Exchange constants and spin waves of the orbital-ordered noncollinear spinel MnV$_2$O$_4$

R. Nanguneri and S. Y. Savrasov

Phys. Rev. B 86, 085138 — Published 24 August 2012

DOI: 10.1103/PhysRevB.86.085138
Exchange constants and spin waves of the orbital ordered, non–collinear spinel MnV$_2$O$_4$

R. Nanguneri, S. Y. Savrasov

Physics Department, University of California, Davis, California 95616, USA

We study the exchange constants of MnV$_2$O$_4$ using magnetic force theorem and local spin density approximation of density functional theory supplemented with a correction due to on–site Hubbard interaction $U$. We obtain the exchanges for three different orbital orderings of the Vanadium atoms of the spinel, two sizes of trigonal distortion, and several values of Coulomb parameter $U$. We then map the exchange constants to a Heisenberg model with single–ion anisotropy and solve for the spin–wave excitations in the non–collinear, low temperature phase of the spinel. The single–ion anisotropy parameters are obtained from an atomic multiplet exact–diagonalization program, taking into effect the crystal–field splitting and the spin–orbit coupling. We find good agreement between the spin waves of one of our orbital ordered setups with previously reported experimental spin waves as determined by neutron scattering. We can therefore determine the correct orbital order from various proposals that exist in the literature.

I. INTRODUCTION

Transition metal oxides (TMO) are a class of solid–state materials that exhibit a rich variety of physical phenomena. Among them, magnetic cubic spinels AV$_2$O$_4$ have recently attracted much attention due to geometrically frustrated corner sharing tetrahedral network formed by the V atoms (also known as a pyrochlore lattice). An interesting example is represented by MnV$_2$O$_4$ which is the spinel having additional magnetic Mn ions. It exhibits an orbital ordering (OO) that occurs at finite $T$ as a thermal phase transition: At room temperature, crystalline MnV$_2$O$_4$ is a cubic paramagnet (PM) where Mn sites occupy the centers of oxygen tetrahedra (MnO$_4$ units), while V sites occupy the centers of oxygen octahedra (VO$_6$ units) which exhibit slight trigonal distortions consistent with the $Fd3m$ cubic symmetry. As $T$ is lowered there occur two phase transitions: [1] A magnetic transition at $T_F = 56$ K from the high–T PM phase to a cubic ferrimagnetic (FEM) phase, with the Mn and V moments anti–aligned; [2] followed by a second transition at $T_S = 53$ K to a tetragonal, non–collinear FEM with orbital ordering of V$^{3+}$ 3d$^2$ electrons. The orbital ordered phase is accompanied by a reduction of the V magnetic moments due to the formation of the electron orbital moment (finite orbital angular momentum). The orbital moment, $m_o \approx 0.34$, is anti–aligned with the spin moment, $m_s \approx 1.65$, giving the total moment of $m \approx 1.3$. The reduced value of V moment has been reproduced by an earlier first–principles work in Ref. [2] and is explained by the spin–orbit coupling (SOC) on the V 3d$^2$ which generally favors anti–alignment of spin and orbital angular momenta for $T$ below the energy scale of SOC.

The local tetrahedral and octahedral coordination of the Mn and V sites results in the crystal–field (CF) splitting of their 5-fold 3d orbital degeneracy. Tetrahedrally coordinated Mn has an $e_g$ lower in energy than $t_{2g}$, while the splitting is opposite for octahedrally coordinated V. Inter–electron Coulomb interactions and exchange anti–symmetry lead to Hund’s rule splitting of up and down spins, which is greater than the CF splitting. In the stoichiometric crystalline environment, Mn has an outer shell high–spin $S = 5/2$ configuration of 3d$^5$ and a valence of +2: all 5 up/spin 3d orbitals are occupied giving $L_z = 0$ (quenched total orbital moment), and the down spin ones are empty. V has a valence of +3, an outer shell configuration of 3d$^2$, and $S = 1$: in this case, 2 electrons must occupy the 3 $t_{2g}$ orbitals. In the high temperature cubic phase, these latter three are nearly degenerate, while in the low temperature tetragonal phase, where the unit cell is slightly compressed along the c–axis, the $xy$ is lowered in energy while $yz, zx$ remain degenerate. Thus, in the tetragonal (low–T) phase, one electron on V occupies the $xy$, and the second electron has the freedom to occupy either $yz$ or $zx$. Unlike Mn, the orbital angular momentum of V is not fully quenched: The partial occupation of the $yz$ and $zx$ gives an effective orbital angular momentum $L = 1$ for V. The fact that $L \neq 0$ implies that there may be non–negligible effects of SOC in the V atoms. Further, this is a hint that the $yz$ and $zx$ could form complex linear combinations of one–electron states if it happens that $L_z = \pm 1$, since only such a complex state can have a non–zero $L_z$. The freedom of the second electron of V to occupy $yz, zx$, or some linear combination of the two gives rise to the possibility of long–range orbital order in the low–T phase.

Two simple choices have been proposed for the orbital ordering in this spinel and both have been studied theoretically in mean field models. One is the Antiferro–Orbital Order (AFOO) with alternate occupation of the $yz$ and $zx$ along the c–axis, i.e.: the same orbital is occupied in the adjacent planes above and below, as shown in Fig. (1a). This order has the space–group symmetry $I4_1/a$. The second is the Ferro–Orbital Order (FOO) where the same orbital is occupied on all V–atoms, giving...
ing the space–group \(I4_1/amd\), as shown in Fig. 1(b). In the latter, if the orbital order is a complex linear combination of \(yz\) and \(zx\) there will be a non–zero orbital angular momentum and a magnetic moment associated with it. Spin–orbit coupling can stabilize the finite orbital moment, since the energy is lower for anti–parallel alignment of \(\vec{L}\) and \(\vec{S}\).

