Fully symmetrized valence-bond based technique for solving exchange Hamiltonians of molecular magnets

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Generally, the first step in modeling molecular magnets involves obtaining the low-lying eigenstates of a Heisenberg exchange Hamiltonian which conserves total spin and belongs usually to a non-Abelian point group. In quantum chemistry, it has been a long-standing problem to target a state which has definite total spin and also belongs to a definite irreducible representation of the point group. Many attempts have been made over the years, but unfortunately these have not resulted in methods that are easy to implement, or even applicable to all point groups. Here we present a general technique which is a hybrid method based on valence-bond basis and the basis of the *z*-component of the total spin, which is applicable to all types of point groups and is easy to implement on a computer. We illustrate the power of the method by applying it to the molecular magnetic system, $Cu₆Fe₈$, with cubic symmetry. We emphasize that our method is applicable to spin clusters with arbitrary site spins and is easily extended to fermionic systems.

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I. INTRODUCTION

The field of molecular magnetism has witnessed an explosion in the number of systems that exhibit molecular magnetic phenomena such as quantum resonant tunneling and photomagnetism (see reviews^{1[,2](#page-6-5)}). This explosive growth has also presented challenges to theorists modeling these systems. The problems encountered by theorists begin with modeling the nature of exchange interactions between pairs of magnetic ions. While an electronic many-body Hamiltonian has to be solved for determining the nature of exchange, this is often circumvented by guessing the nature of exchange based on the knowledge of the ligands, electron configuration of the transition metal ion and the geometry of the complex. The second problem concerns obtaining the eigenstates of the exchange Hamiltonian,

$$
H_{\text{ex}} = -\sum_{\langle ij \rangle} J_{ij} \hat{S}_i \cdot \hat{S}_j,\tag{1}
$$

describing the coupling between pairs of magnetic ions *ij* with exchange constant J_{ii} . Often, the Fock space of the Hamiltonian of the magnetic system could be very large (in the case of $Mn₁₂Ac$, it is as large as a hundred million⁴ and in Fe_{12} ferric wheel, it is more than two billion³) and obtaining even a few low-lying states of the Hamiltonian could pose a challenge. Since the exchange Hamiltonian conserves both total spin and *z*-component of total spin (M_S) , the problem can be simplified by specializing the basis, in which the matrix representation of the Hamiltonian is sought, to the case of fixed *z*-component of the total spin. Further simplification could come from exploiting spatial symmetries of the model. An ideal situation would correspond to one in which all the spin and spatial symmetries are utilized to construct a fully symmetrized basis to minimize the size of the Hamiltonian matrix that needs to be diagonalized.

The conservation of the S_{tot}^z , the total *z*-component of spin is trivially achieved by choosing from the Fock space, states whose total M_S value corresponds to the desired value. This is possible since individual S_i^z operators commute with the S_{tot}^z operator. It is also quite straightforward to set up the Hamiltonian matrix in this basis and solve for a few lowlying states in cases where the Hilbert space is spanned by a few hundred million states[.30](#page-6-8) However, the exchange Hamiltonian of molecular magnet often consists of interactions that are geometrically frustrated. In such a system, the groundstate spin is often not predictable and one needs to obtain the lowest state in each total spin subspace to fix the spin of the ground state. Besides, low energy states with different total spins lie close in energy and it is numerically difficult to achieve convergence to nearly degenerate eigenstates unless they can be dispersed into orthogonal Hilbert spaces. We can partially alleviate this problem by employing the parity symmetry of the exchange Hamiltonian. This symmetry corresponds to rotation of all the spins in the system around the *x* or *y* axis by an angle π which leaves the Hamiltonian invariant in the $M_S=0$ sector.

$$
[H, R_{y}(\pi)] = 0, \quad \text{where} \quad R_{y}(\theta) = e^{-i\theta \hat{S}_{y}/\hbar}.
$$
 (2)

