Quantum phases of a frustrated spin-1 system: The 5/7 skewed ladder

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The quantum phases in a spin-1 skewed ladder system formed by alternately fusing five- and seven-membered rings are studied numerically using the exact diagonalization technique up to 16 spins and using the density matrix renormalization group method for larger system sizes. The ladder has a fixed isotropic antiferromagnetic (AF) exchange interaction (J2 = 1) between the nearest-neighbor spins along the legs and a varying isotropic AF exchange interaction (J1) along the rungs. As a function of J1, the system shows many interesting ground states (gs) which vary from different types of nonmagnetic and ferrimagnetic gs. The study of diverse gs properties such as spin gap, spin-spin correlations, spin density and bond order reveal that the system has four distinct phases, namely, the AF phase at small J1; the ferrimagnetic phase with gs spin Sgs = n for 1.44 < J1 < 4.74 and with Sgs = 2n for J1 > 5.63, where n is the number of unit cells; and a reentrant nonmagnetic phase at 4.74 < J1 < 5.44. The system also shows the presence of spin current at specific J1 values due to simultaneous breaking of both reflection and spin parity symmetries.

I. INTRODUCTION

In low-dimensional magnetic systems, confinement leads to strong quantum fluctuations, and these systems can show many exotic phases in the presence of frustration induced by the topology of exchange interactions1–21. Even in a one-dimensional (1D) spin system, with only a nearest-neighbor Heisenberg antiferromagnetic (HAF) exchange interaction, the ground state (gs) can be gapped or gapless for integer or half odd-integer spins, respectively, as pointed out in a seminal paper by Haldane22. The gs of the HAF integer spin chain can be represented as a valance bond solid (VBS)23–25, and in 1987 Affleck, Kennedy, Lieb, and Tasaki (AKLT) showed that perfect VBS state may exist on various geometries with specific spins23,24. The AKLT state still continues to inspire physicists for various reasons; for example, the AKLT state has led to many recent developments such as the matrix product states technique26–28 which is a form of the density matrix renormalization group (DMRG) method29–32, the tensor network method33 and the projected entangled pair states ansatz27,28. AKLT states can also be represented as cluster states which can be used in measurement-based quantum computation34,35, and recently these states have been explored in a spin-3/2 on a hexagonal lattice36,37.

The HAF spin-1 chain exhibits a topological phase, spin-1/2 edge modes, and the gs is four fold degenerate in the thermodynamic limit. The correlation length in the gs of spin-1 is 6.05 lattice units and the eigenvalue spectrum has large spin gaps38,39. The gs can be represented as a VBS, which belongs to the same universality class of AKLT states23,24. The two leg HAF spin-1 ladder shows interesting properties like plaquette-singlet solid state, where two spin-1/2 singlet dimers are sitting at each rung and there is no overlap between the VBS states in the large rung exchange limit40. The AKLT state in the system breaks down for any finite value of rung exchange interaction40. The spin-1 zigzag ladder shows a transition from a Haldane phase to a double Haldane phase41,42. In fact the zigzag ladder can be mapped into a chain system with nearest-neighbor and next nearest-neighbor exchange interactions, and the gs of the frustrated systems is a singlet. In this work, we explore the magnetic phases of a spin-1 system on a 5/7-skewed ladder system; it has been demonstrated that a spin-1/2 system on this lattice shows many exotic phases43.

The 5/7-skewed ladder is inspired by fused Azulene, a ladder like structure made up of 5- and 7-membered carbon rings alternately fused on a chain, studied by Thomas et al. in which they showed that the gs is ferrimagnetic44. These structures can be mapped to a zigzag ladder structure with some missing bonds43,44. The HAF spin-1/2 system on various lattices such as the 5/7, 3/4, 3/5, and 5/5 is studied and it was shown that the gs of these systems exhibits many interesting magnetic and nonmagnetic gs in their quantum phase diagrams with strength of the rung exchange interaction as a phase parameter43. In the large rung exchange limit, the gs wavefunction of a 5/7 skewed ladder can be represented as a product of rung singlet dimers and two ferromagnetically interacting spins per unit cell45. In various parameter regimes this system shows dimer, spiral and chiral vector phases43. In the presence of an axial magnetic field the HAF spin-1/2 system on the 5/7 skewed ladder exhibits four magnetization plateau phases45.

