Pressure-Driven Metal-Insulator Transition in Hematite from Dynamical Mean-Field Theory

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The local density approximation combined with dynamical mean-field theory is applied to study the paramagnetic and magnetically ordered phases of hematite Fe2O3 as a function of volume. As the volume is decreased, a simultaneous first-order insulator-metal and high-spin to low-spin transition occurs close to the experimental value of the critical volume. The high-spin insulating phase is destroyed by a progressive reduction of the spectral gap with increasing pressure, upon closing of which the high-spin phase becomes unstable. We conclude that the transition in Fe2O3 at ≈ 50 GPa can be described as an electronically driven volume collapse.

The metal-insulator transition (MIT) is one of the central topics in the physics of strongly correlated electron materials. Despite considerable progress in understanding of this phenomenon in the past decade, we are only starting to uncover the multitude of possible transition scenarios in real materials. An important instance of the MIT is the pressure-driven transition accompanied by a change of the local spin state [high spin (HS) to low spin (LS) transition] seen in MnO [1], BiFeO3 [2], or Fe2O3. Understanding of the pressure-driven HS-LS transition and its relationship to the MIT and structural or volume changes is relevant to a broader class of oxides [3], often with geophysical implications.

In this Letter we study the spin and metal-insulator transition in hematite (α-Fe2O3) under pressure using the local density approximation combined with dynamical mean-field theory (LDA + DMFT) approach [4]. We investigate the effect of temperature and magnetic long-range order (LRO) and suggest a new classification of the simultaneous spin and metal-insulator transitions in related materials. At ambient conditions, hematite is an antiferromagnetic (AFM) insulator (TN = 956 K) with the corundum structure [5]. The iron ions having a formal Fe3+ valence with five d electrons give rise to a local HS state. Photoemission spectroscopy (PES) [6,7] classified hematite as a charge-transfer insulator. A charge gap of 2.0–2.7 eV was inferred from the electrical conductivity data [8]. Under pressure, a first-order phase transition is observed at approximately 50 GPa (82% of the equilibrium volume) at which the specific volume decreases by almost 10% and the crystal symmetry is reduced (to the Rh2O3-II structure) [9–11]. The high-pressure phase is characterized by metallic conductivity and the absence of both magnetic LRO and the HS local moment [10]. Badro et al. showed that the structural transition actually precedes the electronic transition, which is, nevertheless, accompanied by a sizable reduction of the bond lengths [12].

Investigations of the electronic structure of hematite using various theoretical approaches lead to different (and frequently controversial) results. Similar to many transition metal oxides, the local spin density (LSDA) and generalized gradient approximations fail to reproduce the ground state under ambient conditions, severely underestimating the magnetic moment on iron and the spectral gap [13]. The static mean-field LSDA + U approach [14] improves the size of the magnetic moment and spectral gap in the HS phase, but fails to capture the metal-insulator transition; thus, LSDA appears more appropriate for the description of the LS high-pressure phase. The LDA + DMFT method, which can describe the evolution from strong to weak correlations, is well suited to study the transition regime [15]. The only DMFT study of Fe2O3 to date was limited to the t2g band manifold finding a low-spin metallic phase at high pressure [16].

Here we use a Hamiltonian including explicitly all O-p and Fe-d orbitals, which allows us to address the following questions. (i) Is there an electronic transition without a change of the crystal structure? (ii) Is there a simultaneous metal-insulator and local moment transition as found in the isoelectronic MnO [15]? (iii) What is the nature of the transition and how is it affected by temperature and magnetic LRO? We employ the implementation of LDA + DMFT, previously applied to NiO [17], to “solve” the first-principles multiband Hubbard Hamiltonian

\[ \hat{H} = \sum_k \hat{a}_k^\dagger \hat{H}_k \hat{a}_k - \varepsilon_{de} (n_d) \hat{N}_d + \sum_{\mathbf{r} \in \mathbf{Fe}} \hat{n}^d_{\mathbf{r}} \hat{U}^{dd}_{\mathbf{r}} \hat{n}^d_{\mathbf{r}} \]

dealing with the dynamical mean-field approximation. Here, \( \hat{H}_k \) is the LDA Hamiltonian, \( 38 \times 38 \) matrices (two Fe2O3 units) in the basis of Fe-d and O-p Wannier orbitals, on a uniform
Below this value only a LS solution characterized by a substantially reduced local moment exists. The difference between the screened and instantaneous values indicates that the LS moment cannot be viewed as a rigid object. The HS and LS solutions coexist in the 0.8–0.85V₀ range. The Fe-d orbital occupancies fit into the iconic picture of a fully spin-polarized d shell in the HS state, and an orbitally polarized d shell in the LS state. Nevertheless, the substantial e₉g population in the LS phase reflects a sizable covalent p–d bonding (hybridization between O-p and Fe-e₉g orbitals).

