February 2008

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Band-structure and correlation effects in the Co(111) planes of CoO

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(Presented on 9 November 2007; received 11 September 2007; accepted 16 October 2007; published online 13 February 2008)

The origin of the striking magnetic anisotropy of cobaltous oxide (CoO) is investigated by model calculations. A key feature of the electronic structure of the antiferromagnetic compound is the accommodation of Co 3d electrons in the (111) Co planes, with parallel and antiparallel interplane spin orientations and reminiscent of Cu 3d electrons in the CuO 2 planes in high-temperature superconductors. The strong correlations of the Co 3d electrons lead to a picture intermediate between traditional crystal-field and itinerant-electron descriptions of the magnetic anisotropy. Using a simple configuration-interaction approach, we analyze the effect of 3d 7-3d 7 bonds and their interplay with 3d 7-3d 7 configurations. © 2008 American Institute of Physics.

[DOI: 10.1063/1.2830958]

Cobaltous oxide, CoO, is an interesting antiferromagnetic (AFM) material, long investigated but comparatively little understood. Its relatively large magnetocrystalline anisotropy (of the order of 1 MJ/m 3) stabilizes the exchange bias in Co/CoO two-phase magnets, and its transport behavior has attracted much attention due to the Mott localization of the Co 3d electrons. While not suitable for permanent magnets, the understanding of its anisotropy is important for other applications, such as magnetic recording and sensors.

The oxide crystallizes in the cubic rocksalt structure, and its AFM spin structure is realized by the (111) Co planes with ferromagnetic order in each plane but antiferromagnetic spin alignment between adjacent planes. Figure 1 shows the corresponding spin structure. Furthermore, there is a slight tetragonal distortion along the c axis, which is also the uniaxial anisotropy axis. The band structure of paramagnetic CoO is similar to that of fcc Co, except for some band narrowing due to the increased interatomic distance and some hybridization with oxygen states. However, when it became clear that the Co 3d electrons are localized, emphasis shifted toward localized models. This refers, in particular, to the magnetic anisotropy of CoO, which is usually considered as a crystal-field effect but has recently attracted renewed attention due to its nontrivial orbital-moment character. In this paper, we investigate how electron correlations affect the magnetic anisotropy, interpolating between the undercorrelated (band structure) and overcorrelated (localized) limit.

There are two extreme models of magnetic anisotropy. The point-charge model (Fig. 2) treats the system as an insulator and ascribes the anisotropy to crystal-field charges, whereas the itinerant limit involves delocalized orbitals formed from atomic orbitals. We are primarily interested in the involved orbitals, but it is worthwhile recalling that anisotropy also involves spin-orbit coupling. The calculations are usually performed using the Stevens coefficients and perturbation theory, but the orbitals in Fig. 2 are chosen to yield zero and maximum orbital moment and anisotropy along the bond axis. This is because the elongated $|z^2\rangle$ and the flat $|x^2-y^2\rangle \pm i|xy\rangle$ shown in the figures have orbital moments of $m=0$ and $m=2$, respectively.

In the CoO structure, each Co atom is coordinated by 6 O nearest neighbors and by 12 Co next-nearest neighbors. The hopping integrals between the Co 3d and O 2p orbitals are large, but the low energy of the oxygen orbitals means that there is little hybridization between Co and O orbitals. The resulting 3d bands are fcc-like, and the bandwidth is largely determined by the hopping between next-nearest Co neighbors, with an oxygen hybridization contribution of somewhat more than 30%. Another difference between fcc...
Co and CoO is that the cubic crystal field changes sign, because the O atoms give rise to octahedral nearest-neighbor coordination. In other words, in CoO the energy of the $t_{2g}$ states is lower than that of the $e_g$ states. This change is of secondary importance to the band structure but reverses the sign of the corresponding anisotropy contribution.

