Exchange coupling in transition metal monoxides: Electronic structure calculations

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An ab initio study of magnetic-exchange interactions in antiferromagnetic and strongly correlated 3d transition metal monoxides is presented. Their electronic structure is calculated using the local self-interaction correction approach, implemented within the Korringa-Kohn-Rostoker band-structure method, which is based on multiple scattering theory. The Heisenberg exchange constants are calculated with the magnetic force theorem. Based on these the corresponding Néel temperatures and spin-wave dispersions are calculated. The Néel temperatures are obtained using mean-field approximation, random-phase approximation and Monte Carlo simulations. The pressure dependence of TN is investigated using exchange constants calculated for different lattice constants. All the calculated results are compared to experimental data.

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I. INTRODUCTION

In the last years there has been a general strong interest in finding materials with specific or even parametrisable magnetic properties. Such materials could be useful in the field of spintronics. A lot of the promising candidates are strongly correlated electronic systems which in many ways are still a challenge to be properly described theoretically regarding their electronic ground-state properties. On the other hand, for reliable predictions about magnetic properties of materials, it is essential to have theories describing the magnetism adequately by quantitative and qualitative means. One of these theories is the Heisenberg theory of magnetism, which we shall apply in the present paper. Its central quantities are the Heisenberg exchange constants Jij, which are of general fundamental interest. In particular they provide information about the magnetic periodicity (via their Fourier transform), the spin-wave dispersion, magnetic critical temperatures and also allow predictions on structural effects caused by magnetism.12

In this paper we concentrate on the study of the magnetic exchange interactions of transition-metal monoxides (TMOs), specifically MnO, FeO, CoO, and NiO. They are charge-transfer insulators, well known for strong correlation effects associated with the TM 3d electrons. Originating from the Anderson-type superexchange, their equilibrium magnetic structures are of the antiferromagnetic II (AFII) order, characterized by planes of opposite magnetization which are stacked in (111)-direction. Recently, a Mott transition has been observed in MnO at high pressure of about 105 GPa, in resistivity3 and x-ray spectroscopy measurements,4,5 which stimulated new theoretical studies in this high-pressure region.6,7 There already exists a large body of neutron-scattering measurements of magnetic structures and magnetic excitations in transition-metal monoxides. However, the development of new experimental techniques such as neutron powder diffraction8–10 and polarized neutron reflectivity11 has renewed interest in studying TMOs as antiferromagnetic benchmark materials. Modern neutron spectrometers operate with such a high efficiency that also high-angle diffraction experiments can be performed to unravel complex magnetic order e. g. in thin films.12

From the theory point of view conventional methods such as the local spin-density approximation (LSDA) to density-functional theory (DFT), treating electron correlations at the level of the homogeneous electron gas, fail to provide an adequate description of the electronic structure of these oxides. Over the years a number of approaches have been developed, aiming at improvements to the LSDA treatment of electron correlations, and applied to TMOs with varying degrees of success. Among them are: the LSDA+U method,13,14 GGA+U,15 self-interaction corrected (SIC)-LSDA,16–21 hybrid functionals,22,23 and finally dynamical mean-field theory.24 In general, they have improved lattice constants, band gaps and magnetic properties, some of them have also obtained good agreement with spectroscopies.

In the present paper we shall use the so-called local self-interaction correction25 (LSIC) scheme for the calculation of the electronic ground states of the TMOs. As the aim is the investigation of magnetic interactions we combine the LSIC scheme with the magnetic force theorem (MFT)26 in order to obtain the Heisenberg exchange parameters Jij.

The LSIC scheme27 is based on the implementation of the SIC-LSDA formalism16,17,25 within multiple scattering theory in the framework of the Korringa-Kohn-Rostoker (KKR) band-structure method. It was first applied to f-electron systems,27,28 but recently also to TMOs.21 Within the KKR method one calculates the Green function of the investigated system. This Green function is then straightforwardly used in the application of the MFT. This combined approach is applied for calculating exchange constants of the transition metal monoxides. The results of that are compared to the exchange constants extracted from the total-energy differences for a number of magnetic structures and mapping them...
onto a Heisenberg Hamiltonian. Most of the earlier applications of the latter approach have been based on the assumption that only the first two exchange interaction constants are nonzero. Although the present combined approach also relies on the mapping onto a Heisenberg Hamiltonian the assumptions regarding the number of nonzero exchange constants are not needed, which is advantageous to systems with reduced symmetry such as thin films and layered structures (where the justification for such an assumption is not clear from the very beginning).

Having calculated the $J_{ij}$ for the ground states of the TMOs we also calculate and discuss them as a function of external pressure for moderate values of the latter. This is mainly inspired by the recent high-pressure measurements of TMOs.3–5

Based on the calculated magnetic exchange interactions the transition temperatures can be obtained. In this paper it is done in three different ways, namely, by applying mean-field approximation (MFA), random phase approximation (RPA), and using classical Monte Carlo (MC) simulations. The respective results are then compared to those obtained from the disordered local moments (DLM) method,29 which does not involve mapping onto a Heisenberg Hamiltonian, but is based on the same ground-state electronic structure calculations as the present paper.21

The last subject we focus on is magnetic excitations. On one hand, with given $J_{ij}$, one can calculate the magnon spectrum of any material. On the other hand, measuring the latter experimentally is a direct method to examine its exchange constants. Thus, comparing calculated to experimental spin wave dispersions provides a straightforward tool for determining the accuracy of the calculated $J_{ij}$.