The action of parity operator (\hat{P}) on a basis state with site m_s values m_1 , m_2 , m_3 m_n is to flip all the spins in the system, i.e.,

 $\hat{P}|m_1m_2...m_n\rangle = (-1)^{\eta}|-m_1-m_2...-m_n\rangle,$ (3)

where

$$
\eta = \sum_i s_i.
$$

Thus, the parity operator which conserves the total M_S value, only when $M_S=0$, is a symmetry element of the Hamiltonian matrix in the $M_S=0$ sector. In the general case where the individual objects have spin s_i , if $\Sigma_i s_i$ is even then symmetric (antisymmetric) combination of the basis states,

under parity, will span a space of even (odd) total spin states. The method can be extended easily to systems which allow only half odd-integer total spin. Since in most cases, the lowest excited state usually has a spin which is one different from that of the ground state, this symmetry renders it easy to obtain the spin gap accurately. However, the size of the Hilbert space is only reduced by approximately half of the size of the full $M_S=0$ space by using this symmetry. Use of parity is still advantageous as exploiting spatial symmetries is straightforward, even when the point group is not Abelian.

Construction of spin adapted configuration state functions (CSFs) has been a problem of long-standing interest in quantum chemistry. The CSFs are simultaneous eigenstates of total $S_{tot}²$ and *S*_{tot} and setting up the Hamiltonian matrix in this basis leads to matrices of smaller size besides allowing automatic labeling of the states by the total spin. Besides, the eigenvalue spectrum is enriched, since we can obtain several low-lying states in each total spin sector. This is in contrast to obtaining several low-lying states in a given total M_S sector as the latter would have states with total spin $S_{\text{tot}} \geq M_S$. There are many ways of achieving this task.⁵ The simplest method involves setting up the matrix of the total spin operator, $S_{tot}²$, in the CSF basis of fixed M_S and obtaining the eigenstates corresponding to a given total spin value; these eigenstates which are linear combinations of the constant M_S CSFs are then the spin adapted CSFs. For large systems this method is not practical. Another method which is sometimes used is the Löwdin projection method $6,7$ $6,7$ in which a projection operator P_S , given by

$$
P_S = \Pi_{S' \neq S} [\hat{S}^2 - S'(S' + 1)], \tag{4}
$$

is used to project out all undesired states of spins *S'* except the spin *S* of interest from a given CSF. The methods that have been extensively in vogue for construction of the spin adapted CSFs are the graphical unitary group approach $(GUGA), ^{8-10}$ $(GUGA), ^{8-10}$ $(GUGA), ^{8-10}$ symmetry group graphical approach $(SGGA), ^{11}$ $(SGGA), ^{11}$ $(SGGA), ^{11}$ and the valence-bond (VB) approach. $12-15,17$ $12-15,17$ $12-15,17$ In the GUGA method, the total spin adapted CSFs are represented as Shavitt graphs or Paldus arrays and the matrix element of a term (which corresponds to a generator of the unitary group) in the Hamiltonian between any two CSFs is obtained by comparing the two arrays or graphs corresponding to the CSFs. Similarly, graphs are used to represent spin adapted CSFs in the SGGA method and rules for computing matrix elements between two CSFs of a term in the Hamiltonian have been derived (see for example reviews^{18[–20](#page-6-19)}). Using these methods it is possible to carry out large scale configuration interaction (CI) calculations. While in all these methods the total spin symmetry is fully exploited, spatial symmetry adaptation is not an easy task.¹⁸ The CSFs are each built up of several orbitals with each orbital in general transforming according to some specific irreducible representation of the point group of the system. The direct product of the irreducible representations of a general symmetry group is not a single irreducible representation of the same group. Thus it is not possible to associate an irreducible representation with a given CSF unless the point group to which the system belongs is an Abelian group. $2^{1,22}$ $2^{1,22}$ $2^{1,22}$ Otherwise, symmetry operation on a CSF leads to a linear combination of many CSFs which is in general difficult to construct. For small dimensionalities of the Hilbert spaces, matrix representation of the symmetry operators can be obtained in the space of CSFs. The projection matrix for a given irreducible representation can be constructed from these matrices and from these, the symmetry adapted CSFs.²³ However, this approach is of limited value in real large scale problems. 18 In quantum chemical literature this difficulty is bypassed by dealing with Abelian subgroups of the system's point group.^{24,[25](#page-6-24)} However, this can lead to ambiguities in assigning the irreducible representation of a state[.26](#page-6-25) Among the methods for constructing spin adapted CSFs, the VB method is the simplest and will be considered in Sec. II.

