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Modelling Magnetic Multipolar Phases in Density Functional Theory

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Multipolar magnetic phases in correlated insulators represent a great challenge for Density Functional Theory (DFT) due to the coexistence of intermingled interactions, typically spin-orbit coupling, crystal field and complex non-collinear and high-rank inter-site exchange, creating a complexed configurational space with multiple minima. Though the +U correction to DFT allows, in principle, the modelling of such magnetic ground states, its results strongly depend on the initially symmetry breaking, constraining the nature of order parameter in the converged DFT+U solution. As a rule, DFT+U calculations starting from a set of initial on-site magnetic moments result in a conventional dipolar order. A more sophisticated approach is clearly needed in the case of magnetic multipolar ordering, which is revealed by a null integral of the magnetization density over spheres centered on magnetic atoms, but with non-zero local contributions. Here we show how such phases can be efficiently captured using an educated constrained initialisation of the onsite density matrix, which is derived from the multipolar-ordered ground state of an ab initio effective Hamiltonian. Various properties of such exotic ground states, like their one-electron spectra, become therefore accessible by all-electron DFT+U methods. We assess the reliability of this procedure on the Ferro-Octupolar ground state recently predicted in Ba$_2$Mo$_5$O$_{16}$ (M = Ca, Mg, Zn) [Phys. Rev. Lett.127, 237201 (2021)].

Transition Metal Oxides (TMOs) with strong Spin Orbital Coupling effect (SOC) have attracted great attention due to the realization of unconventional magnetic phases, ranging from canted antiferromagnetic (AFM) orders observed in Ba$_2$NaOsO$_6$ [1–3] and Sr$_2$IrO$_4$ [4] to high-rank magnetic multipoles [5, 6]. Remarkable examples of multipolar orderings have been reported for URu$_2$Si$_2$ and NpO$_2$, where the onset of hidden ordered phase transitions have been connected with possible transitions towards a multipolar magnetic phase [6, 7]. More recently, 5d-based TMOs have attracted considerable interest due to the interplay between an unexpectedly high electronic correlation and SOC [8], with several reports providing evidence on the possible formation of multipolar ground states [9–11]. The majority of these works are based on microscopic low-energy effective Hamiltonians, solved by a variety of many-body methods. In fact, the search of multipolar magnetism by means of DFT electronic structure schemes with the +U correction inevitably faces the problem of being trapped in local minima corresponding to conventional dipolar solutions. In a pioneering work S.-T. Pi and coworkers [12] addressed this problem by calculating exchange interactions through flipping of the expansion coefficients of the onsite matrix expanded in terms of multipolar tensor components. The change in band energies reflects the energy cost of the corresponding flipping that can be afterwards mapped onto the exchange constant via Andersen force theorem (FT) [13]. However, this method becomes computationally intensive for multipolar “hidden” order systems, where the space of possible order parameters is large.

An alternative approach [14] is based on a FT formulated for the symmetry-unbroken paramagnetic electronic structure. The latter is obtained within the DFT+DMFT (dynamical mean-field theory [15, 16]) framework using the quasiatomic Hubbard-I (HI) approximation. Even for complex “hidden-order” systems, the full magnetic Hamiltonian can be derived using this FT-HI method from post-processing of a single DFT+HI calculation for the paramagnetic state [7, 17]. However, in contrast to DFT+U, the DFT+HI method cannot directly model the electronic structure of multipolar-ordered phases, since leading inter-site interactions in correlated insulators arising through hybridization of localized electrons (e. g. superexchange) are neglected in DFT+HI.

In this work, we develop a framework for calculating multipolar-ordered phases with the DFT+U method by initializing those calculations using the output provided by the FT-HI effective-Hamiltonian method. Within our scheme we first identify competing multipolar phases from an ab initio effective Hamiltonian. In this Hamiltonian, the relevant ground-state multiplet (GSM) of low-energy electronic states is represented by a pseudo-spin; the inter-site interactions between various moments of this GSM space are calculated by the FT-HI method [14] from the paramagnetic DFT+HI electronic structure. The effective Hamiltonian is then solved either by mean-field or by more sophisticated many-body techniques to obtain the transition temperatures and order
parameters for low-temperature phases.

Using explicit (Fock state) representations of the GSM many-electron states, as calculated by DFT+HI, one may transform those order parameters into on-site one-electron density matrices (ODM). Then, the DFT+U+SOC run is initialized with such ODMs corresponding to a chosen multipolar order. The selective initialization of the ODM allows the direct total energy calculations of a specific multipolar magnetic ground state, thus avoiding the risk of falling in a metastable dipolar state.

