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Single Molecule Magnets Grafted on Gold: Magnetic Properties From Ab Initio Molecular Dynamics†

Alessandro Lunghi\textsuperscript{a}, Marcella Iannuzzi\textsuperscript{b}, Roberta Sessoli\textsuperscript{a} and Federico Totti\textsuperscript{*a}

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A computational characterization of the Single Molecule Magnet (SMM) [Fe\textsubscript{4}(AcS(CH\textsubscript{2})\textsubscript{3}(CH\textsubscript{2}O)\textsubscript{3})\textsubscript{2}(dpm)\textsubscript{6}](Fe\textsubscript{4}C\textsubscript{5}) with Ac = CH\textsubscript{3}CO and Hdpm = dipivaloylmethane, grafted on Au(111) surface (Fe\textsubscript{4}C\textsubscript{5}@Au(111)) is presented. For the first time Ab initio Molecular Dynamics (AIMD) calculations have been used to study the evolution of the structural properties of a SMM once adsorbed on a metallic substrate. Important structural rearrangements induced by the grafting process can be observed thanks to the AIMD approach. This allows to elucidate some possible mechanisms that govern the SMM’s magnetism on surface, which wouldn’t be revealed by the standar static approach. The sampling of the SMM@Au(111)’s configurational space at finite temperature, together with magneto-structural correlations, makes it possible to study the fluctuation amplitude of magnetic properties, thus revealing that a single minimum of the free energy surface (FES) is populated at 200 K. Starting from the finite temperature scenario, the potential energy surface (PES) was sampled by means of thermal annealing calculations showing that multiple local minima could be populated at 0 K. Both isotropic exchange coupling constants and zero field splitting (ZFS) parameters are calculated for the identified Fe\textsubscript{4}C\textsubscript{5}@Au(111) local minima, thereby allowing an in-depth characterization of the Fe\textsubscript{4}C\textsubscript{5} magnetic properties from the isolated to the grafted scenario. The cutting-edge computational protocol used here shows that the experimentally observed retention of a spin \(S = 5\) ground state upon grafting results from balancing the major modifications on spin Hamiltonian (SH) parameters. Electronic effects of the metal on Fe\textsubscript{4}C\textsubscript{5}’s magnetic properties have been also discussed.

1 Introduction

The possibility to employ just few atoms to observe magnetic bistability, and thus retain information, has been recently demonstrated,\textsuperscript{1,2} opening new frontiers in the high-density data storing with a potential increase of orders of magnitude with respect to capabilities available in nowadays magnetic data supports. In order to reach such great technological performances a system able to show an intrinsic magnetic hysteresis loop coming from its intramolecular properties is needed. In this context Single Molecule Magnets (SMMs) represent one of the best candidates to be used for magnetic data recording and in spintronic devices in general, though up to now only at cryogenic temperature.\textsuperscript{3,4} Their peculiar magnetic properties have been intensively studied as far as from the first observation of their molecular hysteresis loop\textsuperscript{5} and still, after almost twenty years, the interest on this class of molecular compounds has not diminished yet. However, in order to be integrated into real devices, SMMs must show enough robustness upon grafting or absorption process on surfaces.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{Fe4C5.png}
\caption{Sketch of Fe\textsubscript{4}C\textsubscript{5} molecule. Iron atoms are colored in blue, oxygen atoms in red, sulfur atoms in purple, carbon atoms in green and hydrogen atoms in white.}
\end{figure}

The SMM studied in this work, the propeller-shaped Fe\textsubscript{4}C\textsubscript{5}\textsuperscript{6} (see Fig. 1), showed to be robust enough to retain its magnetic hysteresis loop\textsuperscript{7} once grafted on Au(111). There-
Therefore, it is an appealing candidate for throughout analysis on how the magnetic properties change upon deposition on a surface. Fe$_4$C$_5$, in its crystalline environment, has an $S = 5$ ground state which originates from the leading antiferromagnetic coupling between the central high spin Fe$^{3+}$ ion (Fe$_c$ or Fe$_1$) and the three peripheral high spin Fe$^{3+}$ ions (Fe$_2$, Fe$_3$ and Fe$_4$). Although experimental data are usually interpreted assuming a threefold symmetry, the Fe$_4$C$_5$ has only a two-fold symmetry passing through the Fe$_1$-Fe$_2$ direction. However, in order to account for the effects on the magnetic properties caused by possible geometrical deviations from the crystalline symmetry, the Hamiltonian describing isotropic interactions is written as

$$H = J_{12}\hat{S}_1 \cdot \hat{S}_2 + J_{13}\hat{S}_1 \cdot \hat{S}_3 + J_{14}\hat{S}_1 \cdot \hat{S}_4 + J_2(\hat{S}_2 \cdot \hat{S}_3 + \hat{S}_2 \cdot \hat{S}_4 + \hat{S}_3 \cdot \hat{S}_4).$$  

(1)

Here, $J_{1x}$ ($x = 2-4$) are the first and $J_2$ the second neighbor exchange interactions ($J_2$ is left three-fold since it is usually below 1 cm$^{-1}$). The magnetic memory effect, i.e., the opening of magnetic hysteresis loop, is originated by the degeneracy lifting of the ground state multiplet due to the anisotropy term of the spin Hamiltonian,

$$H = \sum_i \hat{S}_i \cdot \vec{D}_i \cdot \hat{S}_i + \sum_{i,j \neq i} \vec{S}_i \cdot \vec{D}_{ij} \cdot \vec{S}_j,$$  

(2)

In the above expression, $i$ and $j$ run over the spin ($\hat{S}$) of the four iron ions. Alternatively, in the Giant Spin approximation (GSA), the same anisotropic physical behavior could be modeled as

$$H = \vec{S} \cdot \vec{D}_{\text{SMM}} \cdot \vec{S},$$  

(3)

where $\vec{S}$ now stands for the ground state giant spin $S = 5$ vector.

