites in the outer triangles of a Kagome unit cell. Through suitable analysis, it was also shown that it is possible to construct a ground-state function that has a significant and negative correlation only for the adjacent sites on the outer triangles. Using the said information, we defined two auxiliary Hamiltonians with reduced measurements whose ground state is identical to the system of interest. Equipped with such an operator and an ansatz, we performed a thorough analysis of the choice of optimizer required for a variational computation, the choice of initial state, and the error-mitigation techniques to be employed. Lastly unlike in a previous report, we validate all our inferences by showing the results of computations within 1\% error in energy even on a real quantum device *ibm_hanoi*.

The ansatz we designed can be generalized to larger Kagome lattices naturally. This could be attributed to the existence of long-range order in kagome Heisenberg antiferromagnets as discussed in ref 44. The authors of ref 44 consider a kagome lattice as a set of stars with 12 spins arranged in a triangular lattice as in Figure 10. The authors showed that the interaction between the stars leads to a band of singlet excitations and can be considered a perturbation in the low-energy sector. Extensive work exists in the literature confirming the resonating valence bond state nature of such ground states. To quantify the scalability, consider a larger lattice with \(N\) qubits. Then, one can construct \(N/12\) disjoint stars (since each qubit is part of exactly one star). So, our final ansatz would require \(O(N/12)\) single and two-qubit gates at constant depth \((O(1))\). This is definitely advantageous compared to a generic ansatz (with larger depth) that does not exploit the hidden long-range order in the Kagome lattices or even the ansatz proposed in ref 35 which uses a specialized \(f_{Sim}\) gate implementable using a SWAP gate (requiring three CNOT gates on average for each bond) unlike ours that requires just 1 CNOT gate per bond. There is also an advantage in terms of the unique parameter count (in our case, it is just 2 for each unit cell) and also auxiliary Hamiltonians with reduced number of measurables (see Auxiliary Hamiltonians with Reduced Measurements section) and reduced number of shots, thereby reducing statistical error. The simplification we used for the reduction in the unique parameter count based on symmetry arguments can be adopted for other larger lattices with other symmetries depending on the topology of the lattice (for example, looks at cylindrical and torus topology out of many) and also for simulating correlated states in other problems like exciton condensates and simultaneous Fermion-exciton condensates that has been recently realized.

Since our method entails preparing RVBs in real quantum devices with high fidelity, this holds significance across multiple areas of physics and chemistry, even beyond the domain of quantum spin liquids. This is because the ground-state preparation carried out in this paper would serve as a starting step for various studies one could conduct on systems that support RVBs as their ground states and be an impetus for areas in physical chemistry wherein they feature extensively. In physical chemistry, RVBs arose fairly early due to Pauling in the theory of conjugated electrons in \(\pi\)-bonded organic frameworks. Thereafter, it has been used to qualitatively understand bonding

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**Table 1. Summary of the Results from Actual Hardware *ibm_hanoi*\(^*\)**

<table>
<thead>
<tr>
<th>device (error-mitigation scheme)</th>
<th>job description</th>
<th>(E_{avg}/J)</th>
<th>(\sigma_{E_{avg}}/J)</th>
<th>(E_{\text{avg}}-E_0) %</th>
<th>parameters ((\theta \pm \sigma_\theta, \phi \pm \sigma_\phi))</th>
</tr>
</thead>
<tbody>
<tr>
<td>IBM Hanoi (T-Rex)</td>
<td>run 1 (Figure 9a)</td>
<td>-18.0216</td>
<td>0.1307</td>
<td>0.12</td>
<td>((1.607 \pm 0.004, 3.141 \pm 0.003))</td>
</tr>
<tr>
<td></td>
<td>run 2 (Figure 9b)</td>
<td>-17.7963</td>
<td>0.1591</td>
<td>1.13</td>
<td>((1.844 \pm 0.004, 3.083 \pm 0.002))</td>
</tr>
<tr>
<td></td>
<td>run 3 (Figure 9c)</td>
<td>-17.9182</td>
<td>0.1427</td>
<td>0.45</td>
<td>((1.509 \pm 0.003, 2.984 \pm 0.004))</td>
</tr>
<tr>
<td>IBM Hanoi (ZNE)</td>
<td>run 1 (Figure 9d)</td>
<td>-17.8233</td>
<td>0.2394</td>
<td>0.98</td>
<td>((1.644 \pm 0.004, 3.092 \pm 0.005))</td>
</tr>
<tr>
<td></td>
<td>run 2 (Figure 9e)</td>
<td>-17.8996</td>
<td>0.2266</td>
<td>0.56</td>
<td>((1.549 \pm 0.002, 2.974 \pm 0.003))</td>
</tr>
<tr>
<td></td>
<td>run 3 (Figure 9f)</td>
<td>-17.8257</td>
<td>0.2267</td>
<td>0.97</td>
<td>((1.598 \pm 0.003, 3.236 \pm 0.004))</td>
</tr>
</tbody>
</table>