The structure of zigzag and 5/7 skewed ladder are shown in Figs. 1(a) and 1(b) and by periodically removing some of the rung bonds, shown in red, from Fig. 1(a) to give 5/7 skewed ladder in Fig. 1(b). In this paper, we are interested in the gs phases of a spin-1 5/7 skewed ladder as a function of the ratio of rung-to-leg exchanges J1 and J2, respectively. We show that this system is highly frustrated, and in the small rung interaction limit, J1/J2 < 0.06, singlet dimers along the rung are weak and correlations along the leg remain short ranged, whereas,
for $J_1/J_2 > 1.44$, the gs is magnetic and each unit cell contributes spin-1 to the gs spin $S_G$, and spin densities are distributed over the whole unit cell, with spin density at sites 3 and 7 being large. For $4.74 < J_1/J_2 < 5.44$ the system is nonmagnetic but for $J_1 > 5.63$ gs of the system is magnetic with each unit cell contributing spin 2 to $S_G$ with prominent rung dimers and site spin densities.

This paper is divided into four sections. In Section II we discuss the model Hamiltonian and the numerical methods. The results are presented and discussed in Section III under four subsections. Section IV provides a summary of results and conclusions.

II. MODEL AND METHOD

The site numbering used in this paper for the 5/7 skewed ladder is shown in Fig. 1(b). All nonzero exchange interactions between spins are antiferromagnetic (AF). The sites are numbered such that odd numbered sites are on the bottom leg and even numbered sites are on the top leg. Thus the rung bonds are the nearest-neighbor exchanges $J_1$ and the bonds on the legs are the next-nearest-neighbor exchanges $J_2$. The exchange $J_2$ is set to 1 and it defines the energy scale. The model Hamiltonian of the 5/7 skewed ladder can be written as

$$H_{5/7} = J_1 \sum_i (\vec{S}_{i,1} \cdot \vec{S}_{i,2} + \vec{S}_{i,4} \cdot \vec{S}_{i,5}) + J_2 \sum_i \left(\vec{S}_{i,7} \cdot \vec{S}_{i+1,1} + \vec{S}_{i,8} \cdot \vec{S}_{i+1,2} + \sum_{k=1}^{6} \vec{S}_{i,k} \cdot \vec{S}_{i,k+2}\right).$$

where $i$ labels the unit cell and $k$ are the spins within the unit cell (Fig. 1). The first term denotes the rung exchange terms, and the second term denotes the exchange interactions along the legs.

We use the exact diagonalization technique for finite ladders with up to 16 spins and impose periodic boundary condition (PBC). There are mirror planes perpendicular to the ladder, for example, the plane perpendicular to the ladder and passing through site 3 and the perpendicular bisector of sites 2 and 4, as well as the one passing through site 7 and the perpendicular bisector of sites 6 and 8, again perpendicular to the ladder. An extra rung is needed when the open boundary condition (OBC) is used. For larger system sizes we use the DMRG method to handle the large degrees of freedom in the many body Hamiltonian. We retain up to 500 block states ($m = 500$), which are the eigenvectors of the block density matrix with dominant eigenvalues.

The chosen value of “$m$” keeps the truncation error to less than $10^{-10}$. We also carry out 6–10 finite sweeps for improved convergence. The details of building the 5/7 ladder for the DMRG method is the same as in Ref. [43]. The largest system size studied is a system with 130 sites or 16 unit cells with OBC. The DMRG calculations are carried out for different $S^z$ values of ladders. The gs spin is $S_G = l$, for $l$ that satisfies $\Gamma_l = 0$ and $\Gamma_{l+1} > 0$, where $\Gamma_l$ is given by

$$\Gamma_l = E_0(S^z = l) - E_0(S^z = 0),$$

with $E_0$ being the lowest energy state in the chosen $S^z$ sector. The correlation function and bond orders are computed in the gs, with $S^z = S$.