In both of the above proposals, trigonal distortion of the VO\(_6\) octahedra in the low–\(T\) phase is not taken into account, but it is known to be large in MnV\(_2\)O\(_4\) as compared to other vanadates. While a slight trigonal distortion is present even in the high–\(T\) cubic phase, there is a qualitative symmetry–lowering change and an increase in this distortion in the low–\(T\) phase which lifts the residual degeneracy between \(yz\), \(zx\), and of the \(e_g\) manifold, and combined with the tetragonal distortion results in the mixing all 5 \(3d\) orbitals. In this case, the above OO proposals are not necessarily correct as these assume degeneracy between the \(yz\), \(zx\) orbitals. This low–\(T\) trigonal distortion has indeed been observed in the previous first–principles work\(^{11}\) that used local spin density approximation (LSDA) of density functional theory (DFT)\(^{10}\) supplemented by the correction due to on–site Hubbard interaction \(U \) for correlation strengths \(U > 2 \) eV. In that work, in addition to a tetragonal relaxation (compression) along the \(c\)–axis, structural relaxation of the O positions is performed and a trigonal distortion of the VO\(_6\) octahedron with a concomitant lowering of symmetry from \(I4_1/amd\) to \(I4_1/a\) is found. By projecting the converged density onto an atomic orbital basis using so called N–th order muffin–tin orbital (NMTO) downfolding\(^{10}\) the authors of Ref. 4 find a different electron occupation order from the ones proposed above, namely, the first electron occupies the lowest energy eigenstate, and the second occupies the next higher energy eigenstate. The 3\(d\) energy eigenstates are the same on all V sites, but rotated alternatively by 45° along the \(ab\)–chains due to the staggered trigonal distortion. Thus, the same orbitals are occupied on all V sites, akin to the FOO, but nevertheless the space–group symmetry is \(I4_1/a\) expected of AFOO due to the trigonal distortion.

The low–\(T\) magnetic excitations of the compound have been mapped along high–symmetry directions using inelastic neutron scattering\(^{3,13}\). At the \(\Gamma\) point, these excitations are gapped for the acoustic modes, indicating the presence of single–ion anisotropy, which essentially occurs due to the interplay between SOC and crystal–fields\(^{13}\). In Ref. 13, the authors start with a nearest–neighbor Heisenberg Hamiltonian including the anisotropy term and calculate spin–wave spectra and corresponding eigenmodes using linear spin–wave theory (LSWT) for the non–collinear, tetragonal phase. By fitting the spectrum to inelastic neutron scattering data, they were able to determine the exchange couplings between Mn–V, V–V in \(ab\)–plane, and V–V between \(ab\)–planes along the \(c\)–axis. They find all exchanges to be AFM with the following values:

- \(J_{\text{Mn–V}} = -2.82 \) meV
- \(J_{\text{V–V}}^{ab} = -9.89 \) meV
- \(J_{\text{V–V}} = -3.08 \) meV

The authors point out the interplanar coupling between V atoms, \(J_{\text{V–V}}\), along the \(c\)–axis is unusually large for AFOO because such an alternate orbital occupation in the vertical direction would yield negligible orbital overlap, and would also be ferromagnetic (wrong sign) by the Goodenough–Kanamori rules. The alternate proposal, FOO, would be consistent with these results, but would have the wrong symmetry, \(I4_1/amd\). The symmetry group of this spinel vanadate has been established conclusively as \(I4_1/a\) by a synchrotron x–ray study\(^{15}\) which supports AFOO, but contradicts with the large value of \(J_{\text{V–V}}\). A possible resolution of this puzzle is that trigonal distortion has been ignored in these simple proposals. With trigonal distortion, we expect a more complex orbital ordering which has the requisite symmetry \(I4_1/a\).
and would give the observed (or fitted) $J_{\text{trigonal}}$ along the $c$-axis. This is exactly what has been found in the ab-initio work of Ref. [4]. Their physical picture has received some support by a recent $^{51}$V NMR work of Ref. [10] and by analytical model of Ref. [11].

In this work, we report our study of MnV$_2$O$_4$ based on the LSDA+$U$ method and using linear muffin–tin orbital (LMTO) basis set to solve the electronic structure problem.\cite{12,13} We calculate the pair–wise interatomic magnetic exchange interactions ($J$) between all magnetic atoms using linear response theory and magnetic force theorem\cite{14,15,16}, including the single–ion anisotropies ($D$) for Mn and V found by the exact diagonalization procedure\cite{17}. We then use the obtained $J$ and $D$ as parameters in a Heisenberg Hamiltonian with anisotropy to derive the spin–wave spectra in a semiclassical approximation. We explore three initial orbital ordering scenarios: [1] Antiferro, [2] Ferro, and [3] Complex ferro + SOC in the density matrix of the 3$d$ shell of V to see how they affect the obtained exchange interactions. We also performed non–collinear magnetic electronic structure calculations.

In our low-$T$ tetragonal structures, we explore the effects of two types of trigonal distortions of the VO$_6$ octahedra: A small trigonal distortion, of order 2% of the undistorted structure, with $I4_1/amd$ symmetry; and a larger trigonal distortion, of the type used in the relaxed structure of Ref. [18] (about 10% of the undistorted structure), with $I4_1/a$ symmetry. We find that the $J$'s depend on both the size of trigonal distortion and Coulomb parameter $U$; we are thus faced with a two-parameter ‘trigonal–distortion/Coulomb–$U$’ space within which to search for a good match between experimental and theoretical $J$'s. We find that SOC complex ferro–orbital order give $J$'s which best match the experimental ones for small trigonal distortion and low-$U$, and also for larger trigonal distortion and higher-$U$.