The ultimate goal of symmetry adaptation is to be able to exploit the full spatial and spin symmetries of the system, both for computational efficiency and for labeling of the state by the irreducible representation to which it belongs. In Sec. II, we give a brief introduction to the symmetrized VB approach that was developed earlier and highlight the difficulties associated with the technique.¹⁶ In Sec. III, we present a hybrid VB-constant M_S method which overcomes these difficulties. In Sec. IV, we illustrate an application of this method to a magnetic spin cluster. In Sec. V, we summarize and discuss the future prospects for the technique.

II. SYMMETRIZED VB APPROACH

Exploiting the invariance of both total spin and its *z*-component is nontrivial, since eigenstates of the S_{tot}^z operator expressed as a product of the eigenstates of all the S_i^z operators are not simultaneously eigenstates of the $S_{tot}²$ operator. The situation is further complicated by the fact that in a molecular magnet, often the spins of all the constituent magnetic centers, s_i , are not the same. In such a situation, the easiest way of constructing the spin adapted functions is the diagrammatic VB method based on modified Rumer-Pauling rules.^{12,[14](#page-6-27)} In this method, a magnetic site with a given spin " s_i " is replaced by $2s_i$ spin-half objects. To obtain a state with total spin *S* from *N* such spin-1/2 objects from all the magnetic centers, $(N-2S)$ of these spin-1/2 objects are singlet spin paired explicitly, subject to the following restrictions: (i) there should be no singlet pairing of any two spinhalf objects belonging to the same magnetic center (this ensures that the $2s_i$ objects are in a totally symmetric combination²⁹), (ii) a total of 2*S* spin-half objects are left unpaired, (iii) when all the spin-half objects are arranged at the vertices of a regular polygon with number of vertices equal to number of spin-half objects, *N*, and straight lines are drawn between spin paired vertices, there should be no intersecting lines in the resulting diagram and (iv) when all the spin-half objects are arranged on a straight line and lines are drawn between spin paired objects, these lines should not enclose any unpaired spin-1/2 object. These rules follow from the generalization of the Rumer-Pauling rules to objects with spin greater than 1/2 and total spin greater than zero. The set of diagrams which obey these rules would henceforth be called "legal" VB diagrams. Some legal VB diagrams are shown in Fig. [1.](#page-2-0) A line in the VB diagram between two

FIG. 1. Above VB diagram shows spin pairings to yield a total spin $S_{tot} = 0$ state from ten spin-1/2 objects, constituent elementary spins of two spin 1 and two spin 3/2. Its bit representation corresponds to unique integer *I*=856. The bottom VB diagram shows a $S_{tot}=1$ state, the corresponding unique integer is $I=888$.

