Unconventional magnetism as a consequence of the charge disproportionation and the molecular orbital formation in Ba$_4$Ru$_3$O$_{10}$

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The magnetic and electronic properties of Ba$_4$Ru$_3$O$_{10}$ were investigated by the ab-initio and model calculations. It is shown that nonmagnetic ground state of the one-third Ru$^{4+}$ ions is not due to the correlation effects. It is rather caused by the charge disproportionation between crystallographically different Ru and the molecular orbital formation in the Ru’s trimer.

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I. INTRODUCTION

The complex Ru oxides are widely investigated in the last decades because of their unusual electronic and magnetic properties. The odd-parity pairing was found in the superconductor Sr$_2$RuO$_4$, while Ca$_2$RuO$_4$ became one of the model systems for study of the orbital-selective Mott transition. The integer spins on the Ru$^{4+}$ ions likely form the Haldane chains in Tl$_2$Ru$_2$O$_7$, which leads to a drastic decrease of the magnetic susceptibility at low temperatures. Similar behavior of susceptibility was recently observed in another Ru oxide - Ba$_4$Ru$_3$O$_{10}$. This drastic decrease was initially attributed to the formation of the unconventional antiferromagnetic (AFM) state, but the mechanism of its stabilization is still unknown.

Structurally Ba$_4$Ru$_3$O$_{10}$ is made of the Ru-trimers. These trimers are formed by three RuO$_6$ octahedra sharing their faces. Each trimer is connected with four neighboring trimers (see Fig. 1) via corners of the outer RuO$_6$ octahedra to build corrugated layers. We denote the middle Ru ion in the trimer Ru$_m$, while two outer Ru – Ru$_o$.

In Ba$_4$Ru$_3$O$_{10}$ the Ru ions must have 4+ oxidation state with 4 electrons in the 4$d$ shell, since Ba is 2+ and O is 2-. Due to a large $t_{2g} - e_g$ crystal-field splitting (see Sec. III for details) these four 4$d$–electrons should be stabilized in the $t_{2g}$ sub-shell resulting in the state with $S$=1. Indeed, at room temperature Ba$_4$Ru$_3$O$_{10}$ was found to be paramagnetic with the effective moment $\mu_{eff} = 2.83 \mu_B$ typical for the S=1 ions, while the decrease of the temperature below $T_N = 105$ K leads to the formation of the unconventional AFM state, where one-third of the Ru$^{4+}$ ions are not ordered (tentatively these are Ru$_m$).

Several models were proposed in Ref. to explain this physical phenomenon. One may think that (1) due to some reason a part of the Ru ions stays paramagnetic down to a very low temperature; that (2) they are randomly frozen; or that (3) because of a strong spin-orbit coupling a nonmagnetic state of the Ru ion with $J = 0$ is stabilized.

In order to check all these possibilities, the ab-initio band structure calculations were performed for this compound. We found that the true reason for the absence of the magnetic moments on one-third of the Ru ions is the charge disproportionation between two crystallographically nonequivalent Ru (Ru$_m$ and Ru$_o$) and formation of the molecular orbital in the trimers. In the band-structure picture it is manifested in the suppression of the nonmagnetic 4$d$ density of states (DOS) of the middle Ru$_m$ in the vicinity of the Fermi level.

FIG. 1: (color online). The crystal structure of Ba$_4$Ru$_3$O$_{10}$. Projection on the $bc$–plane. Oxygen ions are in blue, Ruthenium - in red, Ba - not shown for the simplicity. Ru$_m$ are in the middle of the trimers, Ru$_o$ - on the corners.

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II. CALCULATION DETAILS

For the band structure calculations we primary used the pseudo-potential PWscf code. We utilized generalized gradient approximation (GGA) with Perdew-Burke-Ernzerhof version of the exchange-correlation potential and ultrasoft scalar-relativistic pseudo-potentials with nonlinear core correction (for better description of the magnetic interactions). The charge density and kinetic energy cut-offs were taken 40 Ry and 200 Ry, respectively. 664 \( k \)-points in a full Brillouin zone were used in the calculation.

The generalized gradient approximation with the account of the on-site Coulomb repulsion (GGA+U method) was used to test an importance of the electronic correlations. The Hubbard’s on-site repulsion parameter \( U \) and Hund’s rule intra-atomic exchange \( J_H \) for Ru were calculated in Ref. 3 to be 3.0 eV and 0.7 eV.

The Wannier function projection was performed within the linearized muffin-tin orbitals (LMTO) method as it was described in Ref. 10.

All the calculations were performed for the crystal structure corresponding to \( T=10 \) K.