III. RESULTS AND DISCUSSIONS

In the gs, the spin-1 5/7 skewed ladder, like the spin-1/2 system, also shows many exotic phases like the bond order wave (BOW) phase, chiral points in parameter space of the Hamiltonian and nonmagnetic to magnetic phase transition on tuning the value of $J_1$. However, there are significant differences from the spin-1/2 system. To analyze the magnetic transitions in the quantum phase diagram, various quantities are analyzed as a function of $J_1/J_2$ which is the only variable model parameter in this system and $J_2$ is set to 1. Besides the spin gaps $\Gamma_l$, we have the computed correlation function

![Schematic diagram of (a) a zigzag ladder. The nearest-neighbor or rung interaction is $J_1$ and the next-nearest neighbor (along the leg) interaction is $J_2$. (b) The 5/7 skewed ladder: Some rung bonds shown in red of the zigzag ladder are periodically removed to give a 5/7 skewed ladder. All nonzero exchanges are antiferromagnetic (AF). The sites are numbered such that odd numbered sites are on the bottom leg and even numbered sites are on the top leg. Thus the rung bonds are the nearest-neighbor exchanges $J_1$ and the bonds on the legs are the next-nearest-neighbor exchanges $J_2$. The exchange $J_2$ is set to 1 and it defines the energy scale. The model Hamiltonian of the 5/7 skewed ladder can be written as

$$H_{5/7} = J_1 \sum_i (\vec{S}_{i,1} \cdot \vec{S}_{i,2} + \vec{S}_{i,4} \cdot \vec{S}_{i,5}) + J_2 \sum_i \left(\vec{S}_{i,7} \cdot \vec{S}_{i+1,1} + \vec{S}_{i,8} \cdot \vec{S}_{i+1,2} + \sum_{k=1}^{6} \vec{S}_{i,k} \cdot \vec{S}_{i,k+2}\right).$$

where $i$ labels the unit cell and $k$ are the spins within the unit cell (Fig. 1). The first term denotes the rung exchange terms, and the second term denotes the exchange interactions along the legs. We use the exact diagonalization technique for finite ladders with up to 16 spins and impose periodic boundary condition (PBC). There are mirror planes perpendicular to the ladder, for example, the plane perpendicular to the ladder and passing through site 3 and the perpendicular bisector of sites 2 and 4, as well as the one passing through site 7 and the perpendicular bisector of sites 6 and 8, again perpendicular to the ladder. An extra rung is needed when the open boundary condition (OBC) is used. For larger system sizes we use the DMRG method to handle the large degrees of freedom in the many body Hamiltonian. We retain up to 500 block states ($m = 500$), which are the eigenvectors of the block density matrix with dominant eigenvalues. The chosen value of “$m$” keeps the truncation error to less than $10^{-10}$. We also carry out 6–10 finite sweeps for improved convergence. The details of building the 5/7 ladder for the DMRG method is the same as in Ref. [43]. The largest system size studied is a system with 130 sites or 16 unit cells with OBC. The DMRG calculations are carried out for different $S^z$ values of ladders. The gs spin is $S_G = l$, for $l$ that satisfies $\Gamma_l = 0$ and $\Gamma_{l+1} > 0$, where $\Gamma_l$ is given by

$$\Gamma_l = E_0(S^z = l) - E_0(S^z = 0),$$

with $E_0$ being the lowest energy state in the chosen $S^z$ sector. The correlation function and bond orders are computed in the gs, with $S^z = S$. In the gs, the spin-1 5/7 skewed ladder, like the spin-1/2 system, also shows many exotic phases like the bond order wave (BOW) phase, chiral points in parameter space of the Hamiltonian and nonmagnetic to magnetic phase transition on tuning the value of $J_1$. However, there are significant differences from the spin-1/2 system. To analyze the magnetic transitions in the quantum phase diagram, various quantities are analyzed as a function of $J_1/J_2$ which is the only variable model parameter in this system and $J_2$ is set to 1. Besides the spin gaps $\Gamma_l$, we have the computed correlation function...
\[ C(r) = \langle \vec{S}_i \cdot \vec{S}_{i+r} \rangle \]
to study the behavior of the spins in the system. The bond order between bonded neighbors 
\(-\langle \vec{S}_i \cdot \vec{S}_i \rangle\) where sites \(i\) and \(i'\) are bonded neighbors, 
and spin-density \(\langle \vec{S}_i^z \rangle\) within a unit cell are also calculated and compared with the results for a 1D spin-1 chain 
where appropriate.