The present paper is organized as follows: In Sec. II the theoretical approaches for the calculation of electronic structure, exchange interactions and Néel temperatures are presented. The computational details are described in Sec. III. Section IV contains the results and discussion. The exchange parameters and the Néel temperatures are presented for the ground states of the TMOs. The paper is concluded in Sec. V.

II. THEORY

A. Electronic structure

For the electronic structure calculations of TMOs we use a multiple scattering theory-based implementation of the SIC-LSDA method,25 whose total-energy functional is

$$
E_{\text{SIC-LSDA}}[\{n_{a\sigma}\}] = E_{\text{LSDA}}[\{n_{\uparrow\downarrow}\}] - \sum_{a\sigma} (E_{\text{H}}[n_{a\sigma}])
+ E_{\text{xc}}^{\text{LSDA}}[n_{a\sigma},0],
$$

with the LSDA energy functional in units of Rydberg given by

$$
\tilde{E}_{\text{LSDA}}[n_{\uparrow\downarrow}] = \sum_{a\sigma} \frac{1}{2} \langle \phi_{a\sigma} | -\nabla^{2} | \phi_{a\sigma} \rangle + E_{\text{H}}[n_{\sigma}] + E_{\text{xc}}^{\text{LSDA}}[n_{\uparrow\downarrow}],
$$

Here $\phi_{a\sigma}$ is a Kohn-Sham orbital, $a\sigma$ a multi-index labeling the orbitals and spin ($\uparrow$ or $\downarrow$), respectively, $n_{a\sigma}=|\phi_{a\sigma}|^{2}$, $n_{\sigma}=\sum a\sigma n_{a\sigma}$, and $n=n_{\uparrow}+n_{\downarrow}$. $\tilde{E}_{\text{LSDA}}$ differs from $E_{\text{LSDA}}$ since the kinetic energy is evaluated with respect to the orbitals minimizing the SIC-functional. The summations run over all the occupied orbitals, $E_{\text{H}}$ denotes the external energy functional due to ion cores, $E_{\text{H}}$ is the Hartree energy and $E_{\text{xc}}^{\text{LSDA}}$ is the LSDA exchange-correlation energy functional. The second term in Eq. (1) is the so-called self-interaction correction for all the occupied orbitals $a$. It restores the property

$$
E_{\text{H}}[n_{\sigma}] + E_{\text{xc}}^{\text{exact}}[n_{a,\sigma},0] = 0,
$$

that the exact DFT exchange-correlation functional has, namely, that for any single orbital density the Hartree term should be cancelled by the corresponding exchange-correlation term. The cost paid for restoring the above property is the orbital dependence of the SIC-LSDA energy functional [Eq. (1)]. The correction is only substantial for localized-orbital states, but vanishes for itinerant states. In the limit of all itinerant states the SIC-LSDA total-energy functional is identically equal to the LSDA functional.

The main idea behind the “local” implementation of the SIC-LSDA formalism (LSIC) is that within multiple scattering theory, in the framework of the KKR method, one works with the scattering phase shifts, describing scattering properties of single atoms in a solid. Among them only the resonant phase shifts are relevant, as they refer to localized states. Thus the self-interaction correction is associated with the on-site scattering potentials and leads to modified resonant scattering phase shifts. In particular, they become stronger localized. Details of the LSIC implementation are discussed in Ref. 27.

B. Magnetic interactions

The Heisenberg theory of magnetism assumes that it is possible to map magnetic interactions in a material onto localized spin moments, which in a classical picture can be represented by a vector. The resulting classical Hamiltonian,

$$
H = - \sum_{ij} J_{ij} \mathbf{e}_{i} \cdot \mathbf{e}_{j},
$$

contains only the unit vectors $\mathbf{e}_{i(j)}$ of the spin moments and the exchange parameters $J_{ij}$ describing the interactions between them.30 Here $i$ and $j$ index the sites.

It should be mentioned here that the Hamiltonian (4) can be extended to include additional effects such as magneto-crystalline anisotropy or tetragonal or rhombohedral distortions of the lattice. The latter reflect magnetoelastic effects which result in two different values for the nearest-neighbor (NN) exchange parameters, depending on a parallel or anti-parallel alignment of the moments. Such effects are usually present in experiments. Thus, special care is required when comparing theoretical and experimental results.
Our method of choice for the calculation of the exchange parameters \(J_{ij}\) makes use of the magnetic force theorem, but invokes also mapping onto a Heisenberg Hamiltonian. For comparison, we also apply the most commonly used approach which relies on the calculation of total energy differences between different magnetic configurations and mapping them onto a classical Heisenberg Hamiltonian.

1. Magnetic force theorem approach

The idea behind the magnetic force theorem\(^{26}\) is to consider infinitesimally small rotations of classical spins at two different lattice sites. These give rise to energy changes that are mapped onto the classical Heisenberg Hamiltonian via multiple scattering theory. This approach is based on the assumption that the potentials are unchanged by the rotations. The advantage of the MFT method is that the exchange integrals can be calculated directly in the relevant magnetic structure. The result for the exchange parameter \(J_{ij}\) of the two magnetic moments at sites \(i\) and \(j\) can be written as

\[
J_{ij} = \frac{1}{8\pi} \int dq \text{ Im} \text{Tr}_{\mathcal{L}}(\Delta_{ij} \hat{\sigma}^{\uparrow}_j \hat{\sigma}^{\uparrow}_i + \Delta_{ij} \hat{\sigma}^{\downarrow}_j \hat{\sigma}^{\downarrow}_i),
\]

where \(\hat{\sigma}^{\uparrow}_j\) is the scattering path operator between sites \(i\) and \(j\) and \(\Delta_{ij} = \hat{r}_{ij} - \hat{r}_{ij}^{-1}\), with \(\hat{r}_j\) being a single scattering operator for the atom at site \(j\). If not stated otherwise, all the results discussed later would have been obtained with the exchange parameters calculated using Eq. (5).