A detailed experimental characterization of Fe$_4$C$_5$@Au(111) can be achieved only through an interplay between different techniques. In particular, the difficult access to accurate structural information poses major limitations onto the full understanding of deposited SMMs properties. Indeed, the absence of an atomically resolved structure makes the assessments of the origin of the observed magnetic phenomena difficult to be unambiguously rationalized. In this scenario a cutting-edge theoretical description of SMMs deposited on surfaces becomes mandatory to fill up the missing information and therefore to draw a final picture of the SMM@Au(111) system. In this framework, we present here a complete and detailed DFT computational description of the structural and magnetic properties of Fe$_4$C$_5$@Au(111) employing both static and dynamic approaches. X-ray derived geometries are commonly used for the calculation of magnetic properties of SMMs. In some studies, to account for the effects of adsorption on surfaces, the structures have been optimized at the level of theory of choice. However, static optimization approaches are not adequate to describe a complex configurational space that could present multiple minima, as it is expected to be the case by deposition of SMMs on surfaces. On the other hand, ab initio molecular dynamics (AIMD) provides a proper sampling of the accessible space of configurations at finite temperature. The generated trajectory is expected to visit the accessible structures according to their statistical probability at the selected thermodynamic conditions. In this respect, the MD simulation samples the free energy surface (FES) underlying the dynamic of the system of interest, and reproduces its characteristic thermal fluctuations. Recently AIMD has been applied to the study of Spin crossover magnetic systems containing Fe$^{3+}$ ions. However, at the best of our knowledge, this computational strategy has never been applied before to study hybrid systems (adsorbate@surface) containing hundreds of heavy atoms and with a complex magnetic structure as the one observed for SMMs. Indeed, a similar statistical analysis restricted to only isotropic magnetic properties has been previously computed only at the QM/MM level for a metalloprotein system containing two iron ions bridged by di-sulfuric bridge as catalytic center. The AIMD application to the study of the grafting process of Fe$_4$C$_5$@Au(111) will be reported in the first part of this work where we obtained both dynamical information and steady state structures which could well statistically describe the molecular geometry once the SMM is grafted on Au(111). The so computed accurate atomistic picture of the system is of fundamental importance in order to understand how the deposition on a metallic surface affects the molecular magnetic properties. In the second part of the work, a comprehensive magnetic characterization of the computed geometries will be presented within the framework of previous successfully tested computational protocols based on GGA, GGA+U and hybrid functionals.

## 2 Computational Strategy and Methods

When a trend of physical properties must be evinced, it is of fundamental importance to handle all the elements of the ensemble with the same computational procedure. Such a care must be used in order to enforce the homogeneity of the results and reduce the source of errors coming from unbalanced computational treatments. Since the geometry of grafted species can not be determined at the experimental level (it can only be qualitatively extrapolated in simple cases, i.e., adsorbed porphyrins), the calculation of the relaxed geometry, through structure optimization, of the SMM on the surface becomes mandatory. Therefore, in order to study the evolution of the magnetic properties from the bulk to the grafted scenario, the same optimization procedure must be applied also to the SMM bulk phase. The magnetic data computed on optimized geometries become the reference values to compare to those computed in the grafted scenario. In order to have comparable magnetic properties, each spin Hamiltonian parameter must be computed with the same electronic structure approach before and after the grafting. The choice of the computational
protocol clearly depends on the properties we are interested in. GGA DFT calculations are known to accurately determine optimized structures of large systems. However, the accurate calculation of magnetic properties, e.g. the calculation of exchange coupling parameters, may require the use of hybrid functionals. The mixed computational protocol proposed in this work allows us to determine the evolution of the SMM’s magnetic properties from bulk to the grafted scenario. The reference bulk structure optimization and the AIMD simulations have been carried out with the TPSS functional corrected for the dispersion forces using the Grimme’s formalism. The calculation of exchange coupling parameters would require the use of hybrid functionals, while the anisotropic part of the spin Hamiltonian could be accurately calculated with the less demanding PBE functional. However, applying hybrid functionals to large periodic systems (SMM bulk and grafted on a surface) and over many configurations becomes computationally unaffordable. For this reason, the isotropic exchange coupling constants have been calculated at the PBE+U level of theory.

2.1 Models

The Au(111) surface has been modeled as a four layers slab of gold. Each layer consists of 80 gold atoms. The dimensions of the simulation cell are 23.05 x 25.0 x 60.0 Å. Periodic boundary conditions are always applied, but the size of the box is sufficient to avoid interactions between periodic images of the Fe₄S₅ units, which are about 10 Å apart from each other. From experimental and computational evidences it can be assessed that both thioacetyl and simple thiols undergo an homolithec cleavage of the S-Ac (S-H) bond with the formation of a sulfur radical, which is the species that effectively binds the metallic substrate. The Fe₄S₅ molecule has two side chains, each containing one thioacetyl group. Thanks to the steric hindrance of the magnetic core, only one of the two chains can actually bind to the substrate. According to literature, the initial configuration was prepared grafting the X-ray structure of Fe₄S₅ upon the Au(111) after the removal of one Ac group from one aliphatic chain, while leaving intact the other one.