### A. Nature of gs

The spin in the gs, \(S_G\), of the skewed 5/7 ladder systems is obtained from the magnetic gaps \(\Gamma_l\) defined in Eq. 2. In Fig. 2(a), we plot the gaps \(\Gamma_l\) for different values of "\(l\)”, as a function of \(J_1\) for a system with 24 spins corresponding to 3 unit cells \((n = 3)\) under PBC. The plot shows that there are four distinct regions: In region I with \(0 < J_1 < 1.06\), the gs is a singlet and is nonmagnetic; in region II, \(1.06 < J_1 < 4.74\), \(S_G\) is less than or equal to the number of unit cells in the systems, consequently each unit cell contributes at most spin-1 to \(S_G\) and for \(1.44 < J_1 < 4.74\), the spin \(S_G\) saturates to the number of unit cells. We calculate \(S_G/n\) as a function of \(J_1\) for systems with \(N = 24\) and \(N = 48\) spins to investigate if the transition to \(S_G = n\) is smooth or abrupt. We find that \(S_G/n\) shows a gradual increase in the region \(1.06 < J_1 < 1.44\) for the larger system, shown in Fig. 2(b). The weak finite size effect in \(S_G/n\) can be attributed to the short spin-spin correlation lengths. As shown in Fig. 2(b), the increase in \(S_G/n\) is continuous between regions I and II. In region III, \(4.74 < J_1 < 5.44\), the gs becomes nonmagnetic for the 24 spin system. The spin-spin correlation function reveals that in this region the “free” spins in each unit cell align ferromagnetically while the alignment of the spins across unit cells is AF, with large periodicity. The transition from region II to region III is abrupt for \(N = 24\) spins and we had convergence difficulties even for the \(N = 48\) spins and hence cannot comment on the effect of system size.

We investigated the spin gaps at \(J_1 = 5.1\) (where there is a peak in the \(\Gamma_l\) values for the \(N = 24\) spin system) as a function of system size from \(N = 24\) to \(N = 96\) spins, retaining 2400 block states in the finite DMRG calculations for both PBC and OBC. We find that for this \(J_1\) value, the gaps \(\Gamma_1\) to \(\Gamma_n\) exhibit nonlinear variation with system size as shown in Figs. 2(c) and 2(d) for PBC and OBC, respectively. The convergence for higher values of \(l\) in \(\Gamma_l\) for PBC is poor and hence is not shown in Fig. 2(c). We find that for some system sizes (number of unit cells \(n\)) the excitation gaps \(\Gamma_1\), \(\Gamma_2\) and \(\Gamma_3\) vanish, for both PBC and OBC. We surmise that the vanishing of the gaps is because for these “\(n\)”, the number of unit cells in the system is an integral multiple of the periodicity of the spin-spin correlations. In region IV, the spin of the gs is \(2n\), and indicates that all the “free” spins are ferromagnetically aligned. The transition from region III to region IV was followed for the 24 spin system under PBC, by varying \(J_1\) in small increments. We find a step in the gs spin at an intermediate value \(1 < S_G/n < 2\) with a step

![FIG. 2. (a) The lowest excitation gaps \(\Gamma_l\) for different \(S^z = l\) manifolds are shown as functions of \(J_1\). For \(J_1 < 1.06\), \(\Gamma_1\) is nonzero, whereas for \(1.06 < J_1 < 4.74\), \(\Gamma_1\) is zero for \(l \leq S\) where \(S\) is the total spin of the gs. The system exhibits reentrant nonmagnetic phase for \(4.74 < J_1 < 5.44\) where \(\Gamma_1\) is nonzero. For \(1.44 < J_1 < 4.74\) \(S_G \sim n\) and for \(J_1 > 5.63\) \(S_G \sim 2n\). (b) The variation of the gs spin per unit cell \(S_G/n\) with \(J_1\) for system sizes of 48 (\(n = 6\)) spins in regions I and II, and of 24 (\(n = 3\)) spins in all the regions. (c) The lowest excitation gaps \(\Gamma_1\), \(\Gamma_2\) and \(\Gamma_3\) with the inverse of the number of unit cells of a 5/7 with PBC are shown for \(J_1 = 5.1\). (d) The lowest excitation gaps \(\Gamma_1\) to \(\Gamma_7\) with the inverse of the number of unit cells of a 5/7 with OBC are shown for \(J_1 = 5.1\).](image-url)