Our paper is organized as follows. We begin with a discussion of the proposed orbital orders and their electronic structures in Section II. We present our results for exchange interactions and comparisons with experiment in Section III. We end with the conclusions in Section IV.

## II. PROPOSED ORBITAL ORDERS AND THEIR ELECTRONIC STRUCTURES

We have done LSDA+$U$ calculations to model the electronic structure for all three thermodynamic phases of MnV$_2$O$_4$. We describe our results in the following subsections for the $T = 0$ phase only since this is the phase which exhibits orbital ordering and non–collinear magnetism. Our results for the other finite–$T$ phases may be found in Ref. [19]. For the magnetic phases we use the same values of $U$ and $J_H$ for both Mn and V correlated 3$d$ shells. The use of the same $U$ on Mn and V is justified because these elements have atomic numbers 25 and 23, and are thus expected to have similar interaction strengths.\cite{20} The Coulomb and exchange parameters in the solid state are generally screened, and hence reduced by a considerable amount from their bare atomic values.\cite{21} The structural parameters for all three phases are taken from experiment.\cite{22} In the cubic phase, the lattice constant is 16.0746 a.u., and in the tetragonal phase it is 16.12 a.u. with a small tetragonal distortion ratio of $\xi = 0.98$.

The non–collinear orbital ordered phase occurs when the temperature is reduced below $T_S = 53$ K. This phase transition results simultaneously in: [1] a structural transition from cubic to tetragonal; [2] the canting of V moments from a collinear ferrimagnetic (FEM) to a $q = 0$ non-collinear FEM spin order with non–zero components in the $ab$–plane; and [3] a long–range orbital order in the V $t_{2g}$ manifold. We model the electronic structure of this phase using LSDA+$U$ method with $U = 5$ eV and $J_H = 1$ eV, but starting the self–consistency loop after imposition of the initial orbital order(s) in the Hubbard–$U$ density matrix (further described below), along with tetragonal distortion and two different magnetic configurations: [1] collinear, as in the intermediate phase, and [2] non–collinear, which is in fact the correct magnetic order for this phase. The converged charge density for the low–$T$ collinear calculation was used as the initial charge density for the correct low–$T$ non-collinear calculation. The orbital order that is finally obtained after reaching the self–consistency is taken to be the correct metastable solution within this approximation and specified initial condition(s).

We initialize the V 3$d$ density matrix to a particular orbital order by specifying orbital occupation numbers in the atomic basis. This means we initially specify only the diagonal components (occupation numbers) $\langle n_{\sigma}\rangle$, $\langle n_{\sigma'}\rangle$, $\langle n_{\sigma\sigma'}\rangle$ of the density matrix for all four V atoms’ 3$d$ shells and set the off–diagonal elements to zero. The full complex density matrix in the atomic basis is $\langle n_{m\sigma,m'\sigma'}\rangle$ (where $m, m'$ and $\sigma, \sigma'$ are the 3$d$ orbital and spin indices respectively) and includes off–diagonal components as well. As a result of the electron–electron interactions, during the self–consistent cycle non–zero off–diagonal components of the density matrix develop (since the interactions mix the single–particle 3$d$ orbitals at the Hartree–Fock (HF) mean–field level). This means the true occupied orbitals are some linear combination of the atomic basis functions. After convergence is reached, the final density matrix, which is no longer diagonal in the $(m\sigma,m'\sigma')$ basis is diagonalized. The resulting eigenvectors and eigenvalues give the ‘correct’ single–particle HF wave functions and their occupation numbers respectively. In the basis of these eigenfunctions, the density matrix is once again diagonal, and its non–zero entries signify the true orbitals which are occupied within the mean–field approximation of LSDA+$U$. We are thus able to identify the orbital ordering that results after convergence is attained. We describe the final orbital orders below for the collinear magnetic solutions only, since we find that the electronic structures, and therefore the or-
rifamycin and real antiferro–orbital order of the type I (AFOO-I) tetragonal phase with a collinear spin fat bands of V $t_{2g}$ electrons are shown in Fig. 2(a), and the same for non-collinear spins in Fig. 2(b). The occupations of the $t_{2g}↑$ bands, as shown by the partial characters, reflects the converged orbital order, as well as the FEM spin configuration. We also find that imposition of orbital order opens a gap of about $E_{\text{gap}} = 1.67$ eV at the Fermi level leading to an insulator state. The qualitative features of the band structure and partial characters does not change upon canting the V moments to the non-collinear configuration: The band gap remains robust and the phase is still insulating.

Next we describe the case of large trigonal distortion with the space-group symmetry of $I4_1/a$. In this case we use $U = 4.5$ eV, $J_H = 1$ eV, along with muffin-tin sphere radii specified in Ref. 4. We start with an initial uniform orbital order in which the three $t_{2g}↑$ are equally occupied, but the two $e_g$ are almost empty. We also start with a second initial orbital order consisting of equal occupations of all five $3d−↑$ orbitals. In both cases, we found that the converged density matrices, partial DOS, and fat-bands are identical with those of the calculation with small distortion. Our charge density on the V sites are alternately rotated within and between the V chains in the ab plane, and as well, when we transform to the local trigonal coordinate system at each of the V sites, we obtain the same single-particle wavefunctions, showing that the same orbitals are occupied on each V site, but are rotated alternately by $45^\circ$ due to the trigonal distortion. Thus, both small and large trigonal distortions result in the same orbital order, ‘AFOO-I’.