spin-1/2 objects *i* and *j* corresponds to the state $(\alpha_i \beta_j)$ $-\beta_i \alpha_j$ / $\sqrt{2}$ where we choose α to correspond to $|\uparrow\rangle$ and β to $|\downarrow\rangle$ orientations of the spin. The phase convention assumed is that the ordinal number "*i*" is less than the ordinal number "*j*." The 2*S* spin-1/2 objects $k_1 k_2 k_3 \ldots k_{2S}$ which are left unpaired can be taken to represent the state with $M_S=S$ given by $\alpha_{k_1} \alpha_{k_2} \alpha_{k_3} \dots \alpha_{k_{2s}}$. VB states corresponding to other M_S values for this state with spin *S* can be obtained by operating the S_{tot}^- operator on the state by the required number of times. Since the exchange Hamiltonian is isotropic, each eigenstate in the spin *S* is $(2S+1)$ fold degenerate and it is sufficient to work in the subspace of the chosen M_S value. The VB state corresponding to a given diagram is a product of the states representing the constituent parts of the diagram. On a computer, a legal VB diagram of any spin can be uniquely represented by an integer of 2N bits with a one bit representing the beginning of a singlet line and a zero bit the ending of a singlet line. The unpaired spins are also represented as onebits. Figure [1](#page-2-0) also shows bit representation of typical VB diagrams.

To spatially symmetrize a VB basis, it is necessary to know the result of a symmetry operator operating on a legal VB diagram. In general, the resultant of such an operation on a legal VB diagram is an "illegal" VB diagram. An example of this is shown in Fig. [2.](#page-2-1) [16](#page-6-26) An illegal VB diagram can be decomposed into a linear combination of legal VB diagrams, a process that is computationally demanding. In practice, the

VB space is broken down into smaller invariant subspaces of the symmetry group and within each invariant space, a symmetrized linear combination of the VB basis is constructed. However, the structure of the invariant spaces is very complex and constructing disjoint invariant spaces is not simple. While constructing symmetrized VB basis, a projection operator for a given symmetry representation is employed to project the symmetrized linear combinations by acting on each of the VB states in the invariant space. While the number of linearly independent symmetry combinations of a given representation is known *a priori*, the actual linear combinations are obtained by carrying out Gram-Schmidt orthonormalization of the projected states. However, since the VB diagrams are not orthogonal the orthonormalization process is both computationally involved and time-consuming.

Furthermore, in case of molecular magnets containing magnetic ions with spin greater than half, the exchange operator between such high-spin centers also generates "illegal" VB diagrams as it involves non-nearest-neighbor exchange interactions between constituent elementary spins.¹⁴ To illustrate, exchange between a center A with say spin one and a center B with spin $3/2$, $S_A \cdot S_B$, is expressed as (s_A) $+s_{A_2}$ \cdot $(s_{B_1} + s_{B_2} + s_{B_3})$. These exchange terms operate on a VB diagram with constituent elementary spins which are nonnearest neighbors and in general generate illegal VB diagrams as per the VB rules. Decomposition of the resultant illegal VB diagrams to legal VB diagrams could present a serious bottleneck in computing the Hamiltonian matrix elements. In view of these difficulties, a fully symmetrized VB approach to solving a Heisenberg exchange Hamiltonian particularly in the context of molecular magnets is not feasible.

III. HYBRID METHOD BASED ON VB BASIS AND CONSTANT M_S BASIS

In the constant M_S basis, a basis state of an ensemble of spins s_1, s_2, \dots, s_N is represented by a direct product of the m_s states of each spin such that the total $M_s = \sum m_i$. The basis states in the constant M_S representation are orthonormal by construction. Given the definition of a line in the VB dia-

FIG. 2. The effect of operation by the C_3^1 symmetry operator about the (1,8) axis. The top left portion shows the initial and final VB diagrams with spin couplings between vertices of the cube shown as dark lines. The bottom left portion shows the same states as spin couplings between vertices of a regular octagon. The resultant is an "illegal" diagram. Decomposing the resultant to "legal" VB diagrams yields a sum of five VB diagrams shown on the right, with spin couplings between vertices on a cube.