III. NONMAGNETIC GGA CALCULATIONS

We start with the analysis of the results obtained in the nonmagnetic generalized gradient approximation (GGA), presented in Fig. 2. One may see that the oxygen 2p states are placed primarily from -7 to approximately -1 eV. The top of the valence band is formed by Ru-4d \( (t_{2g}) \) states. The Ru-\( e_g \) orbitals are above 1.5 eV. The \( t_{2g} - e_g \) crystal field splitting estimated by the center of gravity calculation is 3.38 and 3.20 eV for Ru\(_m\) and Ru\(_o\) respectively. Thus one would indeed expect that such a strong crystal field splitting would prevent occupation of the \( e_g \) orbitals and \( d^4 \) configuration of Ru\(^{4+}\) would result in the S=1 ground state of each Ru ion.

However, there is another one quite important feature of the nonmagnetic DOS related to the splitting between the Ru\(_m\) and Ru\(_o\) 4d states, which is clearly seen from Fig. 2. The same center of gravity calculation shows that this splitting exceeds 0.13 eV. The Ru\(_m\)-4d states integrally turns out to be lower than Ru\(_o\)-4d due to a larger mean Ru-O distance. According to Ref. 4, the difference in the mean Ru-O distances for these two Ru exceeds 0.035 Å. This is not a small number, since the difference in ionic radii between Ru\(^{4+}\) and Ru\(^{3+}\) is 0.06 Å. As a result one could expect that there would be more d-electrons on the middle Ru, i.e. it would have on the average a smaller valence. And indeed, according to our nonmagnetic GGA calculations the difference in charge between Ru\(_m\) and Ru\(_o\) is 0.52 electron per atom, which is even larger than the charge disproportionation on the Fe sites in the charge ordered phase of Fe\(_3\)O\(_4\). Thus, Ru\(_m\)-\( t_{2g} \) partial DOS states turns out to be almost completely below the Fermi level, except a small shoulder and a narrow peak at \( \sim 0.9 \) eV. The detailed analysis of the band structure, presented in Fig. 3 confirms that the band corresponding to this peak in DOS at \( 0.9 \) eV is dispersionless. One may also see that each band in the ZT direction is two times degenerate, since there are two formula units (f.u.) in the unit cell. In effect each trimer provides one dispersionless band at \( \sim 0.9 \) eV.

In order to analyze the formation of this band we carried out the calculation within the local density approximation (LDA) and the LMTO method for which one may...
corresponds to the band, which has the energy ∼ \text{tibonding} clearly seen in the Fig. 4, where the highest in energy and allows a direct overlap between packing of the RuO \text{matrix elements}. This is mainly due to a face-sharing local surroundings (oxygens), but also by the inter-site determined not only by the crystal-field created by the in the ZT direction.

The diagonalization of the on-site Hamiltonian in a real space for each Ru leads to the order of the levels predicted by the local distortions. The tetragonal elongation of the RuO \text{octahedra} results in two low-lying almost degenerate t_{2g}-levels split from the higher t_{2g} orbital by ∼330 meV. However, a close inspection of the Hamiltonian shows that the hopping integrals between Ru\text{m} and Ru\text{o} exceed 360 meV, which is, in contrast to naive expectations, even larger than the off-diagonal on-site matrix elements for the t_{2g} orbitals. Thus, the orbitals on which the Ru−4d electrons tend to localize are determined not only by the crystal-field created by the local surroundings (oxygens), but also by the inter-site matrix elements. This is mainly due to a face-sharing packing of the RuO\text{o} octahedra in the trimers, which allows a direct overlap between d−orbitals on the neighboring Ru sites. Indeed, the nearest neighbors Ru-Ru distance across the common face 2.55 \text{Å} is even less than that in ruthenium metal, 2.65 \text{Å}. \text{[13]} It is also quite important that the relevant orbitals here are 4d−orbitals, which are more extended than 3d. \text{[14]}

In order to take into account the hoppings, we constructed the large 15×15 Ru\text{o}-Ru\text{m}-Ru\text{o} Hamiltonian for the trimer in a real space. Diagonalizing this Hamiltonian one obtains that the lowest and highest in energy are the molecular orbitals which have the a_{1g} symmetry with the largest contribution on the central Ru\text{m} ion. This is clearly seen in the Fig. 4 where the highest in energy antibonding a_{1g}^{\dagger} molecular orbital is plotted. This orbital corresponds to the band, which has the energy ∼ 0.9 eV in the ZT direction.