### B. Anti-Ferro Orbital Order II: $I4_1/a$ symmetry

The next simplest initial order has the second $t_{2g}$ electron occupying the same real linear combination of $yz$ and $zx$ on all V sites, with equal weight for both orbitals, see Fig. 1(b). This initial order has $I4_1/amd$ symmetry, and we implement only the small trigonal distortion. The real linear combination implies that the orbital angular momentum is zero, $L = 0$. We implement this by setting the initial mean occupations: $\langle n_{xy} \rangle = 1$, $\langle n_{y^z} \rangle = \langle n_{zx} \rangle = 1/2$ (ferro OO), and the off–diagonal elements to be zero. For this setup, the initial order does not persist until convergence is reached. Instead, there are significant non-zero off–diagonal elements, on the same order as the occupied diagonal elements, in the final density matrix. Upon diagonalizing this final matrix, the orbital order we get has the second electron occupying alternately $\psi_+ = (\psi_{y^z}+\psi_{zx})/\sqrt{2}$ and $\psi_- = (\psi_{y^z}−\psi_{zx})/\sqrt{2}$ along the c-axis, which again has the same $I4_1/a$ symmetry considered in the preceding subsection. Thus, we start with an orbital order with $I4_1/amd$ symmetry, but the self–consistent solution breaks certain discrete symmetries and results in an order with $I4_1/a$ symmetry. We thus label this order ‘AFOO-II’. We note that this order is similar to the one obtained for ZnV$_2$O$_4$ using the same

### A. Anti-Ferro Orbital Order I: $I4_1/a$ symmetry

In the low–$T$ orbitally ordered phase, the tetragonal distortion occurs to break the degeneracy of the $t_{2g}$ in both V and Mn. We first describe the case of small trigonal distortion. There is no orbital freedom to place the electrons in the Mn 3$d$. In the V, the energy of $xy$ gets lowered, so the first electron occupies $xy$. The second electron then has the freedom to occupy the remaining degenerate orbitals $yz$ or $zx$. Figure 1(a) shows the initial orbital occupations with $I4_1/a$ symmetry. In this scenario, the second electron of V occupies either $yz$ or $zx$ alternately along the c-axis (antiferro OO), and the same orbital within each ab plane (EE). (Each V chain within an ab plane has the same orbital occupied.) The final converged density matrices of the V 3$d$ subspace show that the converged orbital order is not the same as the initial order, but one which is similar to that found in Ref. 4. That is, when we rotate the 3$d$ density matrix from the global tetragonal coordinate system to the local trigonal one, a rotation by $45^\circ$, we find the same set of eigenstates for all V atoms, and the lowest two of these states in energy are occupied. We label this order ‘AFOO-I’, since it preserves the $I4_1/a$ symmetry.
LSDA+U scheme

The collinear spin fat bands of V $t_{2g}$ and $e_g$ are shown in Fig. 3(a), and the same for non-collinear spins in Fig. 3(b). Qualitative features of the band structures do not change significantly between the collinear and non-collinear spin configurations. In both plots, the occupations and dispersions of the $yz$ and $zx$ bands are nearly identical since these orbitals contribute equal weights to the true orbitals, although their relative signs in the linear combinations might differ in these depending on the particular V atom. We also find an insulating band-gap, which in this case is smaller than for ‘AFOO-I.’

C. Complex Ferro Orbital Order: $I4_1/amd$ symmetry

We focus first on the case of small trigonal distortion. The last OO has one electron in $xy$ as before, and the second electron in the spherical harmonic $L_z = -1, S_z = +1/2$ state, which is a complex linear combination of $yz$ and $zx$, on all V sites. This is an initial ferro–orbital order, but with SOC switched on and non–zero orbital angular momentum. The initial density matrix configuration persists until convergence. This calculation is carried out using LSDA+$U$+SO. This scenario is also illustrated by Fig. 3(b), except that each V atom now carries a non–zero orbital angular momentum of magnitude one due to the complex linear combination; hence, there is a uniform orbital order on all V atoms with $L = 1$ in the $3d$ density matrix. The reason for choosing the opposite $z$–projections for $\vec{L}$ and $\vec{S}$ is that spin–orbit interaction lowers the energy for such a setup, compared to the case of having the same sign for both $z$–projections.

In Fig. 4(a) we present the band structure of MnV$_2$O$_4$, for the collinear magnetic configuration, with the V $t_{2g}$↑ partial characters of the SOC uniform orbital order. We find that the V $t_{2g}$↑ and $e_g$ characters are above the $E_F$ as expected. For Mn, all the $3d$↓ are below $E_F$, while the $3d$↑ are above. There is a band gap of 1.76 eV. Within LSDA+$U$ a half-metallic solution was found in Ref. 4 with only the ↑-spin bands of V atoms crossing the $E_F$ level, a result which we have also confirmed.

Our result is that inclusion of SOC in LSDA+$U$ opens a band gap, signaling a half-metal-to-insulator transition as the SO coupling parameter is switched on. Since we argue that the uniform complex ferro order is the correct orbital order based on exchange constant calculations, we predict a half-metal-to-insulator transition to occur in single crystalline MnV$_2$O$_4$ as the temperature goes below $T_S$. In Fig. 4(b) we present the corresponding band structure of the non-collinear magnetic configuration for this order, with the partial characters of V $t_{2g}$ shown. We find that the Mn atoms carry no orbital moment as expected, but the V atoms have an orbital moment $m_O = 1.03$. The spin moments are $m_s = 4.33$ for Mn, and $m_s = 1.71$ for V. Since the spin and orbital moments are antiparallel due to SOC coupling, the total moment for V is $m = 0.7$ in this phase.