2. Energy differences approach

In this approach the total energies of the TMOs in the ferromagnetic (FM) and antiferromagnetic (AFI) and AFII configurations are taken into account. The AFI structure is characterized by oppositely magnetized planes which are stacked in (100)-direction. Suppose that magnetic interactions operate only between TM atoms—an assumption which is to be discussed later—the mapping onto the Heisenberg Hamiltonian yields

\[
J_1 = \frac{1}{16} (E_{\text{AFI}} - E_{\text{FM}})
\]

and

\[
J_2 = \frac{1}{48} (4E_{\text{AFI}} - 3E_{\text{AFI}} - E_{\text{FM}}),
\]

where \(J_1\) describes the interaction between the NN and \(J_2\) that of the next nearest neighbors (see Fig. 1). This mapping also assumes that the interaction between NN is independent of the sublattice the TM atoms are located on. Of course the choice of the three above mentioned structures restricts one to the determination of \(J_1\) and \(J_2\) only. Using more magnetic structures and hence calculating more exchange parameters is in principle possible. However, due to the nature of the present exchange mechanism, the super exchange, this has usually not been done for the TMOs. Although with this method we also restrict exclusively to \(J_1\) and \(J_2\), it is hoped that the comparison with the MFT method will shed some light on the validity of the underlying assumptions for the TMOs.

C. Néel temperatures

Having calculated the exchange parameters for an antiferromagnet one is able to calculate the Néel temperatures, \(T_N\). Several different approximations can be used, in particular the mean field approximation, the random phase approximation and classical Monte Carlo simulations.

1. Random-phase approximation

In the RPA one solves the equation of motion for the Green function of the spin operators. Following the approach of Rusz et al.,\(^{31}\) one ends up with a semi-classical formula for the average spin polarization \(\langle \hat{S}_\alpha^z \rangle(T)\) of a sublattice (e.g., A), as a function of temperature \(T\),

\[
\langle \hat{S}_\alpha^z \rangle(T) = \mathcal{L}(x) = \frac{1}{k_B T} \int dq \left( N^{-1}(q) \right)_{AA}^{-1},
\]

Here \(\mathcal{L}(x)\) is the Langevin function, \(\mathcal{L}(x) = \coth(x) - 1/x\), \(x\) is the volume of the first Brillouin zone and \(k_B\) denotes the Boltzmann constant. The matrix elements of \(N(q)\) are defined as

\[
N_{AB}(q) = \sum_C J_{AC}(0) \langle \hat{S}_C^z \rangle - \langle \hat{S}_A^z \rangle J_{AB}(q),
\]

with the Fourier transforms of the exchange parameters given by

\[
J_{AB}(q) = \frac{1}{\nu \sum_{j,k} J_{jk} e^{i \mathbf{q} \cdot (\mathbf{R}_j - \mathbf{R}_k)} x_{ijk}^A y_{jk}^B},
\]

where \(\nu\) denotes the number of interacting magnetic sites and \(x_{ijk}^A\) equals one if site \(j\) is on the magnetic sublattice \(A\) and zero otherwise. Equation (8) has to be solved self-consistently since the unknown quantity appears on its left and implicitly also on the right, via \(N(q)\). The Néel temperature is equal to the highest value of \(T\) at which \(\langle \hat{S}_\alpha^z \rangle(T)\) becomes different from zero.
2. Mean-field approximation

To obtain the MFA estimate of the Néel temperature, a matrix $\Theta$ with elements

$$\Theta_{AB} = \frac{2}{3k_B}J_{AB}(0)$$

is constructed, where $J_{AB}(0)$ stands for the Fourier transform of the exchange parameters, defined via Eq. (10), at $q = 0$. The largest eigenvalue of $\Theta$ yields the Néel temperature. If for any TMO in the AFII structure only the nearest and next-nearest-neighbor interactions are considered ($J_1$ and $J_2$, respectively), then the largest eigenvalue yields the well-known relation, $k_BT_N=4J_2$, indicating that the nearest-neighbor interaction $J_1$ does not have any influence on $T_N$. Since fluctuations are completely neglected in MFA the resulting Néel temperatures are commonly overestimated.

3. Monte Carlo simulations

We give a rather brief summary of the method of MC simulations as they are performed in this paper. For a deeper and complete understanding we refer the reader to the book by Landau and Binder.

To estimate $T_N$ via MC simulations a lattice representing the structure of the investigated system is constructed. The magnetic moment at lattice site $j$ interacts with its neighbors $j'$ via the $J_{ij'}$. During a MC run one picks a lattice site $j$ with the magnetic-moment vector $\mathbf{e}_j$, creates a new random direction $\mathbf{e}_{j'}$ and decides by looking at the energy of the system whether $\mathbf{e}_{j'}$ is accepted or $\mathbf{e}_j$ is kept. Performing this procedure $N$ times on a lattice of $N$ sites is defined as one MC step.