gram, every VB diagram can be broken up into a linear combination of the constant M_S basis states. A VB diagram with *p* singlet lines gives rise to 2^p basis states in the constant M_S basis. To effect the conversion of VB diagrams to constant M_S functions, each singlet line gives two states; in one state, the site at which a singlet line begins is replaced by an α spin and the one at which it ends by a β spin with phase +1 and in the other the spins are reversed and the associated phase is -1 . Once the VB diagrams are converted to constant M_S basis states with constituent spins, it is possible to associate a *m_i* value with a composite spin, given by $m_i = (n_{i\uparrow} - n_{i\downarrow})/2$, where $n_{i\uparrow}$ is the number of up-spin halves and $n_{i\uparrow}$ is the number of down-spin halves at the site *i*. However, there is a normalization constant w_i , which follows from Clebsch-Gordan coefficients, given by

$$
w_i = \left[\frac{(2s_i)!}{(s_i + m_i)! (s_i - m_i)!} \right]^{-1/2}
$$
 (5)

for a site with spin s_i in state m_i ^{[29](#page-6-28)} We can assume without loss of generality that the M_S value of the VB diagram is also *S*. Computationally, finding the transformation of a state in the VB basis to constant M_S basis is straightforward. We initialize the coefficients in the constant M_S basis to zero. We then decompose, sequentially, each VB diagram into constant M_S states and update the coefficient of the basis state of corresponding M_S by adding to it the VB coefficient times the product of Clebsch-Gordan factors with appropriate phases. The matrix relating the VB basis states to constant M_S basis states, **C**, is a $V \times M$ matrix, where *V* is the dimensionality of the VB space and *M* that of the constant M_S space.

The construction of the projection operator for projecting all the states of a given symmetry representation in a given spin space can now be accomplished. We construct the matrix representation of a symmetry operator, \hat{R} , of the point group in the chosen spin space by operating with \hat{R} on each state in the constant M_S basis and searching for the resulting state in the list of M_S basis states. Each basis state in this representation is carried over to another basis state by a symmetry operation of the point group. Thus, the matrix \mathbf{R}_M though an $M \times M$ matrix contains only one nonzero element in each row; this makes manipulations with this matrix computationally fast. The knowledge of the C and the \mathbf{R}_M matrices gives the effect of operating by the symmetry operator \hat{R} on a VB state as a linear combination of the constant M_S basis states via the matrix $\mathbf{B}_R = \mathbf{C} \mathbf{R}_M$. The projection operator for projecting out the basis states onto a chosen irreducible representation of the point group Γ is given by

$$
P_{\Gamma} = \sum_{\hat{R}} \chi_{\Gamma}^{\text{irr}}(\hat{R})\hat{R},\tag{6}
$$

where $\chi_{\Gamma}^{\text{irr}}(\hat{R})$ is the character under the symmetry operation \hat{R} in the character table of the point group of the system.³¹ The matrix representation of P_{Γ} in the mixed VB and constant M_S basis is given by

$$
\mathbf{Q}_{\Gamma} = \sum_{R} \chi_{\Gamma}^{\text{irr}}(R) \mathbf{B}_{R},\tag{7}
$$

where \mathbf{Q}_{Γ} is a $V \times M$ matrix. However, the rows of the matrix \mathbf{Q}_{Γ} are not linearly independent, since the symmetrized basis spans a much smaller dimensional Hilbert space. The exact dimension of the Hilbert space spanned by the system in the irreducible representation Γ can be known *a priori*. The dimension of the space Γ , V_{Γ} , is given by

$$
V_{\Gamma} = (d_{\Gamma}/h) \sum_{\hat{R}} \chi(\hat{R}) \chi_{\Gamma}^{\text{irr}}(\hat{R}), \tag{8}
$$