2.2 AIMD and optimizations

AIMD calculations within the Born-Oppenheimer framework have been performed optimizing the wave function at each MD step. Electronic structure and nuclear forces have been calculated at the meta-GGA DFT level of theory, applying the Gaussian and plane wave (GPW) method as implemented in CP2K. The GPW approach is based on the expansion of the valence electron molecular orbitals in Gaussian type orbital basis sets, for which we use molecule optimized basis sets of the DZVP-MOLOPT-SR-GTH type. The auxiliary plane wave basis set is needed for the representation of the electronic density in the reciprocal space and the efficient solution of the Poisson’s equation. We truncate the plane wave basis set at 400 Ry. The interactions between valence electrons and atomic cores are described by means of Godecker-Teter-Hutter pseudopotentials. In particular, we used the TPSS functional together with the Grimme’s D3 corrections to account for the dispersion forces. The computational set up has been tested on bulk Au, Au(111) surface energy and cohesive energy of Benzene on Au(111) (available on ESI). The selected set up can adequately reproduce the structure of both gold slab and grafted molecule.

Hamiltonian equations of motion are numerically integrated using the velocity Verlet algorithm and a time step of 1 fs. Canonical distribution of momenta at 200 K is enforced with the canonical stochastic rescaled velocity (CSVR) thermostat with a time constant of 100 fs during thermalization and 500 fs during acquisition runs. Independent trajectories starting from the same initial configuration are generated by first initializing the velocities at 100 K and then annealing the system up to 200 K by different ramp procedures. Total energy conservation has been obtained with a smearing of molecular orbitals’ occupation numbers with a Fermi-Dirac distribution at 1500 K and with a convergence threshold criteria on the maximum wave function’s gradient of 1.0E-5.

When dealing with multispin systems, the DFT solution which can be used to evaluate forces is not unequivocally defined. Various spin solutions, described by broken symmetry (BS) states, are available. For what concern the overall structure of the complex and its interaction with the surface, the choices of one spin DFT solution instead of another is not important and no appreciable differences should be expected. For this reason, all the AIMD calculations are carried out with forces evaluated from the electronic configuration that better describes the lowest energy spin multiplet (S = 5): the central iron (Fe₄ or Fe₁) is down polarized while the peripheral irons (Fe₂ or Fe₂₋₄) are up polarized (schematically DUUU).

2.3 Magnetic Properties Calculations

Magnetic properties analysis have been carried out utilizing both CP2K and ORCA software. For what concerns the extrapolated geometries we used the procedure recently tested over a series of different Fe₄ SMMs. This procedure consists in two steps. The first one requires the CP2K code with the hybrid PBE0 functional for the calculation of the isotropic exchange coupling constants. The PW cutoff have been set to 400 Ry, DZVP-MOLOPT-SR basis sets with GTH pseudo potentials have been chosen for all the elements. For the evaluation of exact exchange integrals we take advantage of an auxiliary basis set: the DZVP-MOLOPT-SR basis set has been used for iron ions while the SZV-MOLOPT-SR basis set was chosen for carbon, hydrogen, oxygen and sulfur.
atoms. Convergence criteria over the maximum component of the wavefunction’s gradient were set to 1.0E-6. Following the Noodleman formulation of the BS approach\(^6\),\(^7\), isotropic exchange coupling constants have been extracted solving the linear system

\[
\Delta E(HS - BS(s_i \rangle \langle s_j)) = \sum_{ij} 2J_{ij\lambda} s_i s_j \lambda_{ij}^{BS}
\]

where \(s_i\) and \(s_j\) are the value of the single ion spins, \(J_{ij}\) their exchange interaction and \(\lambda_{ij}\) is equal to one if, for the specific BS state considered with the appropriate \(s_i\) value, the couple of spin \((i,j)\) is misaligned, zero otherwise. The SH considered to model the multiplet structure of the Fe\(_4\)C\(_5\) is given in Equation 1. In order to solve the system 4 for the SH model 1 we used four different energy differences built up by the HS state and the four BS determinants: DUUU, UDUU, UUDU and UUUD. Once the non relativistic multiplets structure is obtained, it is then possible to proceed with the second step, where the calculation of the anisotropy tensors is performed. For this second task the software ORCA has been utilized. The computationally cheap GGA (PBE) functional has been used, since the expensive PBE0 is found not to improve the results.\(^5\) We used the def2-TZVP basis set for all the elements and the RI approximation for the coulomb integral evaluation have been employed with the def2-TZVP/J auxiliary basis set. Speaking in the ORCA notation, Grid 5 and Very Tight convergence criteria were used. The protocol used to compute anisotropy splittings follows the route of the diamagnetic substitution. We evaluated single ion tensors making the substitution of all but one iron ions with Ga\(^{3+}\) ions at turn. Both Spin Orbit Coupling (SOC) and Spin Spin (SS) interactions were taken into accounts for single ion anisotropy tensor calculations.\(^38\),\(^39\) For what concerns the anisotropic exchange coupling interaction between iron ions we decided to take into account only spin dipole pair interaction. Moreover, although a few differences between DFT and point-dipole like calculation of this interaction exist,\(^40\) we decided to use the second one which is less computational demanding. This procedure makes possible to directly map the multispin Hamiltonian \(H=\Sigma \bar{s}_i \cdot D_i \cdot \bar{s}_i + \Sigma_{i\neq j} \bar{s}_i \cdot D_{ij} \cdot \bar{s}_j\) from DFT calculations. Thanks to the quite large \(J\) coupling constants, compared to anisotropy splitting, it was possible to take advantage of the strong exchange limit and use the Giant Spin Hamiltonian (GSH) \(H=\bar{S} \cdot D_{g,5} \cdot \bar{S}\), which is generally used to interpret experimental data. The multispin Hamiltonian and the GSH are related by the equation