When we perform the corresponding LSDA+$U$+SO calculation with an $I4_1/a$ symmetry large trigonal dis-
tortion, and $U = 4.5$ eV, $J_H = 1.0$ eV, we find that
the converged density matrices are not significantly dif-
ferent from the ones obtained with the small $J_{A1}/am\_d$
trigonal distortion, therefore, with respect to the density
matrices, the larger trigonal distortion has a minor ef-
effect. However, the trigonal distortion does seem to have
a rather large effect on the exchange interactions as de-
scribed further below. The magnetic moments with the
larger trigonal distortion are, for Mn atoms: $m_s = 4.26,$
$m_o = 0.0$ (since the orbital moment is quenched); and for
V atoms: $m_s = 1.65,$ $m_o = 0.87,$ giving a total $m = 0.78,$
similar to what we obtained with a small trigonal distor-
tion. We label the order obtained with spin-orbit cou-
pling as ‘SOC-FOO’.

III. RESULTS FOR EXCHANGE INTERACTIONS

Here we outline the spin–wave model, the ground state
spin configuration, and present the results for our calcu-
lated exchange constants $J$ and single–site anisotropy pa-
rameters $D.$ Our obtained spin wave spectra of MnV$_2$O$_4$
and comparisons with the neutron scattering experiments
are also given.

A. Spin Wave Model

The parameters of the model are: [1] the exchange
constants $J$ derived from the LSDA+$U(+SO)$ con-
verged charge densities using linear response theory and
the magnetic force theorem$^{[20]}$ and [2] the single–
ion anisotropy parameters $D$ calculated using an exact–
diagonalization atomic multiplet procedure.$^{[21]}$ We input
these parameters into the Heisenberg model Hamiltonian
with anisotropy terms, minimize the classical energy to
find the stable ground state configuration, and calculate
the spin–wave excitation spectra. The model Hamilton-
ian is:

$$H_{\text{spin}} = -\sum_{(ij)} J_{ij} \vec{S}_i \cdot \vec{S}_j - \sum_{(ik)} J_{ik} \vec{S}_i \cdot \vec{S}_k - \sum_{(il)} J_{il} \vec{S}_i \cdot \vec{S}_l - \sum_{(jk)} J_{jk} \vec{S}_j \cdot \vec{S}_k - \sum_{(jl)} J_{jl} \vec{S}_j \cdot \vec{S}_l - \sum_{(kl)} J_{kl} \vec{S}_k \cdot \vec{S}_l$$

$$- J_{\text{Mn–V}} \sum_{(pq)} (\vec{S}_p + \vec{S}_q) \cdot (\vec{S}_i + \vec{S}_j + \vec{S}_k + \vec{S}_l) - \sum_{(pq)} J_{pq} \vec{S}_p \cdot \vec{S}_q + \sum_{x=i,j,k,l,p,q} \vec{S}_x \cdot D_x \cdot \vec{S}_x. \quad (1)$$

The subscripts on the $J$ label the four inequivalent V
sublattices, i, j, k, l, and two inequivalent Mn sublattices,
p, q. The $J_{\text{Mn–V}}$ is taken outside the summation because
it has the same value for all pairs of Mn and V atoms.
All the $J$ couplings are between nearest–neighbor atoms
of two different sublattices, and each pair is counted only
once in the summation over all sites. We ignore the next–
neighbor–neighbor couplings because we found them to be
much smaller in magnitude.

B. Spin Configuration

The low–$T$ spin structure is non–collinear for V atoms,
and collinear for Mn atoms with respect to the $c$–axis.
The pyrochlore lattice on which the V atoms sit is
geometrically frustrated for nearest–neighbor isotropic
($J^{ab} = J^c$) AFM exchange. The frustrated pyrochlore
interactions mean that there could be a macroscopic
ground state degeneracy. But this frustration is par-
tially relieved in the low–$T$ phase by the presence of ad-
ditional nearest–neighbor exchange interactions with Mn
atoms, tetragonal distortion, and orbital ordering. The
last one has the effect of making the V–V AFM exchange
anisotropic: $J^{ab} \neq J^c.$ It is well–known that the orbital
or magnetic degeneracy can be lifted by the coupling of
these degrees of freedom with the lattice via the Jahn–
Teller effect.$^{[10]}$ The ground–state spin configuration se-
lected by the system in the low–$T$ phase is non–collinear
due to the combined effect of the frustration and coupling
of the V spins to Mn spins, V $t_{2g}$ orbitals, and the lattice.

FIG. 5. The $T = 0$ non–collinear spin configuration of the V
spins from Ref. [3]
In this structure, the V–atom spins develop components in the ab plane perpendicular to each other. The amount of canting away from the c-axis can be characterized by a single canting angle $\theta$. Given the values of all $J$ and $D$ in Eq. (1) one can find the angle $\theta$ as a function of $J$ and $D$ that will minimize the classical ground-state energy of the configuration (derivation given in Ref. [22]):

$$\theta = \arccos\left(-\frac{3J_{V-V}S_{Mn}}{(D_{V}^{a}-D_{V}^{b}) - 2R_{V-V}^{a} - 2R_{V-V}^{b})S_{V}}\right) . \tag{2}$$

The non–collinear spin configuration that achieves this energy minimum is shown in Figure [35]35\textsuperscript{35}.