In Fig. 2 we show the corresponding single-particle spectra obtained with the maximum entropy method for analytic continuation [23]. In the LS phase the e₉g-derived bands are above the chemical potential, carrying some O-p admixture with them, which leads to charge transfer from the O-p orbitals and an increase of nd (see Fig. 1). Relaxation of the electronic states, which compensates this charge redistribution in real material, is mimicked by the self-consistency of the double-counting term in the present scheme. Comparison to PES and inverse PES (see left-hand inset of Fig. 2) reveals a fair agreement between the theory and experiment. The computed ambient pressure HS phase is found to be an insulator with a gap of 2.6 eV. Upon compression the gap gradually shrinks due to broadening of the bands and increase of the crystal-field splitting. At the high-pressure end of the HS stability region (0.8V₀) the gap is reduced to a fraction of an eV. The spectra in the LS phase closely resemble the non-

FIG. 1 (color online). Fe-d occupancies (left-hand panel) at various specific volumes in the paramagnetic (PM) state at T = 580 K. The thick line marks the average d occupancy per orbital. Corresponding values of the screened (diamonds) and instantaneous (circle) local moment are shown in the right-hand panel.
interacting one (see the right-hand inset of Fig. 2). Orbital polarization of the $t_{2g}$ band due to distortion of FeO$_6$ octahedra is seen in the LS spectra.

Unlike LDA or LDA + $U$, the LDA + DMFT allows opening of a gap and formation of local moments without magnetic LRO, as observed in many materials. Reducing the symmetry to allow the experimental AFM LRO we observe the following. At room temperature the HS solutions are magnetically ordered, while all LS solutions remain paramagnetic. The spin-polarized spectra (Fig. 3) show that the gap and the main spectral features are insensitive to the AFM LRO. The magnetization curves reflect increase of the exchange coupling due to shortened interatomic distances.

Our observations are summarized in the $T$-$V$ phase diagram in Fig. 4. We distinguish three phases: (i) paramagnetic LS, (ii) AFM HS, and (iii) paramagnetic HS, separated by the line of Néel temperature $T_N$ and the coexistence region of the HS and LS phases. The calculated $T_N$ is affected by two approximations, both favoring the AFM phase. First, the lack of nonlocal spin fluctuations in DMFT, and second, the approximate form of the local interaction, limited to Ising terms, rendering the HS local state twofold ($S_z = \pm 5/2$) instead of sixfold degenerate ($S = 5/2$). The latter is also reflected in the shape of the magnetization curves, which can be fitted with a mean-field theory based on the Brillouin function $B_{1/2}$ (see the inset of Fig. 3). Given these approximations, the calculated $T_N$ of 1600 K appears to be in reasonable agreement with the experimental value of 956 K. The effect of AFM LRO on the HS-LS transition is minor, consisting in a slight shift of the coexistence region in favor of the HS phase. Interestingly, in the vicinity of the HS-LS boundary we find that with increasing temperature the AFM LRO is not destroyed by magnetic fluctuations, but due to the collapse of the constituting moments.

Next, we discuss how a $d^5$ HS phase can be destabilized by increase of the crystal field (and the bandwidth) due to applied pressure. We simplify the local interaction to $\Sigma_{i,j,\sigma} U n_{i\sigma} n_{j,-\sigma} + (U - J) n_{i\sigma} n_{i,\sigma}$ neglecting the orbital dependence in order to keep the following expressions transparent. The HS gap can be estimated as