![FIG. 3. The variation of the gs spin \(S_G\) with the number of unit cells \(n\) is shown for \(J_1 = 1.0, 1.9\) and 6.0.](image-url)
B. Spin correlations

To understand the spin structure in different regions of the parameter space, we have studied the spin-spin correlations of the total spin \( C(r) = \langle \vec{S}_k \cdot \vec{S}_{k+r} \rangle \), where \( k \) is the reference site of spins in the middle of the system. The \( z \) component of the spin correlations, \( C_{zz}(r) = \langle S^z_k S^z_{k+r} \rangle - \langle S^z_k \rangle \langle S^z_{k+r} \rangle \), shows behavior similar to the total spin correlation. The total spin correlations are shown for a system of 98 spins, which correspond to 12 unit cells for OBC. These are calculated in the gs with \( S^z = S_G \). There are three different spin correlations that we have computed. They correspond to the correlation between spins on the lower leg \( C_1(49, 49 + 2r) \), \( C_2(50, 50 + 2r) \) between spins on the upper leg and \( C_3(51, 51 + 4r) \) between “free” spins which reside on the lower leg. The reference site for the correlations is from the middle unit cell which for \( C_1 \) is site 49, for \( C_2 \) is site 50 and for \( C_3 \) is site 51. For convenience, we classify the spins on the lower leg as of two types, type 1 “bound” spins, which are bound to three nearest-neighbor spins and type 2 as “free” spins, which are the middle sites in the five and seven-membered rings.

In Fig. 4, we show the spin correlations in the four different regions of the parameter space. The correlations \( C_1(r) \) shown in Fig. 4(a) correspond to spins in the lower leg. The correlations are given from site 49 which is in the middle of the system. For \( J_1 = 0 \), we have the correlation length of the Haldane system. We also find that as \( J_1 \) is increased, the correlation length gradually decreases to \( \sim 2.2 \) for \( J_1 = 1.0 \). In the transition region between I and II, the correlations fluctuate rapidly and we can not extract a correlation length. We note that for \( J_1 = 1.0 \), the system has a singlet gs. The correlations fall off rapidly and the correlation length \( \xi \approx \sim 2.2 \) sites of the specified kind. In the spin-1 AF chain the correlation length is longer by almost a factor of three and is approximately six sites. The shorter correlation length can perhaps be attributed to the frustration in exchange interactions in the rings. For \( J_1 = 2.0 \) and 6.0, the system is in a magnetic state, \( C_1(r) \) decays very slowly and is AF in nature. In the reentrant phase the system goes to a nonmagnetic state and the spin correlations between the spins on the lower leg have long wavelength spin oscillations whose amplitude shows an exponential decay and corresponds to a noncollinear spin arrangement. From the spin correlations, it appears that the magnetic unit cell is tripled in this region. In Fig. 4(b), \( C_2(r) \) in the upper leg are shown for the four phases, with the reference spin being the 50th spin in the system. \( C_2(r) \) for both \( J_1 = 1 \) and 2 are AF and exponentially decaying with correlation length, \( \xi \approx 3 \). For \( J_1 = 5 \) and 6, \( C_2(r) \) are vanishingly small, the magnitude is less than \( \approx 0.05 \) even for nearest-neighbor pair, and show long wavelength behavior. However, the amplitude of this wave is too small to definitively conclude this oscillatory behavior. The small correlation in upper leg at high \( J_1 \) is due to the strong dimer formation along the rungs and between \( (i, 6) \) and \( (i, 8) \) sites. The correlations between “free” spins \( C_3(r) \) shows a rapid decay in the nonmagnetic state at \( J_1 = 1 \), while those for \( J_1 = 2 \) and \( J_1 = 6 \), the correlations are ferromagnetic. For the \( J_1 = 6 \), the spins at these sites have almost completely aligned ferromagnetically, while for \( J_1 = 2 \), the alignment is partially ferromagnetic. This reflects in the net spin of the gs which is \( 2n \) for the \( J_1 = 6 \) case and \( n \) for the \( J_1 = 2 \) case. In the reentrant phase, the free spins in each unit cell are aligned ferromagnetically while the alignment of these spins across unit cells is AF, with large periodicity.