Starting from a certain initial configuration the system is brought into thermal equilibrium for a fixed temperature. After this, “measurements” and thermodynamical averaging of the observables of interest are performed. Since it is impossible during a simulation run to go through all possible configurations of the system, which would be formally necessary for averaging, one must ensure that the configurational subspace that one is restricted to is of physical significance. This is done by performing the so-called importance sampling. It is applied when one has to decide between the old and new magnetic moment vectors $\mathbf{e}_{j'}$ and $\mathbf{e}_j$ described above. There exist several methods to do this, in the present paper the Metropolis algorithm is used.

One must be aware of the fact that the finite size of the lattice, despite being periodic in all 3 dimensions, leads to a systematic error in the determination of the critical temperature. This so-called finite-size effect, however, becomes smaller with increasing lattice size. It can therefore be eliminated by extrapolating the critical temperatures for different lattice sizes.

For a magnetic system as in the present case it is straightforward to measure two quantities. One is the staggered magnetization $\mathbf{m}_s$ being some sort of an average of the absolute values of magnetization of the sublattices,

$$\mathbf{m}_s = \frac{1}{N} \sum_{j=1}^{N} \mathbf{e}_j e^{iQ\cdot\mathbf{r}_j}. \tag{12}$$

Here, $j$ labels the lattice sites being $N$ in total, $\mathbf{e}_j$ is the unit vector of the magnetic moment at lattice site $j$ and $Q$ is the normal vector of the planes of equal magnetization, in the AFII structure being $(1, 1, 1)$ for example. As a note, $\mathbf{m}_s$ is used instead of the total magnetization since the latter is equal to zero in antiferromagnets. The other quantity measured is the inner, i.e., magnetic, energy of the system $E$, which is given by Eq. (4). This whole procedure of relaxing into thermal equilibrium and thermodynamical averaging is repeated for different temperatures. In principle one can determine $T_N$ from the slope of the temperature dependence of $\mathbf{m}_s(T)$ and $E(T)$. However, there are quantities that show the critical temperatures more clearly. These are the magnetic susceptibility derived from $\mathbf{m}_s$, the specific heat derived from $E$, and the 4th-order cumulant $U_4$. The first two have a singularity at $T=T_N$, the last one has the property that the curves $U_4(T)$ calculated for different lattice sizes have a crossing point at $T=T_N$.

4. Quantum effects

As already mentioned, the three approaches described above are based on mapping of the single magnetic-moment interactions onto a classical Heisenberg Hamiltonian. These moments, however, are quantum objects and this should in some way be accounted for in the calculations. Wan, Yin, and Savrasov did this by replacing the classical $S^2$ in the Heisenberg Hamiltonian with the quantum mechanical expectation value $S(S+1)$, when calculating magnetic properties. Since in Eq. (4) $S^2$ is included in the $J_{ij}$, then to be consistent, one has to divide again by $S^2$. This gives rise to a factor $(S+1)/S$ for the energy and, eventually, also for the Néel temperature. This factor is, however, not needed when the Néel temperature is obtained based on the DLM method since it does not explicitly use the $J_{ij}$.

D. Magnon spectra

With the exchange interactions determined one can also calculate the magnon spectra $E(q)$. They are of special interest since they provide the standard method for determining the exchange parameters experimentally. The latter would be done by fitting a Hamiltonian containing the $J_{ij}$ as fitting parameters to a measured spin wave dispersion. As already mentioned in Sec. II, such Hamiltonians usually contain more terms than the one given in Eq. (4), which is why comparisons between different $J_{ij}$ results must be done carefully.

Considering multiple sublattices one can define magnon spectra as the eigenvalues of the matrix $N(q)$ given by Eq. (9). Assuming two magnetic sublattices, with the same absolute magnetization, and considering only the nearest and next-nearest-neighbor interactions, the spectra are given by...
EXCHANGE COUPLING IN TRANSITION METAL...

\[
E(q) = \frac{1}{\mu} \Sigma(J_{+\alpha}(q) - H_0)^2 - J_{-\alpha}^2(q).
\]  

(13)

Here \( \mu \) is the magnetic moment of the two sublattices in units of \( \mu_B \). \( J_{+\alpha}(q) \) [\( J_{-\alpha}(q) \)] are the Fourier transforms of the intra-(inter-)sublattice exchange parameters expressed, respectively, as

\[
J_{+\alpha}(q) = 2J_1 \times \left[ \cos \alpha(q_y + q_z) + \cos \alpha(q_z + q_y) \right]
\]
\[
+ \cos \alpha(q_x + q_y) \tag{14}
\]

and

\[
J_{-\alpha}(q) = 2J_1 \times \left[ \cos \alpha(q_y - q_z) + \cos \alpha(q_z - q_y) \right]
\]
\[
+ \cos \alpha(q_x - q_y) + 2J_2 (\cos 2\alpha q_x + \cos 2\alpha q_y + \cos 2\alpha q_z), \tag{15}
\]

and \( H_0 = J_{+\alpha}(0) - J_{-\alpha}(0) = -6J_2 \). In Eqs. (14) and (15) the vector \( q \) and accordingly its components are needed in units of \( 2\pi/a \) with \( a \) being the lattice constant of the TMO considered.

III. COMPUTATIONAL DETAILS

The transition metal monoxides crystallize in the rocksalt structure (B1, Fm\( \bar{3} \)m, space group 225). At low temperatures they show small lattice distortions (<2%). However, these distortions are not considered in the present calculations. The crystal potentials for the ground state calculations are constructed in the atomic sphere approximation (ASA). The ASA radii for the TM and oxygen atoms are chosen as 0.2895 \( \text{Å} \), with \( a \) being the lattice constant of a given TMO. To reduce the ASA overlap while keeping a good space filling, empty spheres are used with the ASA radii equal to 0.1774 \( \text{Å} \). The ratios of the respective ASA radii are kept constant across the TMO series.