C. Exchange Constants

In Table [I] we present the $J$ parameters that we calculate using LSDA+U(+SO) method and magnetic force theorem for the small trigonal distortions and the indicated $U$ values. In Tables [II] and [III] we present the $J$'s for the large trigonal distortion: Table [II] for ‘AFOO-I’ and Table [III] for ‘SOC-FOO’. As the method computes the exchange constants in reciprocal space, we Fourier transform them and show only nearest–neighbor exchange interactions between atoms of each sublattice. For the spinel structure, any of nearest–neighbor pairs always belongs to a different sublattice. The values of $J_{V-V}$ for no orbital order and $U = 0$ eV are the same for all V-V pairs; but when $U = 5$ eV, there is a tendency for anisotropy to develop: the in–plane $J_{V-V}$ becomes unequal from the out–of–plane $J_{V-V}$. This shows that the anisotropy in the $J_{V-V}$, and the orbital ordering which causes it, could both be interaction driven. When there is an orbital order, $J_{V-V}$ is different along the ab V chains and between the chains (along the c–axis), as expected. This is true even for the case of the uniform orbital orders, because the exchange matrix elements of the Coulomb operator will be different within the ab plane and between the planes, as can be seen from the shapes of the occupied orbitals in Fig. 1(a,b).

We have calculated the exchange constants within LSDA+U and LSDA+U+SO taking into account both small and large trigonal distortions with $I4_{1}/amd$ and $I4_{1}/a$ symmetry respectively. The larger trigonal distortion result in $J$'s that are 50% – 80% larger, with and without SOC. Thus the effect of increasing trigonal distortion keeping $U = 4.5$ eV affects the agreement with the experimental spin-waves. One explanation for why this is so could be that $U = 4.5$ eV is too small to describe the correlation effects in V. In order to check this, we also tried $U = 4.5, 5.0, 5.5, 6.0$ eV for large trigonal distortions and found that the $J$'s indeed decrease as $U$ increases, see Tables [I] [III]. By varying both the size of the trigonal distortion and the $U$, we are dealing with a two-parameter problem. Since neither the trigonal distortion nor $U$ are exactly known, we have presented our results as an exploration of the trends in $J$'s within this two-parameter space. The increase in $U$ will bring down the values of the $J$'s as they typically scale as $t_{2g}^{2}/U$ for direct exchange between the V atoms. The trends in the variation of $J$'s within the two-parameter space indicate that the $J$'s for SOC-FOO best describes the experimental spin waves for small trigonal distortion with $U = 5$ eV, Table [II] and larger trigonal distortion with $U = 6$ eV, Table [III].

D. Single–Ion Anisotropy

The calculation of single–ion anisotropy requires first the total energies of the interacting atomic shell in a crystal–field environment along with the spin–orbit coupling. The method for its computation is described in Ref. [14] and [22] so here we merely present our results. The input parameters used in the total energy calculation are: the SOC parameter 0.15 eV, and Slater integrals $F_{0} = 5.0$ eV, $F_{2} = 7.6$ eV, $F_{4} = 4.7$ eV. We then vary the direction of the magnetic moment by applying a small external magnetic field. The CF levels have the following values: the energies of $yz$ and $zx$ are both set to the reference value of 0.0 eV. The $e_{y}$ level is varied from 0.2 eV to 1.0 eV in steps of 0.2 eV; and the $xy$ level has the energies $-0.024$ eV, $-0.016$ eV, $-0.008$ eV, 0.0 eV. This gives a set of 20 different CF configurations. The $E_{xy} = E_{yz=zx}$ represents cubic CF, and $E_{xy} \neq E_{yz=zx}$ represents tetragonal CF.

The total atomic shell energies thus obtained are fitted to a parabolic function of the polar angle $\theta$ representing moment orientation, centered at $\theta = 0$ in the case of z–axis anisotropy, and centered at $\theta = \pi/2$ in the case of x/y–axis anisotropy. The results of the parabolic fit that best match the experimentally known D values are given in Table [IV] for both the V 3d$^{2}$ and Mn 3d$^{5}$ shells.

The easy axis for Mn is z(c)–axis, and for V it is either x or y. The easy axis always has a negative anisotropy parameter, which means the energy is lowered when the spin projection along the easy axis is maximized. For Mn, the spin projection along z tends to be maximized. However, V also has a positive anisotropy parameter along the z–axis. So, V spin projection like to be maximized along y, and minimized along x. Thus, the V spin moment has a tendency to be in a non–collinear direction with respect to the z–axis. Our anisotropy computation is able to reproduce these signs as well as magnitudes for Mn (3d$^{5}$) and V (3d$^{2}$) shells.

Looking at the anisotropy fit values we see that the value of Mn anisotropy reported in Ref. [13] are obtained for $E_{xy} = -0.016$ eV, $E_{yz} = 1.0$ eV, namely $D_{Mn} = -0.1123$ meV, which is similar to the literature value. For our fitted values of V anisotropy, we do not find such a close match, but there are several CF values which give the anisotropy of Ref. [13] up to the correct sign and order of magnitude. For example, $E_{xy} = -0.024$ eV, $E_{yz} = 0.4$ eV give $D_{V}^{x} = 7.34$ meV, and $D_{V}^{y} = -4.056$ meV, which can be compared to $D_{V}^{x} = 2.79$ meV, and
TABLE I. Calculated exchange constants in meV for the collinear ferrimagnetic configurations and imposing various orbital orders along with the $J_4$/amd small trigonal distortion. The last two columns list our theoretical $J$s for the spin–orbit coupled ferro–orbital order (SOC–FOO) and experimental $J$’s from Ref. 13 respectively. The experimental $J$’s match the SOC–FOO $J$’s better than for the other theoretical $J$’s.