Inspecting Fig. 4 one may see that the contribution of the Ru\text{m}−4d states to the antibonding a_{1g}^{\dagger} orbital should be the largest (the same is actually correct also for the bonding a_{1g} wave function). This is easy to understand by solving the three-site (i = 1, 2, 3) problem with the single orbital per site in the tight-binding approximation. For simplicity the energies of each orbital are taken the same, \varepsilon_i = 0, and the hoppings, t > 0, are nonzero only between the nearest sites. For the linear cluster of three atoms the Hamiltonian is written as

\begin{equation}
H = \begin{pmatrix}
0 & t & 0 \\
t & 0 & t \\
0 & t & 0
\end{pmatrix}.
\end{equation}

The energy spectrum consists of the bonding (b), antibonding (ab) and nonbonding (nb) orbitals:

\begin{align}
E_{\text{ab}} &= t\sqrt{2}, \quad \psi_{\text{ab}} = \frac{1}{\sqrt{2}}(\psi_1 - \sqrt{2}\psi_2 + \psi_3), \\
E_{\text{nb}} &= 0, \quad \psi_{\text{nb}} = \frac{1}{\sqrt{2}}(\psi_1 - \psi_3), \\
E_b &= -t\sqrt{2}, \quad \psi_b = \frac{1}{2}(\psi_1 + \sqrt{2}\psi_2 + \psi_3).
\end{align}

As a result one may see that the weight of the states corresponding to the middle, i = 2, atom to the bonding and antibonding wave functions equals to 50%, while they do not contribute to the nonbonding molecular orbital at all. Thus, the Ru\text{m}−t_{2g} states of the a_{1g} symmetry are pushed away from the Fermi level due to the molecular orbital formation. Moreover, the shift of the rest of the Ru\text{m}−t_{2g} states downwards, with the cor-
TABLE I: Total energies, magnetic moments on the Ru₄ and Ru₄m ions and band gaps. Results of the GGA in the PWscf code. Total energies are per formula unit (f.u.). The notation \( \uparrow m \downarrow \) means that two Ru ions are AFM, while moment for the Ru₄m can be found in the 3rd column. Symbol \(-\) denotes the bond with adjacent trimers. The unit cell vectors are chosen in such a way, that the type of inter-trimer magnetic order (ferro- or antiferromagnetic) for presented configurations is the same for all 4 inter-trimer bonds.

| Total energy (meV) | Spin moments Ru₄/Ru₄m (\(\mu_B\)) | Band gap (eV) |
|--------------------|--------------------------------------|--------------|
| \( \uparrow m \downarrow - \uparrow m \downarrow \) | 0 | 0.95/0 | 0.12 |
| \( \uparrow m \uparrow - \uparrow m \uparrow \) | 22.7 | 0.98/0.33 | Metal |
| \( \uparrow m \downarrow - \downarrow m \uparrow \) | 20.1 | 0.94/0 | Metal |
| \( \uparrow m \uparrow - \downarrow m \downarrow \) | 61.0 | 0.69/0.15 | Metal |
| 0m0 - 0m0 | 137.6 | 0/0 | Metal |

responsible charge disproportionation, leads to the suppression of the Ru₄m 4d partial DOS in the vicinity of the Fermi level. The Ru₄ - 4d states in contrast form well defined peak exactly at the Fermi energy.

The Stoner criteria cannot be applied directly to the partial DOS, but it is clear that the gain in the magnetic energy due to the spin splitting on Ru₄m will be minimal (because of the pseudogap in its partial DOS), while the loss of the band energy will be substantial. Thus in the band picture the Ru₄m - 4d states are not expected to be polarized. In contrast, the Ru₄ - 4d states can be easily magnetized. This is exactly what we observe in the magnetic GGA calculations.

IV. MAGNETIC GGA CALCULATIONS

Several configurations were investigated in our magnetic GGA calculations (see Tab. I). The lowest in energy is the state when the spins on the Ru₄ both in the same and in the adjacent trimers are antiparallel (\( \uparrow m \downarrow - \uparrow m \downarrow \)). The partial DOS corresponding to this configuration are plotted in Fig. V. One may see that there is no splitting between majority and minority spins of the Ru₄m. Thus according to our calculation 1/3 of the Ru atoms indeed are nonmagnetic, exactly as it was observed in the experiment.

The magnetic moment on the Ru₄m appears only in the configurations with the FM order of the Ru₄ in the trimers. The ferromagnetically ordered spins of the Ru₄ create an exchange field on the Ru₄m, which slightly magnetize these ions. On the band-structure language the appearance of some magnetization on Ru₄m in a ferromagnetic state can be explained by the spin splitting of the Ru₄m - 4d states due to their hybridization with the Ru₄ - 4d states, which as we have shown above is large enough.