In summary, all the nearest-neighbor spin correlations are always AF, in the nonmagnetic gs for \( J_1 \) in regime I, and the correlation lengths are much shorter than the Haldane chain. For \( J_1 \) values in regime-II where \( S_G \sim n \), the correlations in the lower leg are AF and very long ranged while those in the upper leg are AF and fall off rapidly. The “free” spin correlations are ferromagnetic with an amplitude of about \( 0.6 \) and show very slow decay. When the \( J_1 \) value is in the reentrant regime III, the lower leg spin correlations show formation of wave packets over approximately three unit cells. The “free” spin correlations show a long wavelength oscillatory behavior with about five unit cell wavelength, which corresponds to a long period Néel arrangement of “free” spins. While in the classical, frustrated \( J_1 - J_2 \) model, the pitch angle is dependent on \( J_1/J_2 \), in the skewed ladder we have not been able to obtain a similar relationship. Besides, it is unlikely that a classical model will exhibit a reentrant phase. In regime IV, the “bound” spins on the lower leg are antiferromagnetically aligned and the correlations fall off very slowly with distance. The correlations between spins on the upper leg show weak long period Néel structure. The “free” spins are aligned ferromagnetically with very long correlation length.

C. Spin densities and bond order

The correlation lengths in the system are short, often less than distance to a third equivalent nearest-neighbors. Hence, we can get qualitatively correct behavior of the system in the thermodynamic limit from high accuracy DMRG studies on a system with three unit cells. We have carried out studies on a 48 site spin-1 system, with PBC corresponding to six unit cells \( (n = 6) \). We have retained 2400 block states for high accuracy in our DMRG computations. We have computed the spin densities in the gs for \( S^z = S_G \) and the bond orders of all the nearest-neighbor bonds. The system has reflection symmetry and hence there are five unique bonds and five unique sites. The spin densities are computed as the expectation value of \( \langle \psi_{gs} \mid S^z_i \mid \psi_{gs} \rangle \). They are uniformly zero in the singlet gs. The bond orders are computed as \( b_{i,j} = -\langle \psi_{gs} \mid S_i \cdot S_j \mid \psi_{gs} \rangle \), where \( i \) and \( j \) are nearest-neighbor bonds. When the gs of the system is a singlet (region I), from the bond orders (Fig. 5) we can describe the system as weakly coupled spin-1 HAF chains. The
upper leg has a BOW with a periodicity of four bonds, while the lower leg has a BOW with a periodicity of two bonds. The rung bonds are weak and the leg bond orders vary between 1.285 and 1.370. For comparison, in the spin-1 HAF, all the bond orders are uniform and have a value of 1.40. In region II, where \( S_G = n \), the upper leg bond between the sites in the pentagon become weak, the rung bonds become strong and the bonds in the lower leg also become slightly weak. Besides, the bond on the upper leg between the sites which entirely belongs to the seven-membered ring also becomes strong. The rung bonds become much stronger while the ladder bonds become weaker. The spin densities at the “free” spin sites are nearly equal and there is a net negative spin density on the rung bonds with the spin density of the sites on the lower leg being large negative. The “bonded” spin sites on the seven-membered ring on the upper leg acquire small negative spin densities. In region IV, where the \( S_G = 2n \), the rung bonds and the bond in the upper leg of the seven-membered ring almost form singlets, with a bond order close to 2.0. All other bonds are very weak. The spin densities of the sites in the seven-membered ring which form the singlet are very nearly zero, while the rung bonds are qualitatively different with large negative (on the lower leg) and positive (on the upper leg) spin densities. The “free” spins are almost completely polarized and have spin densities that are very nearly unity.

D. Vector chirality

Broken symmetry states give rise to different quantum phases whose properties depend on the type of symmetry that is broken in the system. In general, broken spatial inversion/reflection symmetry gives rise to bond the BOW phase, whereas broken spin inversion symmetry gives rise to spin density wave (SDW). If both the spatial and spin inversion symmetries in the system are broken then the vector chiral phase arises and it leads to a spontaneous spin current in the system. For these symmetries to break simultaneously, the lowest energy states in the two subspaces that the symmetry element divides the appropriate Hilbert space should be degenerate. In this case, any linear combination of the two low-lying states in the two subspaces which are even (odd) under both reflection and spin inversion will be degenerate resulting in symmetry breaking. The symmetry group of the 5/7 skewed ladder system consists of four elements: \( E, P, \sigma \), and \( \sigma P \), where \( E \) is identity, \( \sigma \) is the reflection symmetry, and \( P \) is the spin inversion symmetry and all
In this section, we focus on the lowest energy levels belonging to different reflection symmetry subspaces. These levels are shown as a function of \( J_1 \) in Fig. 6. The energy gap \( \Gamma_\sigma \) (see Eq. 3) between two lowest energy levels belonging to different reflection symmetry subspaces. \( \Gamma_\sigma \) vanishes at \( J_1 = 1.07, 1.408, 4.601 \) and 5.550 indicating degeneracies at these \( J_1 \) values.

