Exchange parameters in Fe-based molecular magnets

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Abstract

The calculation of interatomic magnetic exchange interactions entering the Heisenberg model from the standpoint of the density functional theory (DFT) is outlined for two Fe-based molecular magnets: a trinuclear complex with a Schiff base ligand, which makes an antiferromagnetically coupled frustrated system, and a model bipyrimidine-connected planar network of Fe ions. First-principles electronic structure calculations are performed using the real-space method SIESTA and the full-potential linearized augmented plane wave FLAPW method FLEUR, correspondingly. We discuss the application of fixed spin moment technique for preparing the system in a given magnetic configuration, and the effect of intraatomic Coulomb correlation, approximated by the LDA+U technique, on the values of interaction parameters.

Key words: Ab initio calculations, single-molecule magnets, magnetic interactions

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1 Introduction

Since the first systematizations of their chemistry and basic properties by, e.g., Kahn in 1993 [1], molecular magnets are now developing into a promising class of magnetic nanomaterials with a great potential in applications in areas such...
as magnetic storage, quantum computing, or magneto-optical devices. An up-
to-date review on microscopic properties (spin density, exchange interactions,
magnetic anisotropy) of molecular magnets accessible in experiment and from
first-principles calculations, is going to appear in Ref. [2]. Its major part is sum-
marized as an internet publication [3]. In the present work we address the issue of
extracting the interatomic exchange interaction parameters entering the Heisen-
berg model form the density functional theory (DFT). We discuss two chemical
systems of different complexity. The actual calculations are done by two differ-
ent methods, SIESTA [4] using compact and strictly confined atom-centered basis
functions (see Ref. [5] for details), and the FLEUR [6] code, a realization of
the highly accurate full-potential linearized augmented plane wave (FLAPW)
method. Both calculations used the generalized gradient approximation to the
exchange-correlation functionals.

We skip the subtleties in the formulation of exchange parameters within the DFT;
a detailed discussion can be found in Ref. [3]. Basically, we refer to total ener-
gies in different specially prepared magnetic configurations, calculated from first
principles. Sec. 2 covers the extraction of meaningful results for a Fe-trinuclear
system by applying the fixed spin moment formalism. Sec. 3 addresses a model
Fe-binuclear system and, specifically, the importance of intraatomic correlation,
treated beyond the conventional DFT formalism.

2 Fixed Spin Moment treatment of a Fe-trinuclear system

One of the ways to arrive at the values of interaction parameters in a magnetic
system whose properties are well described by the Heisenberg model is to compare
total energies for different magnetic configurations. This is rather straightforward
if, say, ferromagnetic (FM) and antiferromagnetic (AFM) states are well defined
and at least metastable. A more complicated case is a frustrated system, an
example of which is made of three Fe atoms incorporated into organic molecule.
We consider specifically Fe$_3$(OAc)$_3$L$_3$, a synthesized by Boskovic et al. [7] derivate
of the Schiff base H$_2$L = salicylidene-2-ethanolamine. The molecular unit used in
the electronic structure calculation by the SIESTA method is shown in Fig. 2 (a) of
Ref. [7]. The magnetization measurements reported in Ref. [7] indicate an (almost
AFM) ground state with the total spin $S$=1/2, coming about from the interaction
of three $s$=5/2 spins of Fe(III) ions. Experimentally, the interaction parameters
$J$ of the Heisenberg model, introduced in Ref. [7] as $H = -2 \sum_{\text{Fe pairs}} J_{ij} \mathbf{s}_i \cdot \mathbf{s}_j$, were
reported to be necessarily different for an acceptable fitting of data, even as the
molecule nearly maintains a trifold symmetry axis: $J_{ij} = -15.1$ K, $-13.6$ K, and
$-12.4$ K.