\[
D_S = \sum_i d_i^S D_i + \sum_{ij} d_{ij}^S D_{ij}
\]

where the projection coefficients \(d_i^S\) and \(d_{ij}^S\) were evaluated

\[
d_i^S = \frac{\langle \alpha S \rangle |T_2(s_i, s_i)| \alpha S \rangle}{\langle \alpha S |T_2(S, S)| \alpha S \rangle}
\]

\[
d_{ij}^S = \frac{\langle \alpha S |T_2(s_i, s_j)| \alpha S \rangle}{\langle \alpha S |T_2(S, S)| \alpha S \rangle}
\]

where \(\langle \alpha S | \cdots | \alpha S \rangle\) stands for an irreducible matrix element of spherical tensor \(T_2\). Once the eigenket of the Heisenberg Hamiltonian (Eq. 1) \(|\alpha SM\rangle\) have been evaluated by numerical diagonalization, \(d_i^S\) and \(d_{ij}^S\) could be calculated taking advantage of the Wigner-Eckart theorem. It is well known that DFT at the GGA level overestimates the delocalization of the charge density. This issue leads to unacceptable overestimation of antiferromagnetic interactions. For this reason, in order to study the Fe\(_4\)C\(_5\) and the Au(111) substrate at the same time, we employed a GGA+U methodology\(^21\) as implemented in CP2K. This procedure makes possible to correct part of the GGA deficiency without loosing high speed performances of non-hybrids functionals. In this study we used the U correction for Fe, O and Au elements of 4.1 eV, 3.0 eV and 0.60 eV, respectively. These values were taken from the works of Nino-vana and Malavoliti et al.\(^36\),\(^37\). Spin moments for the iron ions have been calculated and reported on Table 5 of ESI for the different functionals used.

## 3 Results and Discussion

### 3.1 Bulk Structural and Magnetic Properties

One of the main issues related to the calculations of the magnetic properties at the quantum-mechanical level is the choice of the geometrical data. X-ray structures have often been considered as the best choice\(^41\), since optimization procedures might be not accurate enough. Slight distortions of the most important geometrical parameters have significant effects on the computed magnetic properties. Indeed, the geometry optimization of molecules in the gas phase can hardly provide structures corresponding to the experimentally determined magnetic properties, which are generally measured in the solid state. Hence, in order to get a reliable bulk reference state, it is crucial to quantify the effects on the computed magnetic properties ascribed to the optimization at the DFT level and to the crystalline environment. To this purpose, the Fe\(_4\)C\(_5\) molecule has been optimized both in vacuum (Opt-Isol model) and inside the periodic crystallographic cell (Opt-Bulk model). Optimizations have been carried out with the TPSS+D3 functional converging the electronic structures on the \(S = 5\) BS state (see the Methods section for more details). The resulting Root Mean Square Deviation (RMSD) of the the Opt-Bulk model with respect to the X-ray structure is 0.10 Å. The larger contribution to the RMSD value comes from the C\(_5\) aliphatic chains, which, however, are not well resolved in experiment. Instead, the RMSD of the iron and oxygen atoms of the magnetic core is only 0.03 Å. For the Opt-Isol model
the global RMSD amounts to 0.12 Å, while it is reduced to 0.04 Å considering only the core of the molecule. Considering the strong dependence of the magnetic properties on structural parameters, particular attention has been devoted to check the agreement between experimental and computed Fe\textsubscript{O}Fe\textsubscript{p} and γ-pitch angles of the propeller structure.

![Diagram](image)

**Fig. 2** (a): the γ-pitch angle; (b): the FeOFe angle. Color code as in Fig. 1.

The latter is defined as the dihedral angle between the irons’ plane and the plane Fe\textsubscript{O}Fe\textsubscript{p} defining the blades of the propeller structure, see Fig. 2a. Previous studies on magnetostuctural correlation\cite{15,42} evidenced that the isotropic exchange coupling between central and peripheral irons and the axial anisotropy intensity, strongly depends on the Fe\textsubscript{O}Fe\textsubscript{p} and the γ-pitch angles, respectively. The mean value of the Fe\textsubscript{O}Fe\textsubscript{p} angle for the Opt-Isol and the Opt-Bulk models deviates from the corresponding X-ray values of 1.5% and 0.9%, respectively. The average γ-pitch angle deviates of about 1.5% for both Opt models. To verify the effect of these deviations on the magnetic properties, the isotropic exchange coupling and the anisotropy tensors have been evaluated for the Opt-Bulk and Opt-Isol models (see Tables 1 and 2). We employ the same successful approach used to calculate these properties for the X-ray structures of a Fe\textsubscript{4} family, as reported in our previous paper\cite{15}. The isotropic exchange coupling constants are obtained at both PBE0 and PBE+U level of theory. The PBE+U results show a slight overestimation of the antiferromagnetic contributions comparing to the PBE0 ones(see Table 1). The difference between PBE+U and PBE0 can be ascribed to the fact that the used Hubbard U parameters for Fe and O were fitted at the DFT rev-PBE level.\cite{16} Nevertheless, our results show the good transferability of the U parameters within the Fe\textsubscript{4} class of SMMs.