This framework enables the study of materials-specific electronic and magnetic properties of a genuine multipolar state in DFT at the atomic scale without adjustable parameters, and allows for a DFT-based analysis of the response of the multipolar ground state to external stimuli such as local structural distortions or doping, which are difficult to treat at DFT+HI level.

We employ the proposed computational protocol to study the competition between conventional dipolar and magnetic multipolar order in the cubic 5d2 double perovskites (DPs) Ba2MOSO6 (M = Ca, Mg, Zn) (BCOO, BMOO and BZOO from now on). With a t2g2 configuration exhibiting a S=1 spin state and an effective orbital moment l=1, the low energy physics of these spin-orbit coupled systems can be represented by a total effective momentum (pseudo-spin) J_{eff}=2, analogous to a single d-electron l = 2 level. As a consequence, in a cubic symmetry, the J_{eff}=2 level splits due to the remnant crystal field (RCF) into a lower E_g doublet and a higher-in-energy T_{2g} triplet. Since the non-Kramers E_g doublet is isomorphic to a e_g doublet, it carries no dipole moment, thus representing an ideal playground for the realization of high-rank multipole orders.

For these reasons BMOO DPs have recently been in the spotlight, but with conflicting experimental data. On one side muon spin resonance and thermodynamic anomalies show a clear phase transition below T^* ≈ 30-50 K, with broken time reversal symmetry and with large antiferromagnetic Curie-Weiss constant (Θ_{CW} ≈ 130 K) [18, 19], apparently consistent with a weak Néel spin ordering. On the other side no magnetic Bragg peaks were observed in neutron diffraction experiments, establishing an upper limit for the Os dipolar magnetic moment of ≈ 0.1μB [20]. Furthermore, possible quadrupolar orderings are also ruled out by the absence of tetragonal distortion as verified by x-ray diffraction measurements, up to ≈ 0.1 % of the volume, [20].

To shed some light on this complex scenario a few theoretical analyses have been reported providing robust evidence for the formation of a higher-rank order of the octupolar type [17, 21, 22]. We have recently evaluated the effective many-body Hamiltonian for these compounds using the FT-HI method [17] and obtained a ferro alignment of the x-y-z octupoles as their ground (ferro-octupolar, FO) state [17]. This FO order is enabled by a large RCF (an order of magnitude larger than the inter-site exchange interaction) suppressing competing dipolar orders and mediated by superexchange mechanism through O-p and Ba orbitals, [17]. An anti-ferro order of quadrupoles active within the e_g doublet was identified in Refs. [17, 23] as a competing phase. However, neither the electronic structure of the FO state nor its competing phases has been calculated in Ref. [17] due to the above mentioned limitations of DFT+HI.

In this work we make use of the previously obtained FT-HI Hamiltonians and ordered phases to carry out electronic structure calculations for the d^2 DP series using the constrained DFT+U methodology outlined above. We identify signatures of the multipolar order in one-electron spectrum. We also show that this DFT+U methodology is able to qualitatively capture the energetics of multipolar orders; in particular, it correctly predicts the relative magnitude of ordering energies along the 5d^2 DP series.

**METHOD**

We carried out DFT+U calculations by the Vienna Ab initio Simulation Package (VASP) [24, 25] using the generalized gradient approximation of Perdew, Burke and Ernzerhof. We included the onsite Coulomb repulsion at the Os d shells using the rotationally-invariant Lichtenstein formulation of DFT+U [26]. The on-site Coulomb vertex is specified by the Hubbard U and Hund’s rule coupling J; we employed U = 3.2 eV and J = 0.5 eV, in agreement with previous works [17]. The spin-orbit coupling was included in DFT and we estimated, via exact diagonalization of the atomic levels in the local Hamiltonian, the resulting SOC strength λ to be ~ 0.3 eV. For both BCOO, BMOO and BZOO the experimental lattice structures were used from references [19] and [18] respectively; the reciprocal space was sampled with a 6 × 6 × 6 k-mesh and an energy cutoff for the plane wave expansion of 600 eV was applied.