where d_{Γ} is the dimensionality of the irreducible representation Γ , *h* is the number of elements in the point group, and $\chi(\hat{R})$ is the reducible character for the operation \hat{R} . The *V*_T $\times M$ projection matrix, P_{Γ} , of rank V_{Γ} is obtained by Gram-Schmidt orthonormalization of the rows of the matrix Q_{Γ} until V_{Γ} orthonormal rows are obtained. These V_{Γ} orthonormal rows represent the projection matrix P_{Γ} . The $M \times M$ Hamiltonian matrix, \mathbf{H}_M , is constructed in the constant M_S basis which is described elsewhere. 30 Since the basis states in this representation are orthonormal, we do not encounter the problem of "illegal" states as with the VB representation. The $V_{\Gamma} \times V_{\Gamma}$ Hamiltonian matrix in the fully symmetrized basis is given by $P_{\Gamma}H_M (P_{\Gamma})^{\dagger}$ and one could use any of the well-known full diagonalization routines to obtain the full eigenspectrum or use Davidson's algorithm to get a few lowlying states of the symmetrized block Hamiltonian in the chosen spin and symmetry subspace.

The above procedure does not lead to the smallest block of the Hamiltonian matrix, when the irreducible representation for the block is degenerate, such as the *E*, *T*, or *H* representations. In such cases, it is advantageous to work with bases that transform according to one of the components of the irreducible representation. This can be achieved by choosing an axis of quantization and projecting out basis states of the irreducible representation which are diagonal about a rotation about the quantization axes. For example, in the case of the irreducible representation that transforms as *T*, we can choose one of the C_3 axes as a quantization axis and project the basis states of the irreducible representation using $(I + C_3^1 + C_3^{-1})$ as the projection operator. This operator projects states that transform as the Y_1^0 component of the threefold degenerate irreducible tensor operator. Similarly, for the E representation, we could use any of the C_2 axes as a quantization axis and use the projection operator $(I + C_2^1)$ to project out basis states that transform as one of its components. This is equivalent to projecting out the states which transform as a given row of the irreducible representation; the latter are not listed in standard group theoretical character tables.

Computation of static properties such as spin densities and spin-spin correlation functions in the eigenstates of the Hamiltonian is rendered simple due to the orthogonality of the constant M_S basis. The site spin operators such as the *z*-component of the spin are diagonal in this basis, while other operators such as the raising and lowering operators, though not diagonal in this basis, have a very simple matrix representation. In computing various properties, the proce-

TABLE I. Dimensionalities of the total spin spaces of a system of 14 spin-1/2 objects. $D(S)$ is the dimensionality of the constant *S* basis and $D(M_S)$ is the dimensionality of the constant M_S basis.

S/M_S			∸					
D(S)	429	1001	1001	637	273	77		
$D(M_S)$	3432	3003	2002	1001	364	91	14	

dure we follow is to express the eigenstates in the unsymmetrized constant M_S basis and to compute the desired properties using this representation.

The additional steps involved in the hybrid VB-constant M_S method are (i) constructing the **C** matrix, whose ith row contains the coefficients of the constant M_S functions appearing in the *i*th VB basis function. This is a very fast step as the constant M_S states are an ordered sequence of integers and a VB state with *n* lines is a linear combination of 2^n constant M_S functions. (ii) In the hybrid approach, computation of the **C** matrix involves the matrix multiplication, $\mathbf{C}(\Sigma_R \chi_{\Gamma}^{\text{irr}}(R) \mathbf{R}_M) = \mathbf{C} \mathbf{R}'_M$. The number of arithmetic operations involved is however very small, since both **C** and \mathbf{R}'_M are sparse matrices. In both constant M_S and hybrid approaches one has to obtain the projection operator P_{Γ} by retaining only the orthogonal rows of the matrix \mathbf{R}'_M or \mathbf{Q}_Γ , respectively. Since the number of orthogonal rows in \mathbf{Q}_{Γ} is far fewer than in \mathbf{R}'_M , this step is faster in the hybrid approach than in the constant M_S approach by a factor $D(\Gamma_S)/D(\Gamma_{M_S})$, where $D(\Gamma_s)$ is the dimensionality of the space of the irreducible representation Γ with spin *S* and $D(\Gamma_{M_S})$ is similarly the dimension of the space Γ with constant M_S . This advantage is largely offset by the fact that the \mathbf{R}'_M matrix in constant M_S basis is more sparse than the Q_{Γ} matrix in the hybrid approach. Computation of the eigenvalues in the constant M_S approach is slower than in the hybrid approach, since $D(\Gamma_M_S) > D(\Gamma_S)$ for most *S*. This advantage may be partially offset because the Hamiltonian matrix in the hybrid approach is usually more dense. The memory required for the hybrid approach is not very different from that of the constant M_S approach since the matrices, though smaller in the hybrid approach, are slightly denser. The only additional array required in the hybrid approach is the storage of the **C** matrix, which is very sparse. The major advantage of the hybrid approach is that we can obtain a far richer spectrum, since we are targeting each spin sector separately, unlike in the constant M_S approach. Thus, if we can obtain, say ten states in each *S* sector of the 2*n* spin-1/2 problem, we would have $10(n+1)$ unique states compared to the constant M_S technique where many of these spin states would be repeated in different M_S sectors.