<table>
<thead>
<tr>
<th>meV</th>
<th>No OO $U = 0$ eV</th>
<th>No OO $U = 5$ eV</th>
<th>AFOO-I $U = 5$ eV</th>
<th>AFOO-II $U = 5$ eV</th>
<th>SOC-FOO $U = 5$ eV</th>
<th>Expt. $^{13}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$J_{ab}$</td>
<td>-2.72</td>
<td>0.136</td>
<td>0.3264</td>
<td>-0.04488</td>
<td>0.1496</td>
<td>-</td>
</tr>
<tr>
<td>$J_{Vab}^{xy}$</td>
<td>-20.4</td>
<td>-21.76</td>
<td>-14.96</td>
<td>-19.04</td>
<td>-10.88</td>
<td>-9.89</td>
</tr>
<tr>
<td>$J_{Vc}^{xy}$</td>
<td>-20.4</td>
<td>-18.36</td>
<td>-3.536</td>
<td>-7.072</td>
<td>-2.72</td>
<td>-3.08</td>
</tr>
<tr>
<td>$J_{Mn-n-V}$</td>
<td>-10.2</td>
<td>-2.992</td>
<td>-5.44</td>
<td>-5.44</td>
<td>-4.76</td>
<td>-2.82</td>
</tr>
<tr>
<td>$J_{pp}$</td>
<td>1.2</td>
<td>2.167</td>
<td>2.72</td>
<td>2.72</td>
<td>2.72</td>
<td>-</td>
</tr>
<tr>
<td>$J_{pp}$</td>
<td>-0.476</td>
<td>0.204</td>
<td>0.204</td>
<td>0.272</td>
<td>0.272</td>
<td>-</td>
</tr>
</tbody>
</table>

TABLE II. Calculated exchange constants in meV for the collinear ferrimagnetic configurations with the $I4_1/a$ large trigonal distortion of the VO$_6$ octahedra for $U$=4.5, 5.0, 5.5, 6.0 eV. In all these cases, the self-consistency converges to the orbital order named ‘AFOO-I’, which is also the same orbital order found in Ref. 4. For comparison the last column lists the experimental values. $U = 6$ eV gives theoretical $J$’s which approach the experimental $J$’s.

<table>
<thead>
<tr>
<th>meV</th>
<th>$U = 4.5$ eV</th>
<th>$U = 5.0$ eV</th>
<th>$U = 5.5$ eV</th>
<th>$U = 6.0$ eV</th>
<th>Expt. $^{13}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$J_{ab}$</td>
<td>0.449</td>
<td>0.35</td>
<td>0.3</td>
<td>0.272</td>
<td>-</td>
</tr>
<tr>
<td>$J_{Vab}^{xy}$</td>
<td>-17.7</td>
<td>-14.28</td>
<td>-12.92</td>
<td>-11.56</td>
<td>-9.89</td>
</tr>
<tr>
<td>$J_{Vc}^{xy}$</td>
<td>-4.624</td>
<td>-4.352</td>
<td>-3.808</td>
<td>-3.4</td>
<td>-3.08</td>
</tr>
<tr>
<td>$J_{Mn-n-V}$</td>
<td>-6.8</td>
<td>-5.712</td>
<td>-5.304</td>
<td>-4.896</td>
<td>-2.82</td>
</tr>
<tr>
<td>$J_{pp}$</td>
<td>2.72</td>
<td>2.584</td>
<td>2.448</td>
<td>2.312</td>
<td>-</td>
</tr>
<tr>
<td>$J_{pp}$</td>
<td>0.204</td>
<td>0.2</td>
<td>0.2</td>
<td>0.1768</td>
<td>-</td>
</tr>
</tbody>
</table>

TABLE III. Calculated exchange constants in meV for the collinear ferrimagnetic configurations with the $I4_1/a$ large trigonal distortion of the VO$_6$ octahedra for $U$=4.5, 5.0, 5.5, 6.0 eV and SOC. In all these cases, the self–consistency converges to the orbital order named ‘SOC-FOO’. For comparison the last column lists the experimental values. Again, the theoretical $J$’s for $U = 6$ eV come closest to experiment.

<table>
<thead>
<tr>
<th>meV</th>
<th>$U = 4.5$ eV</th>
<th>$U = 5.0$ eV</th>
<th>$U = 5.5$ eV</th>
<th>$U = 6.0$ eV</th>
<th>Expt. $^{13}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$J_{ab}$</td>
<td>0.272</td>
<td>0.204</td>
<td>0.177</td>
<td>0.15</td>
<td>-</td>
</tr>
<tr>
<td>$J_{Vab}^{xy}$</td>
<td>-15.64</td>
<td>-12.24</td>
<td>-10.61</td>
<td>-9.11</td>
<td>-9.89</td>
</tr>
<tr>
<td>$J_{Vc}^{xy}$</td>
<td>-5.8</td>
<td>-4.352</td>
<td>-3.536</td>
<td>-2.788</td>
<td>-3.08</td>
</tr>
<tr>
<td>$J_{Mn-n-V}$</td>
<td>-6.12</td>
<td>-5.44</td>
<td>-4.896</td>
<td>-4.352</td>
<td>-2.82</td>
</tr>
<tr>
<td>$J_{pp}$</td>
<td>2.72</td>
<td>2.72</td>
<td>2.45</td>
<td>2.329</td>
<td>-</td>
</tr>
<tr>
<td>$J_{pp}$</td>
<td>0.272</td>
<td>0.272</td>
<td>0.231</td>
<td>0.231</td>
<td>-</td>
</tr>
</tbody>
</table>

TABLE IV. Table of calculated anisotropy constants for $V^{3+}$ 3$d^2$ and Mn$^{2+}$ 3$d^5$ atomic shells. The energies of $E_{xy}$ and $E_{ez}$ due to the crystal–fields are measured with respect to $E_{yz,zz} = 0$ eV.