We employ the protocol outlined in the introduction to construct an appropriate ODM for a given multipolar order. This ODM is then used as starting guess for the constrained-ODM implementation of reference [27]. The starting ODM is derived from the order parameters of a given multipolar phase. To that end we represent the one-electron ODM corresponding to a given set of the order parameters in the pseudo-J space as follows

$$\rho_{mm'}_{\alpha}^{\text{ODM}} = \text{Tr} [ \rho_{MM'}(J) \hat{\rho}_\alpha(J) ]$$

where $\rho_{MM'}(J) = \langle JM|c_{m}^{\dagger}c_{m'}|JM'\rangle$ is the mm' matrix element of the ODM operator in the GSM basis |JM⟩, $c_{m}^{\dagger}$, $c_{m'}$ are the one electron creation and annihilation operators and $\hat{\rho}_\alpha(J)$ is the many-electron GSM density matrix (DM) at the site $\alpha$. The DM $\hat{\rho}_\alpha(J)$ is computed
from the ordered moments at site $\alpha$ for a given phase, 
$$\tilde{\rho}_\alpha(J) = \sum_{KQ} O_{KQ}^2(J) \langle \hat{O}_{KQ} \rangle_\alpha,$$
where $O_{KQ}$ is the spherical Hermitian tensor for given $J$ with rank $K = 1...2J$ and projection $Q$ [5] and $\langle \hat{O}_{KQ} \rangle_\alpha$ is its expectation value at the site $\alpha$ in a given ordered state. For the $d^2$ DP series we use the FO order parameters $\langle \hat{O}_{KQ} \rangle_\alpha$ that were obtained in Ref. [17] by solving the corresponding FT-HI effective Hamiltonians for pseudo-spin $J = 2$. To evaluate the matrix elements $\langle JM | c_m^\dagger c_m | JM' \rangle$ we employ the many-electron states of the GSM obtained in DFT+HI calculations [17] of those compounds; these states are expressed in the $ms$-electron Fock basis rendering such evaluation straightforward.

The many electron DM $\tilde{\rho}_\alpha(J)$ contains the nominal number of correlated electrons included in the effective Hamiltonian. DFT calculations, on the other hand, typically overestimate this counting because of the strong hybridization between the $d$ orbitals with the $O$ $p$ states. This enhanced electron counting is particularly problematic for osmium double perovskites, where the formal 5$d^2$ occupation is actually $\sim 6$ electrons in DFT [2, 28]. This unbalanced treatment of the ODM in DFT ($\rho_{ODM}$) is incongruous with the nominal $t_{2g}$ configuration adopted in the FT-HI effective Hamiltonian ($\rho_{HI}$) and needs to be corrected in order to have a consistent mapping between the two models. In our protocol (see Figure 1) we correct this problem by quantifying the nominal excess charge from a preliminary spin-unpolarized DFT calculation ($U=0$), were we obtain a reference DFT-ODM ($\rho_{DFT}$) which is a sum of the contribution coming from $t_{2g}$ orbitals ($\tilde{\rho}_{t_{2g}}$) and the $e_g$ ones ($\tilde{\rho}_{e_g}$) that include both the nominal $d^2$ electrons as well as additional electronic charge coming from hybridization effects.

As in VASP the $\rho_{DFT}$ is calculated in the global coordinate system one has to be sure this matches the local reference frame of the octahedral environment, to avoid mixing contributions from different orbitals. In our unit cells these reference frames coincide, and such a rotation is not needed. However, another change of basis was employed in order to move from the spherical harmonic basis of the $\rho_{HUB}$ to the cubic one defined in VASP (see Supplementary Materials (SM) [29]). At this point the hybridization contribution can be readily obtained by splitting the correlated $t_{2g}$ part from the $e_g$ one and defining

$$\rho_{ODM} = [\rho_{HI} + Id_{t_{2g}} \times Tr(\tilde{\rho}_{t_{2g}} - \rho_{HI})/6] + \tilde{\rho}_{e_g},$$

where $Id_{t_{2g}}$ is the identity acting on the $t_{2g}$ subspace only and the division by 6 is due to the spin degeneracy.

With the new ODM that correctly counts the added electrons, we have an educated initial guess on top of which we were able to run a ODM-constrained calculation at DFT+U+SOC level, with the requirement of keeping the total magnetization along the global components as obtained with $Tr(S_i \rho_{ODM})$, where $S_i$ is the dipolar spin operator. As mentioned above, the nature of the FO ground state is such that all these components average to zero and as such have to be initialised, considering that any other initialization would act as localized magnetic field on the osmium atom, thus pushing towards a conventional dipolar solution. After the initial ODM-constrained calculation, we performed a consistency safety check to control that the correct number of electrons is maintained in the ODM, and subsequently performed a full self-consistent calculation to obtain the FO DFT+U+SOC solution, starting from the pre-converged wavefunction generated in (4) (see SM [29]).