The effective C\textsubscript{2} symmetry of the molecule is correctly retained only in the Opt-Bulk model, which turns out to be less antiferromagnetic than the corresponding X-ray structure (see Table 1). A further decrease of J has been obtained for the Opt-Isol model. The same trend has been observed for both the PBE0 and PBE+U approaches. This confirms the strong dependence of the magnetic properties on small geometrical variations, thus emphasizing the importance of accurate reference structures. Previous studies on iron dimers and Fe\textsubscript{4} molecules\cite{15,43} suggest that the FeOFe angle variations have the largest effect on J\textsubscript{1}. In fact, in the Opt-Bulk model a large decrease of J\textsubscript{1} seems to be induced by a slightly changed Fe\textsubscript{O}Fe\textsubscript{p} angle. Most of the structural differences between X-ray and Opt-Bulk come from the rotation of the two tripodal ligands along the pseudo C\textsubscript{3} symmetry axis. This rotation occurs without changing the position of the oxygen atoms. It rather modifies the out-of-plane angle of the alchoxy-nearest carbon with respect to the Fe\textsubscript{4}-O-Fe\textsubscript{p} plane. Test calculations over an iron dimer model support the hypothesis that also this degree of freedom could affect the isotropic exchange interaction between μ-alchoxy bridged iron clusters (see ESI for further explanations).

### Table 1 Fe\textsubscript{4}C\textsubscript{5} Crystal Magnetic Properties

<table>
<thead>
<tr>
<th>Functional</th>
<th>X-ray\textsuperscript{a}</th>
<th>Opt-Bulk\textsuperscript{a}</th>
<th>Opt-Isol\textsuperscript{a}</th>
<th>Exp</th>
</tr>
</thead>
<tbody>
<tr>
<td>J\textsubscript{12} (cm\textsuperscript{-1})</td>
<td>PBE</td>
<td>17.4</td>
<td>13.2</td>
<td>9.8</td>
</tr>
<tr>
<td>PBE+U</td>
<td>21.5</td>
<td>16.2</td>
<td>13.9</td>
<td>-</td>
</tr>
<tr>
<td>J\textsubscript{13} (cm\textsuperscript{-1})</td>
<td>PBE</td>
<td>15.3</td>
<td>8.1</td>
<td>5.0</td>
</tr>
<tr>
<td>PBE+U</td>
<td>19.2</td>
<td>11.7</td>
<td>8.8</td>
<td>-</td>
</tr>
<tr>
<td>J\textsubscript{14} (cm\textsuperscript{-1})</td>
<td>PBE</td>
<td>15.3</td>
<td>7.8</td>
<td>8.6</td>
</tr>
<tr>
<td>PBE+U</td>
<td>19.1</td>
<td>11.4</td>
<td>12.5</td>
<td>-</td>
</tr>
<tr>
<td>J\textsubscript{1} (cm\textsuperscript{-1}) \textsuperscript{b}</td>
<td>PBE</td>
<td>16.4</td>
<td>9.7</td>
<td>7.8</td>
</tr>
<tr>
<td>PBE+U</td>
<td>19.9</td>
<td>13.4</td>
<td>11.8</td>
<td>-</td>
</tr>
<tr>
<td>J\textsubscript{2} (cm\textsuperscript{-1})</td>
<td>PBE</td>
<td>0.2</td>
<td>0.1</td>
<td>0.1</td>
</tr>
<tr>
<td>PBE+U</td>
<td>0.4</td>
<td>0.2</td>
<td>0.2</td>
<td>-</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Selected molecule inside the Fe\textsubscript{4}C\textsubscript{5} crystal;
\textsuperscript{b} J\textsubscript{12}, J\textsubscript{13} and J\textsubscript{14} average value.

### Table 2 Fe\textsubscript{4}C\textsubscript{5} Crystal Magnetic Properties

<table>
<thead>
<tr>
<th>Functional</th>
<th>X-ray\textsuperscript{a}</th>
<th>Opt-Bulk\textsuperscript{a}</th>
<th>Opt-Isol\textsuperscript{a}</th>
<th>Exp</th>
</tr>
</thead>
<tbody>
<tr>
<td>D\textsubscript{1} (cm\textsuperscript{-1})</td>
<td>PBE</td>
<td>-1.05</td>
<td>-1.050</td>
<td>-1.046</td>
</tr>
<tr>
<td>E/D\textsubscript{1}</td>
<td>0.03</td>
<td>0.01</td>
<td>0.11</td>
<td>-</td>
</tr>
<tr>
<td>D\textsubscript{2} (cm\textsuperscript{-1})</td>
<td>PBE</td>
<td>0.818</td>
<td>0.639</td>
<td>0.650</td>
</tr>
<tr>
<td>E/D\textsubscript{2}</td>
<td>0.16</td>
<td>0.20</td>
<td>0.21</td>
<td>-</td>
</tr>
<tr>
<td>D\textsubscript{3} (cm\textsuperscript{-1})</td>
<td>PBE</td>
<td>0.731</td>
<td>0.619</td>
<td>0.540</td>
</tr>
<tr>
<td>E/D\textsubscript{3}</td>
<td>0.17</td>
<td>0.18</td>
<td>0.22</td>
<td>-</td>
</tr>
<tr>
<td>D\textsubscript{4} (cm\textsuperscript{-1})</td>
<td>PBE</td>
<td>0.731</td>
<td>0.628</td>
<td>0.591</td>
</tr>
<tr>
<td>E/D\textsubscript{4}</td>
<td>0.17</td>
<td>0.18</td>
<td>0.20</td>
<td>-</td>
</tr>
<tr>
<td>D\textsubscript{0,5} (cm\textsuperscript{-1})</td>
<td>PBE</td>
<td>-0.475</td>
<td>-0.426</td>
<td>-0.425</td>
</tr>
<tr>
<td>E/D\textsubscript{0,5}</td>
<td>0.02</td>
<td>0.03</td>
<td>0.06</td>
<td>0.01</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Selected molecule inside the Fe\textsubscript{4}C\textsubscript{5} crystal;