For the electronic structure calculations the complex energy contour has 24 Gaussian quadrature points, and for the Brillouin zone (BZ) integrations a \( 14 \times 14 \times 14 \) \( k \)-points mesh is constructed. For the calculation of the magnetic interactions, using the MFT, 60 energy points on a Gaussian mesh in the complex plane are chosen. Convergence of the \( J_1 \) with respect to the number of \( k \) points is achieved with a \( 20 \times 20 \times 20 \) \( k \)-points mesh per energy point for the first 50 of them, and a \( 60 \times 60 \times 60 \) \( k \) mesh for the last 10 energy points, lying close to the Fermi energy.

For the MC simulations an fcc lattice representing the transition metal atoms in the TMO crystal is constructed. To avoid finite-size effects, the size of the lattice is varied from \( 40 \times 40 \times 40 \) to \( 60 \times 60 \times 60 \) elementary fcc cells. To use all observables described in the MC part of Sec. II one has to restrict the simulations to a relatively small number of MC steps. This is necessary in order to prevent the system from changing the orientation of the ferromagnetic sublattices, for example from (111) to (111), which are degenerate in energy. Thus, starting from the AFII state, the system is assumed to have reached thermal equilibrium after 5,000 MC steps, and for averaging 10,000 MC steps are performed. If one does so all observables yield the same result for the Néel temperatures for each TMO, respectively. To ensure a thorough exploration of phase space, simulations with up to 100,000 MC steps for averaging have also been performed. In this case the specific heat, not affected by reorientations of the magnetic sublattices, has reassuringly indicated the magnetic phase transitions to occur at the same temperatures as in the short simulations.

IV. RESULTS AND DISCUSSION

A. Exchange parameters

The present calculations of the exchange parameters of TMOs use the ground-state electronic structure properties of these materials as input. The latter are obtained self-consistently with the LSIC method, explained in detail in Ref. 21. In particular, as seen in Eq. (5), for the MFT approach the relevant quantities are the scattering properties evaluated at the equilibrium lattice constant of the ground state, AFII, magnetic structure. For the approach based on the energy differences only the total energies of the FM, AFI, and AFII structures, evaluated at the theoretical equilibrium lattice constants of the AFII configuration, are of relevance.

From Table I we can see that the LSIC method, treating localized, and itinerant electrons on equal footing, reproduces well the equilibrium lattice constants and also the corresponding spin magnetic moments in the AFII structure. The overall agreement with the experimental values is reasonable for both quantities. Note, however, that the experimental magnetic moments listed in the table include both the spin and orbital contributions, which are substantial for FeO and CoO, and non-negligible even for NiO. Regarding the calculated spin magnetic moments, they are effectively equal to the spin moments of the TM atoms, as the oxygen atoms are not polarized in the AFII environment, and the induced spin moments on the empty spheres are very small. In addi-

| TMO | \( a_0 \, [\text{Å}] \) | \( \mu \, [\mu_B] \) |
|-----|----------------|----------------|
| \( \text{MnO} \) | 4.49 | 4.63 |
| \( \text{FeO} \) | 4.39 | 3.68 |
| \( \text{CoO} \) | 4.31 | 2.69 |
| \( \text{NiO} \) | 4.24 | 1.68 |

\( ^a \text{Reference 42.} \)
\( ^b \text{Reference 43.} \)
\( ^c \text{Reference 44.} \)
\( ^d \text{Reference 45.} \)
\( ^e \text{Reference 46.} \)
\( ^f \text{Reference 47.} \)
agrees well with the idea of super-exchange\cite{33} and can easily
the higher shells are less than 0.1 meV in magnitude. This

Using the above ground state properties in the MFT ap-
proach, we have calculated the $J_{ij}$ exchange constants. This dependence of the calculated 

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The situation is very different for the $J_1$ exchange para-
ters. From Table II we see that, with the exception of NiO, 
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between the two theoretical approaches, but the signs are 
opposite. Looking more closely at the parameters calculated 
with the MFT we see that the results show the opposite trend 
to that found for $J_2$. Namely, the antiferromagnetic coupling 
is getting weaker as one moves from MnO to CoO, and in 
be explained with it. Consequently, and for the purpose of comparison with the results of the energy difference approach, in Table II we display only the $J_1$ and $J_2$ quantities, as well as the experimental results. Our results also provide justification for one of the assumptions underlying Eqs. (6) and (7), that the interactions between NN atoms are not dependent on the sublattices of the atoms.

From our results in Table II one finds that the $J_2$ parameters constitute the major part of magnetic exchange in TMOs and that in magnitude they agree reasonably well between the two theoretical approaches, MFT and $\Delta E$. The results are about 70–80% of the experimental values, except for FeO, where the agreement for the MFT $J_2$ is almost perfect and the $\Delta E$ value is larger than the experimental one. This rather accidental agreement for FeO can most likely be attributed to the fact that the experimental values are measured for Wüstite samples Fe$_{1-x}$O with $x \neq 0$\cite{50}. Also, the experimentally observed trend of the increasing absolute value of $J_2$ across the series is present in both approaches and is most likely associated with the increasing number of the TM 3$d$ electrons, responsible for the magnetic superexchange. Regarding the quantitative agreement with experiment, the MFT results agree better on average. One could envisage that this agreement could be further improved, if the MFT approach was applied in the DLM state\cite{29,56}. Nevertheless, compared to the other calculations displayed in Tables II and III, our present results may already be considered as being at least as good as those.