FIG. 1. Scheme of the constrained multipolar ODM protocol: 1) FT-HI optimization and calculation of the FT-HI ODM $\rho_{HUB}$; 2) Preliminary DFT + $U$ ($= 0$ eV) calculation for the $t_{2g}$ and $e_g$ density; 3) Hybridization correction and initialization of a $\rho_{ODM}$ consistent with $\rho_{HUB}$; 4) ODM-constrained DFT+$U$+SOC calculation with the new $\rho_{ODM}$; 5) Full self-consistent DFT+$U$+SOC run starting from the pre-converged wavefunction generated in (4) (see SM [29]).

**RESULTS**

We start by discussing the FO solution obtained by DFT using the proposed approach and comparing it with the dipolar AFM-110 state. After that, we present the DFT estimations of the intersite exchange couplings and
explore the possibility for tetragonal transition.

The Ferro-octupolar phase

To extract useful information from the DFT output we fit the final DFT-ODM $\rho_{ODM}$ to the basis of 2-electrons average of multipolar moments $\langle O_m^N(J) \rangle$:

$$\rho_{ODM} = \sum_{n,m} a_m^n \langle O_m^N(J) \rangle,$$

where $a_m^n$ are the fitting coefficients.

We find indeed that the octupolar $O^{-2}_3$ operator remains non zero, even if not fully saturated (see Table I), providing clear evidence of the capability of DFT to model a genuine multipolar order. Most of the other magnetic multipoles are almost zero, apart from $O^2_2$, $O^6_2$ and hexadecapoles, the latter with values that do not impact on the quality of the fit and are therefore neglected from now on. The non zero values obtained for quadrupolar $O^2_2$ and $O^6_2$ terms are a consequence of switching-off all symmetry as required in VASP-based SOC calculations, and do not imply any tendency of the system to undergo a cubic-to-tetragonal transition (we have verified that the systems prefers to preserve the cubic symmetry, see Sec. "Tetragonal Distortions"). We further confirmed the role of symmetry by performing a non-magnetic DFT + SOC calculation from which we extracted the expansion coefficients finding values of $\sim 0.30$ for $O^2_2$ and $\sim 0.43$ for $O^6_2$, then subtracted from the pure DFT + U + SOC values to partially compensate the above-mentioned effect (see Table I for the re-scaled values). The final ODM contains also spurious terms coming from the hybridization between Os and O atoms, such that only 90 % of the total Hilbert space from which $\rho_{ODM}$ is constructed can be correctly mapped in our tensor fit; this imbalance can be easily adjusted by subtracting the non-magnetic DFT+SOC ODM calculation from the final ODM. A visual representation of the obtained FO state is given in Fig. 2 in terms of the magnetization density isosurface along the crystallographic $z$ direction, showing a FM order of magnetic octupoles.

After clarifying the basic multipolar character of the FO order with DFT+U+SOC, we move forward to the analysis of the differences between the FO and the competing dipolar AFM-110 solutions in terms of stability, electronic structure and magnetic properties. Our results show that the FO phase is lower in energy than the dipolar phase, with a difference of $\approx 41, 45$ and 43 meV/f.u. for BCOO, BMOO and BZOO respectively.

The density of states and the band structure displayed in in Fig. 3 show an insulating electronic ground state for both phases with strong qualitative differences in the character of the unoccupied states: FO-ordered BCOO exhibits one broad unoccupied $t_{2g}$ band, which is split into three peaks in the AFM phase. The FO insulating energy gap, 1.16 eV, is significantly larger than the corresponding AFM, 0.55 eV, see Table II). These are well-defined electronic characteristics that identify a clear difference between these two magnetic orders, which can be verified experimentally.

From a magnetic point of view, the difference between the FO and AFM phases become transparent from the ordered magnetic dipolar moments and the magnetization density. The dipolar AFM-110 posses a local spin and orbital moment on the osmium atom of $m_S \sim 1.4 \, \mu_B$, $m_L \sim 0.7 \, \mu_B$ summing up to $m_J \sim 0.7 \, \mu_B$, while they average to zero in the FO case. The diversity between the FO and AFM-110 magnetic orderings is also reflected in the magnetization density along $z$ Figure 2.
FIG. 3. (Color online) Comparison of the electronic properties of BCOO for the two different magnetic phases FO and AFM-110: (a) density of states and (b) band structure.

| Compound          | FO     | AFM-110 |
|-------------------|--------|---------|
| Ba$_2$CaOsO$_6$   | 1.16   | 0.55    |
| Ba$_2$MgOsO$_6$   | 1.16   | 0.52    |
| Ba$_2$ZnOsO$_6$   | 1.24   | 0.64    |

TABLE II. Energy Gap for the different magnetic phases show how the magnetic and electronic properties are strictly related in these compounds. An enhancement of the band gap is found in the multipolar ordered phases, as compared with the conventional dipolar solution.