For what concerns the axial anisotropy parameter, D, the symmetry properties are correctly retained in the Opt-Bulk model and this seems to be sufficient to get a good agreement.
with experiment. Apparently, the structural changes generated by the optimization have a marginal effect on this quantity. These results shed light on the importance of studying the condensed phase to reproduce the experimental properties.

As a summary, the Opt-Bulk model is preferable to reproduce both geometrical and magnetic features of Fe₄C₅ and thus it is selected as reference model.

### 3.2 Fe₄C₅@Au(111) Molecular Dynamics and Structures Relaxation

The DFT-optimized structure published by Mannini et al.⁶ was obtained with the bare TPSS functional as were not available vdW correction methods at the time. This structure has been optimized with the computational set up explained in the Methods section but the inclusion of D3 corrections does not significantly change the Fe₄C₅@Au(111) structure. This geometry (GeoUp model, Fig. 3 left panel), already found by Mannini et al., is the starting configuration for the AIMD simulations. The initial thermalization run at 200 K (≈ 3 ps) has produced important structural changes and a stabilization of about 77 kcal mol⁻¹. The new relaxed geometry (GeoDown model, see Fig. 3 right panel) is characterized by the collapse of the aliphatic C₅ chain on the gold surface, so that the organic scaffold of the magnetic iron core is significantly closer to both the surface and the aliphatic chain itself. Moreover, the angle θ, defined as the angle between the normal to the plane of the four irons and the surface normal, is significantly smaller (see Fig. 3). θ is related to the easy-axis direction of the spin ground state anisotropy. As a consequence of this rearrangement, also the magnetic core undergoes a strong deformation, with the loss of the C₂ symmetry axis along the Fe₁-Fe₂ bond.

In order to sample the configurational landscape of Fe₄C₅@Au(111) at finite temperature, we started eight independent AIMD simulations from the minimum structure found after the thermalization. In order to generate independent tra-
Fig. 5 (a): Molecular average FeOFe angle’s normalized distribution. (b): Single FeOFe angle’s normalized distributions. (c): Molecular average γ-pitch angle’s normalized distribution. (d): Single γ-pitch angle’s normalized distributions.

The statistical distributions of Fe\textsubscript{c}OFe\textsubscript{p}, γ-pitch, and θ angles, reported in Fig. 5 and Fig. 6, provide an interesting overview on the observed distortions. The mean value of the Fe\textsubscript{c}OFe\textsubscript{p} angle, averaged over the three Fe\textsubscript{c}-Fe\textsubscript{p} couples inside the molecule, is 103.8° (Fig. 5a). This value is only slightly larger than the value characterizing the X-ray geometry: 102.6°. Splitting the average into the single angle contributions, both for the oxygen above (O\textsubscript{u}) and below (O\textsubscript{d}) the iron ions’ plane, as showed in Fig. 5b, it is noticed that the Fe\textsubscript{1}O\textsubscript{d}Fe\textsubscript{3} distribution is shifted with respect to the others. On the other hand, the single γ-pitch angles fluctuate around quite different mean values, while the molecular average is strongly peaked at 61.1° (Fig. 5c vs Fig. 5d). In both cases, the larger deviation from the molecular average value is observed for the Fe\textsubscript{3} ion’s properties. Such different behavior can be ascribed to the fact that this ion is the one in direct contact with the C\textsubscript{5} chain.

Even though Fe\textsubscript{c}C\textsubscript{5}@Au(111)’s local magnetic centers undergo severe geometrical distortions upon grafting, the differences are partially averaged at the molecular level. Fig. 6 shows that θ explores angles between 10° and 25° with a
Table 3 AIMD Walkers Final Energies

<table>
<thead>
<tr>
<th></th>
<th>GeoUp</th>
<th>GeoDown</th>
<th>Walker1</th>
<th>Walker2</th>
<th>Walker3</th>
<th>Walker4</th>
<th>Walker5</th>
<th>Walker6</th>
<th>Walker7</th>
<th>Walker8</th>
<th>Avg.</th>
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<tr>
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<td>-83.43</td>
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<td>-79.97</td>
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<tr>
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<td>0.00</td>
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<td>0.25</td>
<td>0.38</td>
<td>0.20</td>
<td>0.21</td>
<td>0.20</td>
<td>0.28</td>
<td>0.34</td>
<td>0.25</td>
</tr>
<tr>
<td>$\theta$</td>
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<td>102.94</td>
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<td>104.88</td>
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<td>104.07</td>
<td>103.73</td>
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<tr>
<td>$\gamma$-pitch</td>
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<td>14.7</td>
<td>18.5</td>
<td>18.0</td>
<td>16.5</td>
<td>16.9</td>
<td>16.3</td>
<td>13.9</td>
<td>14.3</td>
<td>17.5</td>
<td>16.5</td>
</tr>
</tbody>
</table>

$^a$ This averaged value is calculated over the eight final AIMD structure only;
$^b$ Evaluated considering irons and oxygens from magnetic core region only;
$^c$ Averaged over the molecule, Exp. Value 102.89$^\circ$;
$^d$ Averaged over the molecule, Exp. Value 70.45$^\circ$;
$^e$ Estimated upper limit value by Mannini et al. $^b$ 35$^\circ$;
$^f$ Reference value.