\(^*E_{avg}/J\) and \(\sigma_{E_{avg}}/J\) are the mean energy and the associated standard deviation, respectively, computed using the last 15 iterations of each run to account for fluctuations in self-convergence (see inset of Figure 9a-f). The relative energy error \(\%\) in the fifth column is computed using the \(E_{avg}\) in the third column and the true energy value \(E_0 = -18J\). The converged set of unique parameters \((\theta, \phi)\) is also averaged over the same set of iterations and presented in radians along with their respective standard deviations \((\sigma_\theta, \sigma_\phi)\) defining error ranges as \(\pm \sigma_\theta/2\) and \(\pm \sigma_\phi/2\). The exact values are \((\tilde{\theta} = \pi/2, \tilde{\phi} = \pi)\) as discussed for noiseless simulations.

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**Figure 10.** Pinwheel structure of several Kagome unit cells. The ansatz that we picked satisfies the \(C_4\) symmetry and also matches the spin correlations obtained for the ground state of this lattice, thereby making it extendable to handle larger Kagome lattices too.
in graphene within the length scale of few atomic units, and also in explaining the electronic structure of recently synthesized spiro-bis(1,9-disubstituted phenalenyl)boron-based neutral radical conducting solids, which despite having high density of states at the Fermi-level needs activated conductivity (with an activation energy of 0.054 eV) owing to the possible formation of a dimeric valence-bonded ground state. Charge density phases in the benzannulated variant due to interchain interaction thereby competing with the formation of the RVB state making the latter stable at certain temperatures have also been reported. It has also been used for explaining the breakdown of the free-electron Fermi liquid theory observed from Compton scattering experiments in lithium clusters from a quantum Monte Carlo-based study that claims to show significant contribution to the cohesive energy of the cluster from electron pairing, resulting in an RVB-like state. Even in copper carbodiimide (CuNCN), the anomalous temperature dependence of spin susceptibility has been attributed to the formation of a possible RVB-like state at low temperatures, a theory that enjoys semiquantitative agreement with the experiment. In condensed matter physics, the earliest attention that RVBs enjoyed was in the celebrated work of Anderson and Senthil to partially explain the phase diagram of traditional cuprates in high-Tc superconductivity. It was later extended by Kotliar and to account for s-wave and d-wave superconducting parameters and thereafter by Baskaran to MgB$_2$. In addition, our circuit ansatz can also be used for the study of related valence bond solids with localized dimers like in the Mazumdar–Ghosh model with next to the nearest-neighbor interaction or under certain limiting conditions in a higher-dimensional Shraasy and Sutherland model. Quite recently, experimental creation of an RVB state in an artificial magnet has been reported with even single-site addressability in which Ti atom clusters are probed on a MgO surface using a scanning tunneling microscope showing evidence of dimerization. Also, instead of dimerizing to form an overall singlet, dimerization into a triplet (called tRVB) has been proposed as the basic building block in ferromagnetic strange metals like in ref 75 and in certain transition-metal dichalcogenides like 1-TaS$_2$. In light of these studies, we feel that probing the RVB structure on a quantum device should be of profound importance and a timely investigation that can benefit several disciplines.

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