Intersite exchanges

The driving force that couples rank 3 time-odd multipolar operators and stabilizes the multipolar magnetic order in these compounds is the intersite exchange interaction between osmium atoms mediated via superexchange mechanism [17]. To further analyze and compare our result to previous theoretical findings, we start by mapping total energy differences obtained with our constrained ODM approach for distinct multipolar ordered phases, to the following model Hamiltonian

$$H = \sum_{\langle ij \rangle} \sum_{KQK'Q'} V_{KK'}^{QQ'}(\mathbf{R}_{ij}) O_K^Q(\mathbf{R}_i) O_{K'}^{Q'}(\mathbf{R}_j),$$  \hspace{1cm} (4)

where $O_K^Q(\mathbf{R}_i)$ are the Hermitian spherical tensor [5] for $J=2$ of the rank $K = 1...4$, $Q = -K,...,K$ and the sum runs over all nearest-neighbor $\langle ij \rangle$ Os-Os bonds. This Hamiltonian is a reduced form of the one used in Ref. [17], as the remnant crystal field is already taken into account self-consistently. In ref [17] the Hamiltonian is further simplified to an effective pseudo-spin Hamiltonian acting on the low lying doublet. Here we follow the same reasoning and rewrite it as

$$H = \sum_{\langle ij \rangle} \sum_{\alpha\beta} J_{\alpha\beta}(\mathbf{R}_{ij}) \tau_\alpha(\mathbf{R}_i) \tau_\beta(\mathbf{R}_j),$$  \hspace{1cm} (5)

where $\tau_\alpha$ is the corresponding pseudo-spin-$1/2$ operator and, for $\alpha = y$, it coincides with $O_{-2}^2$. To calculate the value of the intersite exchange interaction $J_{yy}$ between time-odd multipoles, we employed the constrained ODM for an antiferro-octupolar (AFO) configuration, with the $O_{-2}^2$ aligned ferromagnetically in [001] planes and AFM out-of-plane. Our result shows that the FO phase is still lowest in energy and that, while the AFO has different signs of the $O_{-2}^2$ operators, all other saturated magnetic multipolar moments keep their values unchanged, with a difference of $\sim 12\%$ in the tensor fit coefficients for $O_0^2$ and $\sim 1\%$ for $O_{2}^2$. By assuming that these changes are not as significant as the ones brought by $O_{-2}^2$ in the total energy, we calculate the exchange constant as

$$J_{yy} = \frac{E_{FO} - E_{AFO}}{2}. \hspace{1cm} (6)$$

Our results, in Table III, are in qualitative agreement with FT-HI results but are overestimated by a multiplicative factor ($\approx 4$), to be attributed to the different electron counting in the two approaches, as discussed above and in a previous work [2]. Still, the relative strength of the computed $J_{yy}$ in the three compounds is reproduced rather consistently.

Tetragonal Distortions

We conclude by reporting data on the possibility for these systems to undergo a cubic-to-tetragonal distortion. Osmate DPs have been theoretically predicted to host quadrupolar moments and recently it has been
proved that unaxial strain along the z axis might lead to a suppression of the FO transition temperature as a consequence of the interplay of a weakened octupolar exchange interactions and the produced transverse field [21]. Minuscule tetragonal distortions are expected to activate transitions between the ground state and the excited singlet, ultimately detectable in Inelastic Neutron Spectra measurements [17]. We investigated whether such distortions might induce a transition towards a competing favourable solutions by studying the change in energy as a function of the $\delta = c/a - 1$ by switching the lattice parameter in steps of 0.01 Å in both positive and negative directions from the experimental value, while keeping the volume fixed (we further compared our result with an automated conjugate gradient minimization algorithm with both fixed and variable lattice parameters).