$$E_g \approx E(d^6) + E(d^4) - 2E(d^5) - W = U + 4J - \Delta - W,$$

where $\Delta$ is the $e_g - t_{2g}$ crystal-field splitting, $W$ is the average of the $e_g$ and $t_{2g}$ bandwidths, $E(d^6)$ is the ground state energy of an ion in the HS state, and $E(d^4)$, $E(d^5)$ are the energies of the corresponding electron addition and removal states, respectively. (In charge-transfer compounds such as hematite, $E_g$ may be substantially reduced by the presence of the ligand bands.) Similarly the energy difference between ions in LS and HS states can be estimated as

$$E(\text{LS}) - E(\text{HS}) = 6J - 2\Delta.$$

The HS state becomes unstable when either (1) or (2) becomes negative. The former is an instability with respect to transfer of an $e_g$ electron into a $t_{2g}$ orbital on the neighboring cation, the latter is an on-site instability with respect to redistribution of the $d$ electrons. We adopt the

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**FIG. 3 (color online).** Spin-polarized Fe-$d$ spectra at the ambient pressure for 580, 1160, and 1450 K (top to bottom). Left-hand inset: The same spectra averaged over spin. Right-hand inset: The staggered magnetization versus temperature curves for $\nu = 1.0$ (circles, black), $\nu = 0.9$ (squares, red), $\nu = 0.85$ (diamonds, green), and $\nu = 0.8$ (triangles, blue). At empty symbols only the LS solution exists. The lines represent mean-field fits.

**FIG. 4 (color online).** Calculated $V$-$T$ phase diagram of hematite. The symbols mark the actual computational results: HS state (circle), LS state (diamond). Empty diamonds mark LS solutions stable only if PM symmetry is constrained; dotted circles mark HS solutions stable only if AFM LRO is allowed. The black line shows the Néel temperature estimated from fits to $\nu = 1.0$, 0.9, and 0.85 data. The shaded area marks the estimated HS-LS coexistence region with PM constraint; the dashed lines mark the coexistence regime when AFM LRO is allowed.

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terms *gap closing* and *local state transition* for the two types of behavior. While the gap continuously disappears as the gap closing transition is approached, in the local state transition the gap remains finite. Whether the transition in this case is insulator to insulator or proceeds through the appearance of in-gap states depends in a first approximation on $U$ and $W$. We point out that analogous estimates for a two band system provide a surprisingly accurate description of the phase diagram in Fig. 2 of Ref. [24], identifying the Mott insulator-to-metal transition as gap closing and the Mott insulator-to-band insulator transition as local state transition.

Comparing the hematite spectra with the behavior of MnO [15] an obvious similarity is that the HS phase is an insulator while the LS phase is metallic. However, there are two important differences in how the transition proceeds. First, in MnO there is a continuous crossover via a sequence of intermediate states while in hematite we find a first-order transition with coexisting HS and LS solutions. Second, the HS gap in hematite shrinks practically to zero at the border line of the HS stability region while in MnO the gap does not close completely and the transition starts with the appearance of an in-gap spectral density. The latter observation leads us to the conclusion that the HS-LS–insulator-metal transition in hematite proceeds through the *gap closing* mechanism while in MnO a *local state transition* takes place as was argued in Ref. [15]. The gap closing scenario in hematite is also supported by the slope of the HS-LS phase boundary. The HS state can be destroyed by thermal fluctuations, contrary to the expectation for the local state transition where the HS state is favored at high temperature due to its higher entropy.

Finally, we briefly address the structural and volume changes [10–12]. The corundum and Rh$_2$O$_3$-II structures have similar local Fe environment, and transitions between the two were reported in other materials without significant impact on the electronic structure. Here we did not attempt to distinguish between the two structure types. Finding the electronic HS-LS–insulator-metal transition in the high-volume phase shows that the electronic transition is not conditioned by the structural transition or volume collapse. Conversely the electronic transition provides a natural explanation of the volume collapse as a consequence of emptying of the antibonding $e_g^*$ bands in the LS phase and corresponding strengthening of the Fe-O bonds.

In summary, we have observed a simultaneous HS-LS–insulator-metal transition in the corundum structure of hematite close to the experimental volume in the numerical results obtained by LDA + DMFT. The magnetic long-range order is found to play a minor role in the transition. We have discussed two possible scenarios, the *gap closing* and the *local state transition*, and concluded that the transition in hematite, unlike in MnO, proceeds through the gap closing mechanism. Our results are consistent with the picture of an electronically driven volume collapse.

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