We proposed earlier [8] to use the Fixed Spin Moment (FSM) method [9] for
selecting a magnetic configuration of a molecular magnet (a six-center “ferric
wheel”), improving simultaneously the stability of the electronic structure calcu-
Two issues might need further discussion. The HOMO-LUMO gaps shown in the middle panel of Fig. 1 are different in the FSM formalism for two spin directions, because the fixing of the total spin amounts to an imposition of an external magnetic field, moving apart the chemical potentials in two spin channels. Normally this would require the consideration of a Zeeman term when dealing with total
energies. However, for a system with a common band gap in both spin channels a unique chemical potential can be found for both spin directions, therefore the FSM scheme would simply fix one or another of the metastable magnetic configurations at no additional energy cost. One can see that, indeed, a common band gap can be found in all situations relevant for our discussion, i.e. with FSM = 1, 5, and 15 \( \mu_B \), thus justifying the extraction of the \( J \) value above. Another observation concerns the local magnetic moments at Fe sites: as the FSM value is changing, and even as the saturation of magnetization (at FSM = 15 \( \mu_B \)) is achieved, the local Fe magnetic moments merely flip, essentially maintaining their magnitudes of 4 \( \mu_B \), and not 5 \( \mu_B \). The nominal value of \( s=5/2 \), associated with a single Fe atom, comes about due to magnetic polarization of ligands. This situation is identical to that described earlier for “ferric wheels” [3,8,10]. In spite of the very different structure of organic ligands, both systems have similarities in the nearest neighbourhood and in the charge state of Fe atoms (O\(_6\) twisted octahedral coordination in “ferric wheels” vs. slightly distorted octahedral O\(_5\)N coordination in the present system).

3 Fe-binuclear system: Effect of intraatomic correlation

We turn now to the discussion of the effect of on-site intraatomic Coulomb correlation on interatomic exchange parameters. Our objective was to find a compact and yet realistic system, for which an electronic structure calculation could be done with the use of an ultimately accurate method within the DFT, that is, the FLAPW method, and to apply a phenomenological correction by the LDA+U formalism [11] on top of it. Specifically, we apply the realization of the method in the FLEUR code [6]. We derive a simple model system from the Fe-binuclear complex described by Real et al. [12] being subject to previous studies [13]. We neglect for the moment the spin crossover property, which makes this system particularly interesting, and make our calculation for the high-spin configuration on both Fe centers. The chemical formula of the substance is \([\text{Fe(bt)}(\text{NCS})_2]_2\text{-bpym}\), with bt = 2,2'-bithazoline and bpm = 2,2'-bipyrimidine, and the molecular unit as shown on the left of Fig. 2. We also performed electronic structure calculation for this substance in its molecular and crystallized form by the SIESTA method, allowing structure relaxation, the results of this study will be reported elsewhere. For the present simulation, as the molecule is yet too large and too low-symmetric for an affordable FLAPW calculation, we simplify the molecular units, substituting the bt fragments by bpm – that is realistic, as this is encountered in related compounds – see, e.g., [13]. Moreover we cut and connect the (NCS) tails into –N=C=N– chains that makes a dense 3-dimensional lattice out of planar Fe–bipyrimidine groups (Fig. 2, right panel). The nearest neighbourhood of each Fe center remains a slightly distorted N\(_6\) octahedron, as was the case in the original molecular unit. An idealization is that all N–Fe–N bonds make now right angles, and the repeated bipyrimidine fragments are coplanar.
Fig. 2. Molecular unit of the Fe-binuclear system \( \text{Fe[(bt)(NCS)]}_2 \text{bpym} \) described in Ref. [12] (left) and a simplified model thereof, used in the calculation (right). The Fe atom is in the middle of a slightly distorted \( N_6 \) octahedron.

The calculated densities of states (per unit cell and per one Fe site) in Fig. 3 clearly indicate a hybridization of Fe3d with the 2p states of neighbouring nitrogen atoms. Since the minority-spin Fe 3d states are not completely empty, the local magnetic moment is less than 5 \( \mu_B \). As for the previous trinuclear system, the magnetic moment induced on nitrogen neighbours substantially contributes to the net value, associated to each Fe site. This resolves, at least in part, a controversy between the spin value \( s=5/2 \), expected at the Fe site from formal valence considerations, and the local magnetic moment of our calculation. The cumulative magnetic moment, distributed over the Fe site and its neighbours, behaves like a “rigid” spin in the sense of the Heisenberg model. However, one can expect that a stronger localization of this distributed magnetic moment at the Fe site will be enforced by an inclusion of intraatomic Coulomb correlation, underestimated in conventional DFT calculations. In order to check this possible effect, we brought in an additional Coulomb correlation applying a semi-empirical...