The results for BCOO are collected in Figure 4 (data for BMOO and BZOO are available in the SM [29]). Our DFT data clearly highlights no deviation from the cubic symmetry for both FM and AFM octupolar FO and AFO phases. In contrast, the AFM-110 phase shows an expansion along the z axis with change in lattice parameter of the order of $\sim 1\%$ and the appearance of JT distortions, both of which would have been detected by X-ray diffraction measurement. These considerations apply also for BMOO and BZOO. An important point is the dependence of the magnetic multipoles on the magnitude of the strain, here analysed for BCOO. We obtain for the $O^2_2$ moment a sharp linear dependence as function of $\delta$ in good qualitative agreement with the DFT+HI results (see Supplementary related to Ref. [17]). We further observe a linear behavior of the $O^{-2}_3$ moment which suggests a strengthening of the FO phase upon tensile stress, with a possible enhancement of the corresponding transition temperature (indeed the evaluation of $J_{yy}$ for the $\delta = +0.01$ structure gives a value $\sim 3\%$ larger). For completeness, we find that the multipolar moment $O^2_2$ remains constant.

These results suggest that tetragonal distortions might play a decisive role on the magnetic properties of these compounds and that further investigations, also via experimental analysis, might be a worthy research path.

| Compound          | DFT      | FT-HI [17] |
|-------------------|----------|------------|
| Ba$_2$CaOsO$_6$   | -10.70   | -2.98      |
| Ba$_2$MgOsO$_6$   | -10.10   | -2.93      |
| Ba$_2$ZnOsO$_6$   | -8.47    | -1.71      |

**TABLE III.** Comparison between DFT and FT-HI intersite exchange constants for the effective Hamiltonian of equation 5.

In conclusion, we have proposed a protocol for obtaining magnetic multipolar ordered phases in DFT+U+SOC based on the constrain of the occupation density matrix as obtained from DFT+DMFT within the FT-HI approximation. We applied this procedure to $5d^2$ double perovskites, correctly reproducing the FO ordering of time-odd octupoles. We proved the FO phase to be the ground state and compared our results with dipolar AFM-110 configuration, that would be the "conventional" DFT ground state solution. We calculated the interstite exchange interaction constant, finding reasonable agreement with other theoretical results. Finally, we explored the possibility for these systems to undergo a tetragonal distortion, finding no such evidence and revealed the dependence of the dipolar moments upon strain from purely ab initio perspective.

**II. ACKNOWLEDGMENTS**

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[29] Supplementary Material.
Supplementary material for 'Modelling Magnetic Multipolar Phases in Density Functional Theory'

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I. FIRST PRINCIPLES METHODS

The two-electron Onsite Density Matrix as obtained via FT-HI method (Wien2k) is in spherical harmonic basis \((Y_{-2}^2, Y_{-1}^2, Y_{1}^0, Y_{1}^2, Y_{1}^{-1}, Y_{1}^0, Y_{2}^0, Y_{1}^1, Y_{2}^2, Y_{1}^{2}, Y_{1}^{-2}, Y_{2}^{-2}, Y_{1}^{-2})\) and is given by the following real part:

\[
\begin{pmatrix}
0.2112 & 0 & 0 & 0 & -0.1609 & 0 & -0.1527 & 0 & 0 & 0 \\
0 & 0.4597 & 0 & 0 & 0 & 0 & -0.0532 & 0 & 0 & 0 \\
0 & 0 & 0.0069 & 0 & 0 & 0 & 0 & -0.0532 & 0 & 0 \\
0 & 0 & 0.1977 & 0 & 0.1093 & 0 & 0 & 0 & 0 & -0.1527 \\
-0.1609 & 0 & 0 & 0 & 0.1244 & 0 & 0.1093 & 0 & 0 & 0 \\
0 & 0 & 0 & 0.1977 & 0 & 0.1093 & 0 & 0 & 0 & 0 \\
0 & -0.0532 & 0 & 0 & 0 & 0.0069 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & -0.1527 & 0 & -0.1609 & 0 & 0 & 0 & 0.2113 \\
\end{pmatrix}
\]

and imaginary component

\[
\begin{pmatrix}
0 & 0 & 0 & -0.0367 & 0 & 0 & 0 & 0 & 0 & 0.2659 \\
0 & 0 & 0 & 0.2270 & 0 & 0.1880 & 0 & 0 & 0 & -0.2660 \\
0.0367 & 0 & 0 & 0 & 0 & 0 & 0 & -0.0318 & 0 & 0 \\
0 & -0.2270 & 0 & 0 & 0 & 0 & 0 & 0 & 0.0318 & 0 \\
0 & 0 & 0.0270 & 0 & 0 & 0 & 0 & 0 & 0 & -0.1880 \\
0 & 0 & 0.0318 & 0 & 0 & 0 & 0 & 0 & 0 & -0.2270 \\
0 & 0 & 0.0318 & 0 & 0 & -0.0270 & 0 & 0 & 0 & 0.0367 \\
0.2659 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & -0.0367 & 0 \\
0 & 0.2660 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\end{pmatrix}
\]