The magnetic moments on Ru₄ are smaller than 2\(\mu_B\) expected for \(S=1\). This is a rather typical situation for the transition metal oxides (especially for those based on the 4d and 5d elements) and can be attributed to a strong hybridization with the oxygens. Because of the hybridization we also observe non zero magnetic moments on some of the oxygens, which vary from 0.04 to 0.26 \(\mu_B\), depending on the magnetic configuration and oxygens crystallographic positions. The largest magnetic moment, 0.26 \(\mu_B\), was found on the oxygen atoms, which belongs to two neighboring trimers in the fully FM solution (\( \uparrow m \uparrow - \uparrow m \uparrow \)).

The analysis of the calculation results presented in Tab. I shows that the magnetic contributions to the total energies of different solutions is not described by purely Heisenberg terms, like \(J_{ij} \vec{S}_i \cdot \vec{S}_j\). Indeed, if one compares the total energies of two solutions with the AFM order in the trimer (e.g. \( \uparrow m \downarrow - \uparrow m \downarrow \) and \( \uparrow m \downarrow - \downarrow m \uparrow \)) then the lowest will be the one with the AFM coupling between trimers. However, if one analyses another two solutions with the FM order in the trimer (e.g. \( \uparrow m \uparrow - \uparrow m \uparrow \) and \( \uparrow m \uparrow - \downarrow m \downarrow \)), it becomes clear that the lowest will be the one with the FM coupling between trimers. This is mostly related with the smaller magnetic moment on the Ru₄m ions in the \( \uparrow m \uparrow - \downarrow m \downarrow \) configuration, and it may be consequence of molecular orbital formation on trimers.

There are also two other contributions to the magnetic energy in addition to the conventional Heisenberg model, which have to be taken into account to describe magnetic properties of Ba₄Ru₄O₁₀. This is the development of the magnetic moments on Ru₄m (when the spins in the trimer are parallel) and on those oxygens, which are shared by two trimers (when the spins on the neighbor trimers are FM ordered). In each case the contribution to the total energy is \(-1M^2/4\), where \(M\) is the magnetic moment on the Ru₄m or O ions, and \(I\) - the Stoner parameter, which is approximately equals to the Hund’s rule exchange parameter for Ru (0.7 eV) and to 1.6 eV, as it was calculated in Ref. [13].

It is important to mention that only fully AFM state, corresponding to the lowest total energy, is insulating in the GGA approach, without including electronic correlations (Hubbard’s U). This is related to the band narrowing caused by the antiferromagnetism. The small gap insulating ground state, obtained in the magnetic GGA calculations agrees with a semiconducting temperature dependence of the electric resistivity, observed in Ref. [4].

V. CORRELATION EFFECTS

It is well known that for the description of the electronic and magnetic properties of the transition metal oxides one often needs to take into account strong Coulomb correlations, which can be incorporated in the calculation scheme via the LDA+U/GGA+U [8] or LDA+DMFT [16] formalism. We repeated the total energy calculation for some of the magnetic configurations within the GGA+U approx-
TABLE II: The total energies of the different magnetic solutions, as a result of the GGA+U calculations in the PWscf code. Total energies are given per formula unit. The notations are the same as in Tab. I.

| Magnetic configuration | Total energy (meV) | Band gap (eV) |
|------------------------|-------------------|--------------|
| m↓ m↑ - m↑ m↓          | 0                 | 0.26 eV      |
| m↑ m↑ - m↓ m↑          | 18.3              | Metal        |
| m↓ m↑ - m↑ m↑          | 19.8              | 0.09 eV      |
| m↑ m↑ - m↓ m↓          | 86.4              | Metal        |
| 0m↑ 0m↑ - 0m0          | 284.6             | Metal        |

VI. SUMMARY

Summarizing, on the basis of ab-initio and model calculations we obtained the explanation of the unusual magnetic properties of Ba$_4$Ru$_3$O$_{10}$, which consist of the Ru trimers coupled via corner-shared oxygens. The ground state is found to be an antiferromagnetic insulator, with antiferromagnetic ordering both within and between Ru trimers. The most surprising fact – the non-magnetic nature of the middle Ru ions in each trimer – is explained to be a result of a combined action of the formation of molecular orbitals in the Ru trimers and of charge redistribution, with the increasing $d$-states occupation on the middle Ru which is sandwiched between two other Ru ions with antiparallel spins. These unusual magnetic properties of Ba$_4$Ru$_3$O$_{10}$ are well explained by the band picture, and the electron correlations do not play significant role in the formation of this magnetic state.

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