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\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|}
\hline
TMO & Exp. & MFT & $\Delta E$ & TB$^a$ & Exp. & MFT & $\Delta E$ & TB$^a$ \\
\hline
MnO & $-2.06, -2.64$ & $-0.91$ & 0.68 & $-4.41$ & $-2.79$ & $-1.99$ & $-1.65$ & $-1.09$ \\
FeO & 1.04, 1.84 & $-0.65$ & 0.48 & $-2.99$ & $-3.24$ & $-3.17$ & $-3.50$ & $-1.56$ \\
CoO & 0.70$^d$, 1.07 & $-0.32$ & 0.53 & $-1.83$ & $-6.30, d$ & $-5.31$ & $-4.84$ & $-4.40$ \\
NiO & $-0.69, 0.69$ & 0.15 & 1.42 & $-1.44$ & $-8.66, 5$ & $-9.51$ & $-6.92$ & $-6.95$ \\
\hline
\end{tabular}
\caption{The exchange parameters, $J_i$, in meV, with $i$ being the shell index, for the first two shells of all the studied TMOs and both the MFT and $\Delta E$, energy difference, methods. The experimental (exp.) values are given in the leftmost column, respectively, according to the Hamiltonian in Eq. (4). For $i > 2$, the absolute magnitudes of the $J_i$’s for all the TMOs have been less than 0.1 meV. For both the MFT and $\Delta E$ approaches the calculated equilibrium lattice constants from Table I have been used. The results of Harrison\cite{39} obtained using a tight-binding (TB) formalism for the complete series of TMOs are listed here for a direct comparison. The other previous results, obtained for selected monoxides, are listed in Table III.}
\end{table}

\begin{footnotesize}
$^a$Reference 39.
$^b$Reference 49.
$^c$Reference 50.
$^d$Reference 51.
$^e$Reference 52.
$^f$Reference 53.
$^g$Reference 54.
\end{footnotesize}
TABLE III. Summary of the first principles results for $J_1$ and $J_2$ in MnO and NiO, based on the Hamiltonian in Eq. (4), from the present and previous theoretical works for comparison. Only those close to experimental values are listed. For details see the corresponding references. We found one result by Feng for CoO, obtained by using the B3LYP hybrid functionals method, $J_1 = -47.12$ meV and $J_2 = -42.56$ meV. To our knowledge no further theoretical papers giving numerical values for the exchange parameters of FeO exist.

| Method  | $J_1$ [meV] | $J_2$ [meV] | Method  | $J_1$ [meV] | $J_2$ [meV] |
|---------|-------------|-------------|---------|-------------|-------------|
| Exp.    | $-2.06$, $-2.64$ | $-2.79$ | Exp.    | $-0.69$, $0.69$ | $-8.66$, $-9.51$ |
| This Work | $-0.91$ | $-1.99$ | This Work | $0.15$ | $-6.92$ |
| LDA+U a | $-2.50$ | $-6.60$ | GGA+U b | $0.87$ | $-9.54$ |
| OEP a | $-2.85$ | $-5.50$ | SIC-LMTO c | $0.90$ | $-5.50$ |
| PBE+U d | $-2.21$ | $-1.16$ | Fock35 e | $0.95$ | $-9.35$ |
| PBE0 d | $-3.10$ | $-3.69$ | B3LYP e | $1.20$ | $-13.35$ |
| HF d | $-0.73$ | $-1.16$ | UHF e | $0.40$ | $-2.30$ |
| B3LYP f | $-2.64$ | $-5.52$ | | | |

aReference 41.
bReference 15.
cReference 20.
dReference 22.
eReference 23.
fReference 55. NiO it becomes ferromagnetic. This can be explained by assuming both kinds of interaction to be present and to be competing, in the direct and indirect exchange between NN TM atoms. The picture is relatively intuitive for the direct case. For Mn, which has half filled d shells, one expects antiferromagnetic coupling since an electron hopping from one Mn atom to the other one keeps its spin. Thus, this transfer clearly prefers antiferromagnetic alignment of the Mn atoms.57 Moving across the TMO series the occupation of the minority spin channels is growing. This increases the probability of an electron hopping, if the TM atoms are ferromagnetically aligned. Thus, the character of the exchange should go toward ferromagnetic, which is what we find for the $J_1$ calculated via the MFT. Regarding the indirect exchange, we can also follow Goodenough.57 Nearest TM neighbors interact antiferromagnetically when two electrons in the same oxygen p or s orbitals are excited to the empty TM $e_g$ orbitals. The strength of this kind of interaction can be assumed not to change a lot along the TMO series, since the occupation of both the oxygen p or s and the TM $e_g$ orbitals does not change either. Ferromagnetic coupling on the other hand is provided by electrons of alike spin that are in different orbitals of the O atom. It is strengthened by a growing occupation of $t_{2g}$ orbitals because this increases intra-atomic exchange. Since the $t_{2g}$ occupancy is rising when moving across the TMO series from MnO to NiO21 one would expect that the magnitude of the ferromagnetic interaction increases while the antiferromagnetic does not. This tendency is clearly present in the MFT values for $J_1$ in Table II.