We employed a change of basis from spherical to cubic as in the VASP conventional form \((d_{xy}, d_{yz}, d_{z^2}, d_{x^2-y^2}, d_{xy}, d_{yz}, d_{z^2}, d_{x^2-y^2})\) using the following transformation matrix (without considering the spin degeneracy)

\[
\begin{pmatrix}
0 & 0.70711 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & -0.70711 \\
0 & 0 & 0.70711 & 0 & 0 & 0 & 0.70711 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0.70711 & 0 & 0 & -0.70711 & 0 & 0 & 0 & 0 \\
0.70711 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0.70711 & 0 \\
\end{pmatrix}
\]

where the columns are alternated real and imaginary parts. After having added the contribution coming from the \(e_g\) orbitals \((0.61347)\) and a charge correction on the \(t_{2g}\) as explained in the main text we obtain the final occupation matrix given in input with the flag \(OCCEXT = 1\), as reported below.
where the Spin Component 1,2,3,4 refer to \( (\uparrow\uparrow), (\uparrow\downarrow), (\downarrow\uparrow) \) and \( (\downarrow\downarrow) \) spinor components respectively. After the final self consistent calculation we obtain the resulting ODMs for BCCO, BMOO and BZOO, which are listed in the following.

### A. Final Onsite Density Matrix BCOO

In the following are listed the final ODM after the self-consistent calculations obtained for BCOO, BMOO and BZOO in the cubic experimental structure.

| Spin component 1 | 0.62239 | 0.0000 | 0.04504 | 0.0000 | 0.0000 | 0.0000 | 0.0000 | 0.0000 | 0.0000 | 0.04340 |
|------------------|---------|--------|---------|--------|--------|--------|--------|--------|--------|--------|
|                  | 0.0000  | 0.62239 | 0.0000  | 0.22700 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|                  | 0.04504 | 0.0000  | 0.60912 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.00686 |
|                  | 0.0000  | 0.22700 | 0.0000  | 0.6239  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|                  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.60912 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |

| Spin component 2 | 0.0000  | -0.13100 | 0.0000  | 0.22695 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|------------------|---------|----------|---------|---------|--------|--------|--------|--------|--------|--------|
|                  | 0.13100 | 0.0000  | -0.02249 | 0.0000  | 0.03900 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|                  | 0.0000  | -0.02249 | 0.0000  | 0.03762 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|                  | 0.22700 | 0.0000  | -0.03762 | 0.0000  | 0.02170 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|                  | 0.0000  | 0.03895  | 0.0000  | -0.02170 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |

| Spin component 3 | 0.0000  | 0.13100 | 0.0000  | 0.22700 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|------------------|---------|----------|---------|---------|--------|--------|--------|--------|--------|--------|
|                  | -0.13100| 0.0000  | -0.02249 | 0.0000  | 0.03895 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|                  | 0.0000  | -0.02249 | 0.0000  | 0.03762 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|                  | 0.22695 | 0.0000  | -0.03762 | 0.0000  | 0.02170 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|                  | 0.0000  | 0.03900  | 0.0000  | -0.02170 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |

| Spin component 4 | 0.62244 | 0.0000  | -0.04504 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|------------------|---------|----------|---------|---------|--------|--------|--------|--------|--------|--------|
|                  | 0.0000  | 0.62244 | 0.0000  | -0.02270 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|                  | -0.04504 | 0.0000  | 0.60912 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.00686 |
|                  | 0.0000  | -0.22700 | 0.0000  | 0.62244 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |
|                  | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.60917 | 0.0000  | 0.0000  | 0.0000  | 0.0000  | 0.0000  |

...
### B. Final Onsite Density Matrix BMOO

| Spin Component 1 | 0.6167 0.0001 -0.0134 -0.0001 -0.0000 -0.0000 -0.0001 -0.0000 0.0004 0.0379 |
|------------------|--------------------------------------------------|
|                  | -0.0000 -0.0001 -0.0000 0.0004 0.0379           |
| Spin Component 2 | -0.0001 0.1095 -0.0001 -0.1959 -0.0000 -0.0000 -0.0000 -0.0000 -0.1104 -0.0000 |
|                  | 0.1094 -0.0000 0.0066 -0.0001 -0.0116 0.0001 -0.0000 -0.0000 0.0000 0.0000 |
|                  | 0.0000 0.0000 -0.0000 -0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 |
| Spin Component 3 | 0.0000 -0.1094 -0.0001 -0.1959 0.0000 0.0000 0.0000 0.0000 -0.1107 -0.0000 |
|                  | 0.0195 -0.0000 0.0066 -0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 |
|                  | 0.0000 0.0000 0.0000 -0.0001 0.6512 -0.0335 -0.0000 0.0016 0.0000 0.0000 |