Fig. 6 The angle between the irons’ plane normal and the surface’s normal.

Fig. 7 Overlapped structure of all the Fe$_4$C$_5$ Walkers. Walkers’ geometries have been translated over the Au(111) slab minimizing the displacement between equivalent iron ions. Color code as in Fig. 3.

mean value of 16.7$^\circ$. This means that the magnetization easy-axis does not sample all the directions inside a cone, as expected.$^6$ Instead, it moves inside an hollow cone, avoiding the structures strictly perpendicular to the surface. The observed behavior is due to the presence of the C$_5$ carbon chain which remains under the organic scaffold of the Fe$_4$ core, thus imposing the tilt. The not innocent influence of the functionalization group of the tripod ligand on the final grafted geometry is even more evident. Indeed, if a flexible aliphatic chain can ensure a lower angle $\theta$, it can also induce larger geometrical perturbations at the magnetic core level, which in turn might alter the Fe$_4$C$_5$ magnetic properties. SMMs magnetism becomes experimentally evident only at temperature of a few K, while AIMD generates a structural sampling at 200 K.

In principle, starting from the sampled configurations, an estimate of the magnetic properties could be extrapolated from structural magneto-correlations. However, it is preferable to get rid of the entropic contributions, i.e. to derive from the FES exploration the corresponding equilibrium structures on the PES. Even though on the FES the system has remained in one single basin of attraction, we found that the underlying PES is topographically corrugated, i.e. is characterized by several local minima. The structural features distinguishing among these minima determine fluctuations of the magnetic properties that might become important for the understanding of the experimentally observed behaviour. In order to identify possible different minima on the PES, we applied simulated
3.3 Magnetic Properties Analysis

As discussed above, in order to employ a consistent electronic structure method to compare magnetic properties of Bulk-Opt model and the Fe$_4$C$_5$@Au(111) model, we adopt the PBE+U approach to evaluate isotropic exchange coupling constants. As for Bulk-Opt, Fe$_4$C$_5$@Au(111) retains antiferromagnetic exchange interaction between iron ions and the $S = 5$ ground state multiplicity. The average values of the $J$s reported in the last column of Table 4 reveal a stronger antiferromagnetic character for Fe$_4$C$_5$@Au(111) than for Opt-Bulk. In particular, $J_1$ increases of about 11 cm$^{-1}$ and $J_2$ of about 0.2 cm$^{-1}$. The bulk pseudo Cs$_5$ symmetry is here lost and the individual $J_{12}, J_{13}$ and $J_{14}$ values are quite different one from each other. According to the structural analysis made in the last section, these $J_{1x}$ differences could be attributed to the removal of any symmetry elements for the four iron ions upon grafting. The $J_1$ values are all larger than the bulk reference, as expected from the increase of the Fe$_x$OFe$_p$ angles. It is interesting to note that the standard deviations for the two exchange coupling parameters are 3.0 cm$^{-1}$ and 0.6 cm$^{-1}$, respectively. This means that the energy separation from the first excited state, $S = 4$, can range from 46 cm$^{-1}$ to 70 cm$^{-1}$. The estimate for the Opt-Bulk model is significantly lower, 32 cm$^{-1}$, once more strongly supporting the not innocent role of the Au(111) surface. These results pose a question on the nature of the effects induced by the grafting process. In order to

<table>
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<tr>
<th>Functional</th>
<th>Walker1</th>
<th>Walker2</th>
<th>Walker3</th>
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<th>Walker5</th>
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<th>Walker8</th>
<th>Avg</th>
</tr>
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<td>29.9</td>
<td>33.1</td>
<td>17.8</td>
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<tr>
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<td>20.2</td>
<td>23.4</td>
<td>22.8</td>
<td>23.3</td>
<td>27.7</td>
<td>30.7</td>
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<td>1.1</td>
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All reported values are in cm$^{-1}$.
* $J_{12}$, $J_{13}$ and $J_{14}$ average value.

<table>
<thead>
<tr>
<th>Functional</th>
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<th>Walker2</th>
<th>Walker3</th>
<th>Walker4</th>
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<td>$J_{1a}$</td>
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<td>21.3</td>
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<td>0.3</td>
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<td>0.2</td>
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</table>

All reported values are in cm$^{-1}$.
* $J_{12}$, $J_{13}$ and $J_{14}$ average value.