Fig. 3. Total density of states per unit cell (outer solid line) and local DOS per Fe site (evaluated within the muffin-tin sphere of 2.3 Bohr; shown by filled black) densities of states calculated for the model Fe-binuclear system, shown in the right panel of Fig. 2, for parallel orientation of Fe magnetic moments. Left panel: LDA calculation, right panel: LDA+\( U \) calculation with \( U=4 \) eV.
Table 1
Local magnetic moments at the Fe site (in $\mu_B$, within muffin-tin sphere of 2.3 Bohr); total magnetic moments per Fe center, total energy difference between FM and AFM configurations, and corresponding extracted values of the Heisenberg exchange parameter $J$, assuming nominal spin value $s=5/2$.

| $U$ (eV) | $M_{Fe}$ | $M/Fe$ |
|----------|----------|--------|
|          | FM       | AFM    | FM $E_{FM} - E_{AFM}$ (meV) | $J$ (K) |
|          | 3.62     | 3.61   | 102.5 | $-95$ |
| 4        | 3.93     | 3.92   | 76.8  | $-71$ |

“LDA+$U$” scheme [11]. We choose the Coulomb interaction constant $U = 4$ eV, which seems reasonable for correlated Fe compounds [14,15]. One sees in the DOS of Fig. 3 (right panel) the known effect of the “+$U$” correction to lower the occupied states in energy whereas shifting unoccupied ones upwards.

The values of magnetic moments and exchange parameters $J$, determined by the formula $E = -J s_1 s_2$, are listed in the Table. It is noteworthy that, whereas the total magnetic moment per Fe site is increased in the LDA+$U$ calculation, the local Fe magnetic moment still does not exceed $4 \mu_B$. One can conclude that the spatial distribution of magnetic density is not confined to the Fe atom even as the intraatomic Coulomb correlation is enhanced; instead, the magnetization maintains its complex spatial distribution, spilling onto neighbouring N atoms.

As the intraatomic Coulomb correlation shifts apart on the energy scale the centers of gravity of occupied and unoccupied states, it reduces the interatomic exchange parameter, because the latter (in one of its formulations, see [16]) has corresponding energy differences in the denominator. The same trend prevails in our case even as we estimate the values of $J$ by comparing the total energies of two magnetic configurations. The effect of Coulomb correlation, included on top of a straightforward DFT calculation, is therefore always towards lowering the absolute values of Heisenberg exchange parameters. The same conclusion has been drawn by Boukhvalov et al. in their calculations for “Mn$_{12}$” [17] and “V$_{15}$” [18] molecular magnets. Comparing with the results of the previous Section, one can conclude that the estimation of exchange parameters in molecular magnets, that is already qualitatively correct (in sign and order of magnitude) in a conventional DFT calculation, can be brought closer to the experimental estimates by reducing their numerical values due to additionally included intraatomic Coulomb correlation.

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[19] It should be explained that the Fe-trinuclear complex reported in Ref. [7] crystallizes with two molecular units per primitive cell, which are slightly different. In particular, there is a formal disproportionality in the nominal electron number between these two fragments. As a consequence, a single molecular unit we took for our calculation possesses an odd number of electrons. Therefore only odd FSM values correspond to integer number of electrons in both majority- and minority-spin channels and may lead to an appearance of HOMO-LUMO gaps, shown in the middle panel of Fig. 1. However, the even FSM values can be formally applied in the calculation as well, notwithstanding that they do not possess the HOMO-LUMO gap. Technically we applied an artificial broadening of molecular orbitals’ eigenvalues by 400 K. Some of the even FSM numbers allow an easily converging calculation, with the total energy and local moments which follow the general trend. Such results are added in the left and right panels of Fig. 1.