### C. Final Onsite Density Matrix BZOO

| Spin Component 1 | 0.6324 0.0007 -0.0155 0.0001 -0.0000 -0.0000 -0.0000 -0.0000 -0.0001 0.0335 |
|------------------|--------------------------------------------------|
|                  | 0.0007 0.6322 -0.0000 -0.2040 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 |
|                  | 0.0155 -0.0000 0.6512 -0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 |
|                  | 0.0001 -0.2040 -0.0000 0.6432 -0.0001 0.0001 -0.1123 -0.0000 0.0000 0.0000 |
| Spin Component 2 | 0.0000 0.1123 0.0000 -0.2041 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 |
|                  | -0.1119 -0.0000 0.0077 -0.0003 -0.0134 0.2043 0.0000 -0.2090 0.0009 -0.0167 |
|                  | -0.0000 0.0000 0.0000 -0.0290 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 |
|                  | -0.2040 -0.0003 0.2290 -0.0001 -0.0167 0.1124 0.0008 -0.0078 0.0001 -0.0134 |
| Spin Component 3 | 0.0000 -0.0134 0.0000 0.0000 0.0167 -0.0000 0.0000 0.0000 0.0000 0.0000 |
|                  | 0.0000 0.0000 -0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 |
| Spin Component 4 | 0.6321 -0.0007 0.0155 0.0000 0.0000 -0.0000 -0.0000 -0.0000 -0.0000 0.0335 |
|                  | -0.0007 0.6321 0.0000 0.2040 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 |
|                  | 0.0155 0.0000 0.6512 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 |
|                  | 0.0000 0.2040 0.0001 0.6322 0.0001 0.0001 0.1123 0.0000 0.0000 0.0000 |
|                  | 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 |
II. DOS AND BANDSTRUCTURES

A. BMOO

Supplementary Figure 1: (Color online) Comparison of the electronic properties of BMOO for the two different magnetic phases FO and AFM [110]. The change in Energy Gap is clear in the Density of States (a) and more explicit in the Bandstructures for the FO phase (b) and the AFM [110] phase (c).

B. BZOO

Supplementary Figure 2: (Color online) Comparison of the electronic properties of BZOO for the two different magnetic phases FO and AFM [110]. The change in Energy Gap is clear in the Density of States (a) and more explicit in the Bandstructures for the FO phase (b) and the AFM [110] phase (c).
III. MAGNETIZATION DENSITY PLOTS X AND Y COMPONENTS BCOO

Supplementary Figure 3: (Color online) Plot Magnetization density for the Octupolar FM phase of BCOO of the x and y components in a) and b) respectively.

IV. TETRAGONAL DISTORTION BZOO AND BMOO

A. Plot BMOO

Supplementary Figure 4: (Color online) Energy as a function of $\delta = c/a - 1$ for BMOO. The total energies are scaled to have the zero for the lowest energy value.
B. Plot BZOO

Supplementary Figure 5: (Color online) Energy as a function of $\delta = c/a - 1$ for BZOO. The total energies are scaled to have the zero for the lowest energy value.

V. DIPOLAR CALCULATIONS

In the following Table I are listed the total energies for dipolar configurations calculated within DFT+U+SOC using the +U correction in the rotationally invariant Dudarev’s scheme and $U = 3.4$ eV. The ”xy” notation refers to magnetic moments aligned ferromagnetically within the xy planes along the [abc] crystallographic direction, and anti-ferromagnetically in between adjacent xy planes. Otherwise the magnetic moments are to be read as anti-ferromagnetically aligned within the xy plane, with magnetic moments pointing along the [abc] crystallographic direction, and ferromagnetically aligned in adjacent xy planes.

| Configuration | Energy (eV) |
|---------------|-------------|
| AFM [001]     | -66.683     |
| AFM-xy [001]  | non converged |
| AFM [100]     | -66.710     |
| AFM-xy [110]  | -66.710     |
| AFM [110]     | -66.708     |
| AFM-xy [110]  | -66.713     |
| AFM [111]     | -66.688     |
| AFM-xy [111]  | -66.708     |
| FM [001]      | non converged |
| FM [100]      | -66.696     |
| FM [110]      | -66.701     |
| FM [111]      | -66.701     |

Supplementary Table I: Comparison total energies calculations with dipolar magnetic configuration.