<table>
<thead>
<tr>
<th></th>
<th>CF $E_{xy}$ eV</th>
<th>CF $E_{ez}$ eV</th>
<th>Theory meV</th>
<th>Expt. $^{13}$ meV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mn $D^z$</td>
<td>-0.016</td>
<td>1.0</td>
<td>-0.1123</td>
<td>-0.1024</td>
</tr>
<tr>
<td>V $D^x/y$</td>
<td>-0.024</td>
<td>0.4</td>
<td>-4.056</td>
<td>-4.04</td>
</tr>
<tr>
<td>V $D^z$</td>
<td>-0.024</td>
<td>0.4</td>
<td>7.34</td>
<td>2.79</td>
</tr>
</tbody>
</table>

$D_{V}^{x/y} = -4.04$ meV of Ref. $^{13}$ One reason why our calculated $D_{V}^{z}$ parameter differs by a large amount from the experimental value is because we have to tune the CF energy levels to simultaneously match two different single–ion anisotropies, and it was not possible to get them both to match the experimental $D$ values of V.

E. Spin–Wave Spectra

We developed a code to compute the linear spin wave spectra for the non–collinear spin configuration. The program takes as an input our computed values of $J$ and $D$. We first find the ground state which will in general be a non–collinear configuration with the spins pointing along the local quantization axis as given by $\theta$ in Eq. 2. We second find the Heisenberg equations of motion, and numerically diagonalize the resulting system of linear equations. The resulting spin–wave spectra are plotted in Fig. 6(a) for the ‘SOC-FOO’ uniform orbital order, along with the experimental spin waves. We find that the spin waves
obtained from the $J$ and $D$ values of the SOC ferro-orbital order with small trigonal distortion and $U = 5$ eV matches well with experiment, although other combinations of trigonal distortion and $U$ could also yield similar $J$'s. We also note that the lower two modes are due to the oscillations of Mn spins: The lower energy being the symmetric mode, and the higher energy the anti-symmetric mode. The upper four modes are oscillations of the V spins.

For comparison, we show the spin waves for the other orbital orders, also obtained with small trigonal distortion.

![Graphs showing spin waves for different orbital orders.](image)

**FIG. 6.** In all panels, the red lines are experimental and black lines are theoretical spin–waves: (a) Spin wave spectrum for $I_4/a$ spin–orbit coupled ferro–orbital order (SOC–FOO) along the high–symmetry lines of the Brillouin zone. We find an excellent match between our theoretical and previous experimental data of Ref. 13. (b) Spin–waves corresponding to the $I_4/mmd$ spin–orbit coupled ferro–orbital order with small trigonal distortion, where the second $t_{2g}$ electron occupies a complex linear combination of $|yz\rangle + i|zx\rangle$ uniformly on all V–sites. The incorporation of the low symmetry $I_4/a$ large trigonal distortion tends to increase the $J_{V-V}$’s by 50%–80%, but we find that by a reasonable increase of the Coulomb parameter to $U = 6$ eV, we can obtain $J$’s that match the experimental ones. The trend we notice is that small trigonal distortion + lower $U$ as well as large trigonal distortion + higher $U$ both give $J$’s that are close to the experimental $J$’s, however, the former case with ‘SOC–FOO’ seems to give the best match of all the combinations we have tried. The other two orbital orders, ‘AFOO–I’ and ‘AFOO–II’, do not give such a good match with experiment throughout the Brillouin zone for the same value of distortion and $U$ so these orders may be ruled out. We further note that the spin–orbit coupling plays an important role in the orbital physics of V–atoms in MnV$_2$O$_4$. This is also justified by the fact that the single–ion anisotropy is relatively high, as evidenced by the large gaps for the would–be acoustic modes at $\Gamma$.

**IV. CONCLUSION**

By theoretical computations of the interatomic exchange constants using LSDA+$U$(+SO) method, magnetic force theorem and by imposing various orbital ordering scenarios we have shown that the orbital order on the V sites of MnV$_2$O$_4$ is similar to a complex linear combination of $zx$ and $yz$ on all V sites. Our calculated spin wave spectra for this order come closest to the experimental data. Further support in evidence of the complex order is the strong single–ion anisotropy experienced by the spin moments on the V sites, as well as the reduction of the V magnetic moment in the low–$T$ phase which could not be captured by LSDA+$U$ alone. We also predict,
based on our $U = 5$ eV, orbital–ordered band–structures, that the low–$T$ phase of MnV$_2$O$_4$ is a Mott–type insulator, and that a half–metal–to–insulator transition accompanies the simultaneous orbital ordering, structural distortion, and non-collinear moment transitions at $T_S = 53$ K.

**Acknowledgements.**

The authors acknowledge useful discussions with Myung Joon Han, Rajiv Singh and Nick Curro. The work was supported by DOE SciDAC Grant No. SE-FC02-06ER25793 and by DOE Computational Material Science Network (CMSN) Grant No. DE-SC0005468.
11 For a review, see, e.g., Strong Correlations in electronic structure calculations, edited by V. I. Anisimov (Gordon and Breach Science Publishers, Amsterdam, 2000).