Since these elements commute with each other leading to an Abelian group. The four irreducible representations correspond to \( A^+, A^-, B^+, \) and \( B^- \). A \( \sigma \) space under both symmetries. The matrix elements of the spin current operator has nonzero eigenvalues. This leads to nonzero spin current in a well-defined \( S_0 \) state, but the total spin is no longer conserved in the eigenstate. This implies spontaneous symmetry breaking, i.e., the eigenstate does not exhibit the full symmetry of the Hamiltonian.

The magnitude of the \( z \) component of the spin current, \( \kappa_z(j,k) \), is given by the eigenvalues of the matrix of the spin current operator for the \( j-k \) bond, viz., \((\hat{S}_j \times \hat{S}_k)_z \) in the \(| \psi_G(\pm) \rangle \) basis. The matrix, in this basis, is given by,

\[
\begin{pmatrix}
\langle \psi(+) | \hat{r}_z & | \psi(+) \rangle \\
\langle \psi(-) | \hat{r}_z & | \psi(-) \rangle \\
\end{pmatrix} \begin{pmatrix}
\langle \psi(+) | \hat{r}_z & | \psi(-) \rangle \\
\langle \psi(-) | \hat{r}_z & | \psi(+) \rangle \\
\end{pmatrix},
\]

where the function \(| \psi_G(\pm) \rangle \) is the lowest energy state in the even subspace for reflection and even subspace for spin inversion, similarly, \(| \psi_G(-) \rangle \) is in the odd subspace under both symmetries. The matrix elements of the spin current operator can be evaluated easily by using the operator identity

\[
\kappa_z(j,k) = -i(\hat{S}_j \times \hat{S}_k)_z = \frac{1}{2}(\hat{S}^+_j \hat{S}^-_k - \hat{S}^+_k \hat{S}^-_j)
\]

The diagonal matrix element in the \( 2 \times 2 \) matrix is zero, and the eigenvalues of the spin current are given by \( \pm \frac{1}{2} \left| \langle \psi_G(+) \mid (\hat{S}^+_j \hat{S}^-_k - \hat{S}^+_k \hat{S}^-_j) \mid \psi_G(-) \rangle \right| \)

In Fig. 7, we show the spin currents for \( J_1 = 1.408 \) and 5.550 for different degenerate \( S^z \) values. Spin currents for all \( J_1 \) values, at which \( \Gamma_\sigma = 0 \) and there is a degeneracy between states of odd and even total spin, are shown in Table II for different \( S^z \) values. We find that the spin currents are large for \( J_1 = 1.408 \) compared with \( J_1 = 5.550 \). At \( J_1 = 1.408 \), the spin current is larger in the five-membered ring compared with that in the seven-membered ring. Also, the direction of spin currents is opposite in the two rings. The spin currents are also not uniform for all the bonds, implying the mean angle between the spins depends upon the bond, as the spin current of a bond is a measure of the angle between

![Graph showing \( \Gamma_\sigma \) as a function of \( J_1 \) for a system size \( N = 16 \) with PBC.](image-url)
or orientations of the spins of the bond. The rung bonds have smaller currents than the leg bonds and in the five-membered ring, the upper leg has larger currents than the lower leg, while it is the opposite in the seven-membered ring. The spin current in the five-membered ring almost vanishes at $J_1 = 5.55$ while it is much weaker in the seven-membered ring. The spin currents of the rung bonds are very small and the spin current on the leg bonds in the seven-membered ring becomes uniform. There is also weak dependence of the spin current on the $S^z$ value of the state for which it is calculated.