Looking at the agreement with experiment not much overlap can be spotted. For the MnO the agreement is satisfactory considering the simplicity of our Hamiltonian. For FeO the sign is opposite. This could be caused by the above mentioned fact that in experiment Wüstite samples of the kind Fe$_{1-x}$O are investigated while our calculations are performed for the ideal FeO system. For CoO and NiO comparison is difficult since experimental values of opposite signs, but similar absolute magnitude have been measured in different experiments. To conclude the comparison of MFT $J_1$ and experimental $J_1$ one can say that the agreement is not as good as for the $J_2$. Possible reasons have just been given, but it also seems that the experimental determination of the $J_1$ is not as accurate as for the $J_2$, as can be seen from the variety of numbers obtained for the same compounds. The lack of agreement between experimental and MFT values for the $J_1$ may influence the calculations based on those. For the magnon spectra it could be expected that, besides quantitative differences, due to the different signs even qualitative changes of the curves might occur. We shall see later, however, that the latter is not the case. The effect on the calculated Néel temperatures should be small in any case since the energy contributions of the NN in the AFII structure are canceled out.

For the $J_1$ calculated with the energy difference approach no obvious trends are seen in Table II, and in addition they are positive for all TMOs. For NiO the latter agrees qualitatively with the MFT-J1 and also with previous theoretical results, and the agreement with those by Ködderitzsch et al.,20 $J_1 = 0.9$ meV and $J_2 = -5.5$ meV, is also quantitatively rather good. For the other TMOs the sign of $J_1$ is opposite to the ones calculated with the MFT, and for MnO and CoO they also do not agree with previous theoretical investigations. The totally different behavior—compared to the MFT values—can be explained by looking at the electronic ground states of the calculated AFI and FM structures. In both of them the oxygen atoms carry a magnetic moment, which
they do not in the AFI structure. This magnetic moment can be assumed to give rise to magnetic interaction with the neighboring TM atoms (as a matter of fact, applying the MFT to AFI or FM structures yields NN exchange parameters of several meV in magnitude between TM and oxygen atoms). This, however, is in strong contradiction to the assumptions underlying Eqs. (6) and (7), stating that magnetic interaction only occurs between TM atoms. Thus, when using these equations anyway, this “artificially” created magnetic exchange is projected onto the $J_1$ and $J_2$. The reason for the latter quantity being relatively close to its counterpart calculated with the MFT is probably due to the large energy differences between the AFI configuration and the AFI as well as the FM configuration for each of the TMOs. This obviously reduces the error made in Eq. (7).

It should be mentioned that for all calculated pairs of $J_1$ and $J_2$, using SIC-LSDA, the resulting ground state magnetic structure is that of AFI,\(^5\)\(^8\) despite the relatively large spread of the $J_1$ parameters. Note that the exchange parameters $J_1$ and $J_2$, obtained for the TMO series by applying the MFT approach to the LSDA ground state electronic structure, show no agreement with experiment, except for NiO, which can perhaps be considered as a lucky coincidence. Furthermore, the $J$'s are longer-ranged, i.e., their character is more metallic. This agrees with the fact that their uncorrected (not SIC) ground states show only very small or no band gaps at all.\(^2\)

Finally, we would like to comment on the variation in the exchange parameters as a function of lattice constants shown in Fig. 3 for all the TMOs. As one can see, the absolute value of $J_2$ increases with decreasing $a$. This is in good agreement with the interpretation of the exchange parameters in terms of overlap integrals. The closer the atoms are, the larger the overlap is between the TM $d$ orbitals and the oxygen $p$ orbitals. A similar behavior is found for the $J_1$, which can be understood in the same way as for $J_2$. However, going through the TMO series and starting with MnO, the change of the $J_1$ gets smaller as the antiferromagnetic character becomes less pronounced. According to Goodenough's arguments,\(^5\)\(^7\) this suggests that the ferromagnetic coupling becomes more prominent than the antiferromagnetic one.

**B. Néel temperatures**

The calculated transition temperatures are summarized in Table IV. One finds that MFA overestimates the experimental Néel temperatures, whereas RPA underestimates them. This is what can be expected from general considerations.\(^5\)\(^5\) One can also see that the Néel temperatures calculated in the RPA approach based only on $J_1$ and $J_2$ do not differ significantly from those calculated using the 11 neighbor shells. This again agrees with the idea of superexchange. What is not expected is that, for MnO and FeO the RPA and MC results are relatively small compared to experiment. In fact, for these TMOs the MFA gives a better estimate. The probable reason for that is the general relative underestimate for the $J_2$. The latter being the main contribution of magnetic exchange, their underestimate is largest for MnO and decreases toward NiO. An exception is FeO. The agreement for $J_2$ is almost perfect, yet the RPA and MC estimates are roughly of the same quality as those for the other TMOs. However, it can again (see discussion of $J_1$ and $J_2$) be argued that due to the experimental imperfect FeO lattice other effects not considered in our approach may play an important role for the formation of magnetic order. The DLM results of Hughes et al.\(^2\)\(^9\) are, with the exception of NiO, in good agreement with

![FIG. 3.](https://example.com/figure3.png) (Color online) (a) The $J_1$ and (b) $J_2$ dependency on the lattice constant $a$ for all TMOs, calculated using MFT. The vertical lines mark the calculated equilibrium lattice constants from Table I.