Table 4: Isotropic Magnetic Properties - Fe$_4$C$_5$@Au(111) Models

Table 5: Isotropic Magnetic Properties - Fe$_4$C$_5$@Au(111) Models

thermal annealing (from 200 K to a few K) to the final configuration of each of the eight generated trajectories. Each annealing has been followed by a structure optimization. Energies, RMSD and selected geometrical parameters for the eight optimized geometries (Walker1-8 from now on) are summarized in Table 3. All the structures are in a range of few kcal mol$^{-1}$ in energy. With respect to the GeoDown model, they are approximately 4 kcal mol$^{-1}$ more stable. RMSD (only for iron and oxygen atoms) values and Fe$_x$OFe$_p$ and θ angles are close to the averages of the corresponding distributions as obtained along the AIMD sampling, and differ substantially from those calculated for the GeoUp model. The most evident structural parameter that distinguishes the eight Walkers is the position of the dangling C@Au(111) aliphatic chain, as illustrated by the overlap of eight sets of coordinates in Fig. 7. Small, but significant from the magnetic point of view, deviations in the core region are also observed, which point to a distribution of the values of certain magnetic properties, even though these structures belong to the same minimum on the FES.
separately address steric effects and electronic contributions of the gold surface, we removed the gold slab and computed the magnetic properties molecule keeping the same coordinates, Fe₄C₅@Au(111) model. In this case the A Agencies group has been reintroduced at the S radical site, in order to avoid spurious spin contributions. Indeed, in the grafted scenario, at this site a strong Au-S bond is present. For a check on the reliability of the DFT+U correction for geometries different than the equilibrium one, the magnetic behavior of Fe₃ is modified. The dpm ligands of Fe₃ are always found to lie above the C₅ aliphatic chain. This causes the change in sign of its axial anisotropy parameter in all cases, except for Walker2, leading to an easy-axis anisotropy, in contrast with the easy plane anisotropy observed for the Opt-Bulk model. To be stressed that in spite of all these not negligible modifications, the orientation of the single ion tensors resemble the bulk one. The easy-axis of both the central iron and the peripheral iron ions are almost parallel to the iron plane normal, e.g. the mean inclination angle of the Fe₃ easy-axis with respect to the normal is 4.8° ± 3.7°. The global molecular anisotropy, evaluated for the S = 5 ground states in the GSH approximation, is an easy-axis anisotropy and, for all the adsorbed molecules except for Walker3, its value is diminished of about the 11%, with respect to the Bulk-Opt model. The easy-axis orientation, with respect to the iron’s plane normal, is retained: the easy-axis direction deviates from the peripherals iron’s plane normal of about 4°. Moreover, the easy-axis direction inclination (ζ) with respect to the surface normal, evaluated from the D₃₅ tensor, well matches with the angle θ for all the Walkers, except for Walker3. For Walker1, Walker3 and Walker7 we also observe the increase of the rhombicity term with respect to the Opt-Bulk model. The significant increase in the rhombicity anisotropy induced by the interaction of the aliphatic chain with the surface, reduc-

<table>
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<tr>
<th>Functional</th>
<th>Walker1</th>
<th>Walker2</th>
<th>Walker3</th>
<th>Walker4</th>
<th>Walker5</th>
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<th>Walker7</th>
<th>Walker8</th>
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<td>-1.105</td>
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<td>0.23</td>
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<td>0.17</td>
<td>0.14</td>
<td>0.23</td>
<td>0.32</td>
<td>0.19</td>
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<tr>
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<tr>
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<td>14.7°</td>
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<td>13.9°</td>
<td>15.3°</td>
</tr>
</tbody>
</table>

All reported D and E/D values are in cm⁻¹.
ing the deformation of the magnetic core.

4 Conclusions and Future Perspectives

In this work we present a computational study of the Fe₄C₅ SMM adsorbed on Au(111). We propose a new computational protocol, which is able to predict the evolution of both structural and magnetic properties going from the bulk to the adsorbed scenario. We emphasize the importance of the AIMD approach to properly take into account the distribution of accessible configurations and to reveal, after annealing, the roughness of the PES. Selected geometrical parameter, crucial for the SMM magnetism, have been monitored both at finite and zero temperature and considerable modifications, with respect to the bulk reference values, have been observed.

The relevant conclusion is that the retention of the S = 5 as ground state in the transition from bulk to Au(111) is not due to the rigidity of the Fe₄ core but on accidental balanced structural distortions of it. In particular the not innocent influence of the C₅ aliphatic carbon linker, on the final adsorbed geometry, have been discussed. Indeed, if a longer spacer chain can ensure a facilitate grafting procedure, it can also induce larger geometrical perturbations at the magnetic core level risking to alter the Fe₄C₅ magnetic structure. For this reason, a more rigid functionalization group, with less degrees of freedom, could reduce such a risk.

The combination of AIMD sampling, annealing procedures, and magnetic properties calculations at the DFT level of theory allowed to shed light on the effects of structural rearrangements and of the surface coupling on grafted SMMs. The fact that the electronic coupling with gold has a marginal role is in contrast to what observed for metal phthalocyanines and metal porphirins, where, instead, the metal is in direct contact with the surface and the metal’s magnetic orbitals could strongly hybridize with surface’s states. 44–46.

From a perspective point of view, this work paves the ground for further experimental and computational studies. Indeed, the confirmation of the importance of the aliphatic carbon spacer and the demonstration of the overcoming importance of structural rearrangements over electronic effects induced by the surface on magnetism show how could be important a chemical tailoring strategy, of the organic scaffold of SMMs, in order to modulate and control the SMMs’ magnetic properties on surface. In this context, further studies on different linkers and on effects of molecular packing on the surface become of primary importance.

5 Acknowledgement

We thank the European Research Council for funding through the Advanced Grant MolNanoMaS (no. 267746), Italian MIUR for support through FIRB project Nanomagnetismo molecolari su superfici metalliche e magnetiche per applicazioni nella spintronica molecolare (RBAP117RWN), and the CINECA award under the PRACE ("DySMoMAu" 5th PRACE call n. 2012061051) initiative, for the availability of high performance computing resources and support.

References

The adsorption process on gold and its consequences on the magnetic behavior of \{\text{Fe}_4\} SMM has been highlighted through AIMD.