**IV. SUMMARY AND CONCLUSION**

In this paper, we study quantum phases of a spin-1 HAF model on a 5/7 ladder shown in Fig. 1(b). This system goes from a nonmagnetic state to a partially magnetized state for $J_1 > 1.06$, and for $J_1 > 1.44$ magnetization per unit cell is $m = S_G/n = 1$. The gs again goes to a nonmagnetic state for $4.74 < J_1 < 5.44$, and spins have noncollinear arrangement in this phase. For large $J_1 > 5.63$, the gs goes to a magnetic state with $m = 2$.

The correlation length in the 5/7 ladder decreases monotonically with $J_1$ and for $J_1 = 1$ the correlation length $\xi \sim 3$ lattice units in the singlet gs. The bond order of the rung bonds increase monotonically with $J_1$, and in the large $J_1$ limit, the gs is a product of rung dimers and free spins at sites 3 and 7 in each unit cell. The uniform VBS state of spin-1 chain disappears even for the small value of $J_1$ as spin-1/2 at the edges of the ladder gets pinned by the rung interaction and forms singlet pairs. In large $J_1$ limit, the spins at sites 8−2 and $8i$ ($i$ is the unit cell index) form a strong singlet dimer, which is comparable to a spin-1 singlet dimer with bond order $\sim 2$.

**TABLE I. Two lowest energy levels from different $S^z$ sectors at specified $J_1$ values (see Fig. 6).**

| $J_1$  | $E(S^z = 0)$ | $E(S^z = 1)$ | $E(S^z = 2)$ | $E(S^z = 3)$ | $E(S^z = 4)$ |
|--------|--------------|--------------|--------------|--------------|--------------|
| 1.07   | -23.8471     | -23.8470     | -23.8470     |              |              |
| 1.408  | -25.0575     | -25.0575     | -25.0574     |              |              |
| 4.601  | -44.7871     | -44.7871     | -44.7871     | -44.7871     |              |
| 5.55   | -51.7948     | -51.7948     | -51.7948     | -51.7948     | -51.7946     |

FIG. 7. Spin currents $\kappa_{z}(j, k)$ of a 5/7 skewed ladder of $N = 16$ spins (see Eq. 4) for (a) $J_1 = 1.408$ in the $S^z = 1$ sector, (b) $J_1 = 5.55$ in the $S^z = 2$ sector, and (c) $J_1 = 5.55$ in the $S^z = 3$ sector. Arrows indicate the direction of the current and the magnitude of the currents is given adjacent to the arrows. No arrowhead for a bond means the spin current is zero along that bond.
This system may be realized in molecular magnets based on transition metal compounds.

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**TABLE II. Spin currents for different $J_1$ values in different $S^z$ sectors.** At $J_1 = 1.07$, $S = 0$ and $S = 1$ are degenerate. At $J_1 = 1.408$, $S = 1$ and $S = 2$ are degenerate. At $J_1 = 4.601$, $S = 2$ and $S = 3$ are degenerate and finally at $J_1 = 5.5$, $S = 4$ and $S = 5$ are degenerate. $\kappa_z$ is defined in Eq. 4.

| $J_1$ | $S^z$ | $\kappa_z(1, 2)$ | $\kappa_z(4, 5)$ | $\kappa_z(1, 3)$ | $\kappa_z(5, 7)$ | $\kappa_z(2, 4)$ | $\kappa_z(4, 6)$ |
|-------|------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| 1.07  | 0    | −0.302          | −0.302          | 0.165           | −0.158          | −0.237          | 0.086           |
| 1.408 | 0    | −0.316          | −0.316          | 0.225           | −0.220          | −0.315          | 0.129           |
|       | 1    | 0.274           | 0.274           | −0.195          | 0.190           | 0.273           | −0.112          |
| 4.601 | 0    | −0.045          | 0.045           | 0               | 0.205           | 0               | −0.205          |
|       | 1    | −0.042          | 0.042           | 0               | 0.193           | 0               | −0.193          |
|       | 2    | 0.033           | −0.033          | 0               | −0.153          | 0               | 0.153           |
| 5.55  | 0    | 0.027           | −0.027          | 0               | −0.147          | 0               | 0.147           |
|       | 1    | −0.026          | 0.026           | 0               | 0.143           | 0               | −0.143          |
|       | 2    | −0.023          | 0.023           | 0               | 0.127           | 0               | −0.127          |
|       | 3    | −0.018          | 0.018           | 0               | 0.097           | 0               | −0.097          |

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