**TABLE IV.** Summary of the Néel temperatures calculated with the $J_{ij}$ from the MFT approach (see Table II). In the top two rows the experimental and the DLM values are listed, followed by the RPA values based on the interaction of the first 11 TM-TM shells and of only the nearest and next-nearest neighbors (i.e., only $J_1$ and $J_2$). In rows 5 and 6 the MFA results shown, again using 11 or 2 shells, respectively. In the last row the results of the Monte Carlo simulations are presented.

| T$_N$ [K] | MnO | FeO | CoO | NiO |
|-----------|-----|-----|-----|-----|
| Experiment | 118 | 192 | 289 | 523 |
| DLM\(^a\) | 126 | 172 | 242 | 336 |
| RPA with $J_{1-11}$ | 81 | 146 | 252 | 440 |
| RPA with $J_{1,2}$ | 87 | 155 | 260 | 448 |
| MFA with $J_{1-11}$ | 122 | 210 | 362 | 628 |
| MFA with $J_{1,2}$ | 129 | 221 | 373 | 644 |
| MC | 90 | 162 | 260 | 458 |

\(^a\)Reference 29.
the experimental values. Their trend, however, is opposite to ours, namely the ratio $T_{N,\text{DLM}}^{\text{obs}}/T_{N,\text{RPA}}^{\text{obs}}$ becomes smaller with increasing atomic number. This could be due to not taking into account the quantum character of the systems, which in the present paper is done via the factor $(S+1)/S$,

where $S$ is calculated according to Hund's rules. Another possible reason especially for the NiO result, as discussed in Ref. 29, might be related to a possible importance of the short range order correlations that a single-site approximation-like DLM would not do justice to. Concentrating on our RPA and MC results, we have to admit that better calculations for the individually selected TMO systems can be found in literature. Among them are the calculations by Zhang et al.,

for NiO (with a rather semiempirical approach) and Towler et al.

for MnO. However, when studying the whole TMO series with the same approach, such as the above DLM application, the work reported by Harrison or Wan, Yin, and Savrasov,

it is hard to find ab initio results, treating electron correlations at the same level of sophistication and predicting the Néel temperatures qualitatively and quantitatively as accurately as in the present paper throughout the whole TMO series.

To finish, we briefly discuss the Néel temperature dependence on pressure shown in Fig. 4 for all the studied TMOs. To calculate the pressure, $p$, the Murnaghan equation of state has been used. Based on the behavior of $J_2$ seen in Fig. 3, it is not surprising that for the whole TMO series the calculated Néel temperatures increase with pressure. Qualitatively, this agrees with previous experimental and theoretical results, indicating a stability of the antiferromagnetic structure up to high pressures (several tens of GPa, at least) before it collapses and a paramagnetic or low-spin configuration takes over. We can compare the pressure dependence of $T_N$ to experiment (for MnO—Ref. 4, FeO—Ref. 65, CoO—Ref. 62, NiO—Ref. 64) for $p > 0$ by assuming them to be linear. Taking the pressure derivative of the normalized Néel temperatures, $\delta (T_N/T_N(p=0)) / \delta p$, we find that our calculated values increase too slowly, roughly by a factor of 1/2.

![FIG. 4. (Color online) The normalized RPA-based Néel temperatures $T_{N,\text{RPA}}^{\text{obs}}/T_{N,\text{RPA}}^{\text{obs}}$ for all TMOs as a function of pressure $p$. $T_{N,\text{RPA}}^{\text{obs}}$ is taken from row 3 of Table IV.](image)

![FIG. 5. Shown are the calculated TMO spin wave dispersions together with experimental data points for MnO (black diamonds, open circles, FeO, CoO, and NiO, respectively. The coordinates are cartesian and in units of $2\pi/a$. The path chosen along several high-symmetry lines starts at $X=(0.25,0.25,0.25)$ and goes along $[qqq]$, $[qq0]$ to $\Gamma=(0,0,0)$, then along $[qq0]$ to $M=(-0.5,0.5,0.5)$, and further along $[00q]$ to $\Gamma$ of the neighboring AFII Brillouin zone, then continuing along $[qq0]$ to $M$. The inlays in the MnO panel show the different branches along the AFII Brillouin zone.)](image)
all the studied TMOs, in the AFII structure, are shown in Fig. 5 together with the experimental results. Generally, the agreement between the calculated dispersion curves and the experimental observations is rather good, considering the Heisenberg Hamiltonian used in this work—anisotropy and alignment energy terms are neglected. This is also the reason why the calculated curves fail to reproduce the nonzero energies at $M=\{-0.5,-0.5,0.5\}$. Besides that, minima, maxima, and curvature are well reproduced. Furthermore it can be seen that except for FeO the theoretical curves generally underestimate the experimental energies, which is due to the underestimate of the $J_2$ parameters. The relative magnitude of the peak along $[q\bar{q}q]$ varies strongly, as one goes through the TMO series. This effect can be ascribed to the changing ratio of $J_2/J_1$.

The qualitative agreement with previous theoretical works, e.g. such as that of Solovyev and Terakura\(^4\) is good, although not in the absolute numerical terms, arising from different values of the Heisenberg exchange parameters $J_{ij}$.

V. CONCLUSION

We have used the local self-interaction correction, implemented in the multiple scattering theory in the framework of KKR in combination with the magnetic force theorem to study magnetic interactions in transition metal oxides. Specifically, we have calculated the $J_1$ and $J_2$ exchange parameters, the corresponding Néel temperatures and the respective magnon spectra for the whole TMO series. The most important conclusion of this work is that the combined approach used here provides an adequate description of magnetic interactions for the series as a whole. Without considering correlation effects the theoretical results in general do not agree with experimental findings. Furthermore, we have shown that our ab initio approach yields upper (MFA) and lower limits (RPA, MC simulations) for the Néel temperatures for the whole TMO series, and the calculated magnon spectra are in good qualitative agreement with experiment and other theoretical calculations.

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