The minority electrons of antiferromagnetic CoO are largely confined to the Co (111) planes, in contrast to paramagnetic CoO. The situation is reminiscent of the behavior of Cu 3$d^9$ in the Cu–O planes of La$_2$CuO$_4$, but both the band structure and the effect of correlations are complicated by the presence of more than one electron or hole per site. Using a configuration-interaction (CI) approximation, we analyze the effect of 3$d^6$–3$d^8$ bonds and their interplay with 3$d^7$–3$d^9$ configurations. This mechanism yields a specific anisotropy contribution, different from but adding to the electrostatic or “point-charge” and LCAO-type ligand-field (or itinerant) anisotropy contributions. Figure 3 shows the corresponding tight-binding band structure of a Co (111), using the $dd\sigma$, $dd\pi$, and $dd\delta$ hopping integrals from Ref. 2. The small bandwidth, less than 2 eV, contains the reduced hybridization between 3$d$ orbitals located in adjacent (111) planes.

Correlation effects in solids are a considerable challenge, because they occur “on top” of the band structure. In a few cases, such as the CuO$_2$ planes of La$_2$CuO$_4$, it is possible to restrict the consideration to a single band with one electron or hole per atom, corresponding to fluctuations between Cu 3$d^8$ and 3$d^{10}$ configurations. However, CoO has two Co $↓$ electrons per atom in the minority band, corresponding to a 3$d^7$ configuration. This means that 40% of the bands shown in Fig. 2 are filled, and we cannot restrict the consideration to low-lying states such as those formed from $|xy\rangle$ orbitals at the $M$ point.

Here, we take a somewhat different approach and focus on pairs of neighboring Co atoms. The representation of eigenfunctions $|\phi\rangle$ by approximate eigenfunctions $|\tilde{\phi}\rangle$ can be considered as a case of downfolding. As emphasized by Löwdin, the Schrödinger equation $H|\psi\rangle=E|\psi\rangle$ can be written as

$$H_{\text{eff}}|\phi\rangle + V|\chi\rangle = E|\phi\rangle$$  \hspace{1cm} (1a)$$
and

$$V^*|\phi\rangle + H_{\text{eff}}|\chi\rangle = E|\chi\rangle.$$  \hspace{1cm} (1b)$$
Here, $|\chi\rangle$ belongs to $|\phi\rangle$ but not to $|\phi\rangle$, and $H_{\text{eff}}$ and $V$ are the respective diagonal and off-diagonal interactions involving $|\chi\rangle$. Substitution of $|\chi\rangle$ into Eq. (1a) and (1b) yields

$$H_{\text{eff}}|\phi\rangle + V \frac{1}{E - H_{\text{eff}}} V^*|\phi\rangle = E|\phi\rangle.$$  \hspace{1cm} (2)$$
This equation is the type $H_{\text{eff}}|\phi\rangle=E|\phi\rangle$, where $H_{\text{eff}}$ describes the interaction of the bare states $|\phi\rangle$ with the environment $|\chi\rangle$.

The downfolding procedure leading to Eq. (2) is exact but leads to a complicated expression for $H_{\text{eff}}$. A crude approach is to treat the matrix elements of $H_{\text{eff}}$ as energy-dependent parameters. To investigate the effect of correlations on the crystal field, we choose the hybridized wave functions shown in Figs. 4(a) and 4(b). In Fig. 4(a), the interatomic hybridization between $z^2$ orbitals ($m=0$) yields bonding and antibonding states $s$ and $s^*$, respectively, whereas in Fig. 4(b) the same occurs for $xy$ and $x^2-y^2$ orbitals ($m=2$). Such hybridization exist in the (111) planes if we use an appropriately rotated coordinate frame. Figure 3 captures essential features of the problem, although anisotropy is obtained by summation over all pairs of orbitals, and we do not claim that the specific orbitals shown in Fig. 3 yield a disproportionately large anisotropy contribution.