IV. APPLICATION TO Cu₆Fe₈

We have applied the above method to model the susceptibility behavior of the molecule $[(Tp)_{8}(H_2O)_{6}Cu_{6}^{II}Fe_{8}^{III}(CN)]$ 24] $(CIO_4)_4 \cdot 12H_2O \cdot 2Et_2O, ^{27}$ where Tp stands for hydrotris (pyrazolyl) borate (Fig. [3](#page-5-0)). In this molecule the Cu*II* ions as well as the Fe*III* are in spin-1/2 state. The eight Fe*III* ions are at cube corners and the six Cu*II* ions are on the outward perpendicular to the face centers of the cube. Each Cu*II* ion is connected to the four nearest Fe*III* ions via ferromagnetic exchange interactions. There are no Fe-Fe or Cu-Cu interactions. The molecule has a spin 7 ground state. This system has a very high symmetry of the cube and incorporates all the complexities that can be encountered in the application of our technique. From the susceptibility data, the strength of the exchange interaction *J* was estimated to be 30 cm^{-1} .^{[27](#page-6-30)} Because of the high symmetry of this system, we chose this as an example for applying our technique. The dimensions of the various subspaces are given in Table [I.](#page-4-0)

Using the hybrid VB-constant M_S method, we have broken down the space in each total spin sector into basis states

Г.												
$S_{\text{tot}} \rightarrow$												
$\Gamma\!\!\downarrow$	$\boldsymbol{0}$	1	2	3	$\overline{4}$	5	6	7				
A_{1g}	6	32	24	24	9	5	$\mathbf{1}$					
A_{2g}	13	15	19	8	5	θ	θ	0				
E_g	34	90	90	60	26	10	2	$\overline{0}$				
$T_{\rm 1g}$	78	165	171	99	39	6	$\overline{0}$	$\overline{0}$				
T_{2g}	66	216	186	138	54	21	3	$\overline{0}$				
A_{1u}	5	19	13	11	$\mathfrak{2}$	$\boldsymbol{0}$	$\boldsymbol{0}$	$\overline{0}$				
A_{2u}	17	20	27	15	10	$\overline{2}$	1	$\overline{0}$				
\mathcal{E}_u	36	78	84	48	20	6	θ	$\overline{0}$				
T_{1u}	105	180	219	123	66	15	6	0				
T_{2u}	69	186	168	111	42	12	$\boldsymbol{0}$	θ				

TABLE II. Dimensionalities of various subspaces of the $Cu_6^HFe_8^H$ cluster for irreducible representation

FIG. 3. Schematic of $Cu₆Fe₈$ cluster. The filled and open circles correspond to Fe and Cu (both spin-1/2) sites, respectively. The lines represent the exchange coupling between various spin sites.

that transform as different irreducible representations of the cubic point group. The dimensionalities of the various sub-spaces are shown in Table [II.](#page-4-1) The subspaces transforming as the *E* representations are broken down into subspaces of half their dimension by quantizing the system along a C_2 axis. Similarly, the subspaces transforming as the *T* representations are broken down into one third their dimensions in Table [II,](#page-4-1) by using a C_3 axis as the axis of quantization.

We have set up the Hamiltonian matrix in each of the subspaces and obtained all the eigenvalues. We have also used a constant M_s basis and using the full cubic symmetry, factored the space into various irreducible representations and obtained all the eigenvalues in each subspace. From the eigenvectors, we have computed the total spin of the state. We find a one to one correspondence to numerical accuracy, between the two sets of calculations. We have also fitted the χT vs *T* experimental plot by using the full spectrum of the Heisenberg Hamiltonian and computing²⁸

$$
\chi T = \frac{3}{8} \left[\frac{g^2 F(J,T)}{1 - zJ'F(J,T)/k_B T} \right],
$$
(9)

where we have taken the *g* factor to be 2.1, and the ferromagnetic exchange constant *J* to be 27.2 cm⁻¹. Here, χT is in units of $N\mu_B$. The function $F(J, T)$ is given by

$$
F(J,T) = \frac{\sum_{S} \sum_{M_{S}} M_{S}^{2} \exp[-E_{0}(S,M_{S})/k_{B}T]}{\sum_{S} \sum_{M_{S}} \exp[-E_{0}(S,M_{S})/k_{B}T]},
$$
 (10)

where $E_0(S, M_S)$ is the eigenvalue of the sum of the exchange Hamiltonian and the magnetic anisotropy term $DS_Z²$ and zJ' is the intermolecular exchange interaction. Here we have assumed that the molecular anisotropy is along the global *z* axis, and this term is treated as a perturbation to the exchange Hamiltonian in Eq. (1) (1) (1) . In Fig. [4,](#page-5-1) we show the fit of the experimental data to the model.

V. SUMMARY AND OUTLOOK

The problem of exploiting total spin invariance together with spatial symmetries especially of non-Abelian point

FIG. 4. Fit of the χT vs *T* plot for the Cu₆^{*II*_Re₈^{*III*} cluster. The best} fit parameters are $J=27.2$ cm⁻¹ (ferromagnetic), $zJ' =$ −0.008 cm⁻¹ (antiferromagnetic), *D*=−0.15 cm⁻¹, and *g*=2.1.

groups has been a long-standing one. While full spin symmetry adaptation can be achieved by various techniques such as the use of permutation groups, unitary groups and the VB method, the last mentioned is the easiest and provides easy chemical visualization of the basis states. The main objection to the VB technique had been that the basis is nonorthogonal and leads to nonsymmetric representation of the Hamiltonian matrix. However, with the modification of the Davidson's algorithm³² to nonsymmetric matrices by Rettrup, 33 this objection has ceased to be important. The ease with which VB states with any given total spin can be generated from objects with assorted individual spins is an advantage which far outweighs the other historical objections to the VB method.

However, even the VB basis suffers from the serious disadvantage like all other spin adapted methods when the question of full spatial symmetry adaptation comes up. The constant M_S basis methods do not suffer from this disadvantage. In this paper, we have demonstrated, how by combining the ease of spin symmetry adaptation of the VB method with the spatial symmetry exploitation of the constant M_S methods, we can devise a scheme which is fully spin and spatial symmetry adapted. This has been possible because of the ease of transformation of the VB basis to the constant M_S basis. We have demonstrated the power of the method by applying it to the exchange Hamiltonian of the molecular magnet $Cu₆Fe₈$ which has cubic symmetry. We note that the hybrid VBconstant M_S method also allows easy manipulation of the eigenstates of the Hamiltonian for computing properties. The method described here can easily be extended to fermionic systems and should provide a significant improvement for obtaining exact eigenstates of spin conserving model Hamiltonians.

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