In Fig. 3, the $\sigma$ and $\delta$ bonds yield to zero and easy-axis anisotropy, respectively. Note that $m=0$ ( $\sigma$ bond) implies...
zero orbital moment, and therefore zero spin-orbit coupling and zero anisotropy. The Hund’s rule interactions with fully occupied majority \( \uparrow \) band ensure that all spin states are \( \downarrow \), but there are several possibilities of accommodating the spins. This is a common feature of correlations, and a conceptually simple approach is actually the consideration of CI and compared to Fig. 2, the case (a) has now the lower energy, because it amounts to bonding \( \sigma \) orbital.

![Figure 4](image_url)

FIG. 4. Interatomic hybridization and crystal field: (a) \( dd\sigma \) overlap, (b) \( dd\delta \) overlap, and (c) and (d) two spin configurations derived from (a) and (b). Note that \( dd\sigma \) hopping integrals are much larger than \( dd\delta \) hopping integrals, and compared to Fig. 2, the case (a) has now the lower energy, because it amounts to bonding \( \sigma \) orbital.

Why does the second electron occupy the antibonding \( \delta^* \) level rather than the bonding and low-lying \( \sigma \) level, as in Fig. 4(d)? The reason is that Fig. 4(d) implies pronounced charge fluctuations, amounting to the coexistence of \( 3d^7-3d^7 \) and \( 3d^6-3d^8 \) configurations. This can be shown by explicit consideration of the wave functions

\[
\begin{align*}
|\sigma\rangle &= |\sigma\rangle + |\sigma\rangle, \\
|\sigma^*\rangle &= |\sigma\rangle - |\sigma\rangle,
\end{align*}
\]

where the indices refer to the top and bottom orbitals shown in Fig. 4(a). The \( \delta-\delta^* \) configuration has the two-electron wave function

\[
|\delta\delta^*\rangle = |\delta\rangle|\delta^*\rangle - |\delta^*\rangle|\delta\rangle = |\delta\rangle|\delta\rangle - |\delta\rangle|\delta\rangle,
\]

that is, each of the two atomic \( \delta \) orbitals (top and bottom) is \( \delta^* \) occupied. The same calculation for

\[
|\sigma\rangle \delta \sim |\delta\rangle|\sigma\rangle - |\delta\rangle|\sigma\rangle,
\]

yields a 50% admixture of states such as \( |\sigma\rangle|\delta\rangle - |\delta\rangle|\sigma\rangle \), meaning that both electrons are accommodated in the top or bottom atom. This costs correlation energy \( (U) \), is energetically unfavorable, and causes the transition from Figs. 4(d) and 4(c).

From a many-electron point of view, each one-electron calculation corresponds to one Slater determinant where available low-lying one-electron states are filled like liquid which is poured into a jar, as in Fig. 4(d). A simple example is the tight-binding (or LCAO) approximation, where the one-electron states are linear combinations of atomic orbitals, but advanced local spin-density approximation (LSDA) electronic-structure calculations are based on the same principle. This includes LSDA+U calculations, which contain a self-energy correction \( (U) \) and yield improved predictions on a quantum-mechanical mean-field level. However, the treatment of specific correlation effects, such as spin-charge separation, requires the explicit or implicit consideration of several Slater determinants. In the present paper, this is done in an approximate way, by considering the two configurations Figs. 4(c) and 4(d), but a detailed analysis of configurations and their interactions in CoO remains a challenge to future research.

In summary, we have investigated how electron correlations affect the magnetocrystalline anisotropy of CoO. Correlations suppress the charge fluctuations \( (3d^7 \) and \( 3d^8 \) configurations) associated with the formation of bonding states and thereby change the character of the crystal-field interactions responsible for magnetic anisotropy. The resulting scenario is intermediate between the electrostatic or point-charge and itinerant (or LCAO-type ligand-field) mechanisms. In the highly simplified model of Fig. 3, this leads to a change from zero anisotropy to easy-axis anisotropy.

This research is supported by NSF MRSEC and NCMN.

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