First-principles study of an iron-based molecule grafted on graphene

A. Reserbat-Plantey, P. Gava, N. Bendib, and A. M. Saitta

1 Institut Néel, CNRS-UJF-INP - BP 166, 38042 Grenoble cedex 9, France, EU
2 Institut de Minéralogie et Physique des Milieux Condensés, CNRS-UMR 7590, Université Pierre et Marie Curie-Paris 6, Université Denis Diderot-Paris 7, IPGP - F-75252 Paris, France, EU

received 22 September 2011; accepted in final form 14 October 2011
published online 8 November 2011

PACS 73.22.Pr – Electronic structure of graphene
PACS 75.30.Hx – Magnetic impurity interactions
PACS 71.15.Mb – Density functional theory, local density approximation, gradient and other corrections

Abstract – Motivated by recent experimental studies on single molecular magnets grafted on graphene and single-walled carbon nanotubes, we investigate the structural, electronic, and magnetic properties of an iron-based magnetic molecule grafted on a graphene sheet using \textit{ab initio} calculations. In particular, the induced charge transfer and magnetization are described in terms of the coupling between the molecule and the graphene orbitals. This interaction and its effects on graphene electronic properties are determined and discussed in view of the potential utilization of graphene in spintronics.

The interaction between the graphene monolayer and a magnetic molecule is an important point in order to have insights into local effects such as induced magnetization and charge transfer, which are of interest for spintronic and nanoelectronic applications. In the present work, we chose the iron tetraphthalic acid molecule —FeTPA$_4$— deposited on monolayer graphene as a case-study of this interaction. Indeed, Gambardella et al. [20] have succeeded in the manipulation of the magnetic anisotropy of a supramolecular assembly of FeTPA$_4$ self-assembled on a Cu surface. However, to speed up our DFT calculations, we modeled FeTPA$_4$ by explicitly studying its magnetic core Fe(OH)$_4$, as we will discuss in the following, where this choice will be justified on the basis of structural, electronic, and magnetic properties.

Computational details. – First-principles DFT calculations were performed within the plane-wave approximation (PW) and pseudopotential scheme, as implemented in the quantum ESPRESSO code [21,22]. We adopt a Perdew-Burke-Ernzerhof (PBE) [23] gradient corrected functional for the exchange and correlation potential, and ultrasoft pseudopotential technique is used to describe C, H, O and Fe atoms. Electronic wave functions are written in terms of plane waves, with an

1 Spin-orbit coupling is not considered in this study.
A. Reserbat-Plantey et al.

Fig. 1: (Color online) Top and middle panels: structural description of a) Fe(TPA)$_4$ isolated molecule, b) Fe(OH)$_4$ isolated molecule, c) hybrid system composed of graphene and Fe(OH)$_4$. Iron and oxygen atoms are indicated in red, hydrogen atoms in blue, and carbon atoms in yellow. The high-symmetry adsorption sites are indicated with respect to the hexagonal cell of graphene d). Bottom panel: charge density map within the xy-plane for e) Fe(TPA)$_4$ and f) Fe(OH)$_4$.

energy up to 30 Ry, which is sufficient to ensure convergence of structural, electronic and magnetic properties. The electronic occupation is computed using Fermi-Dirac distribution, with a smearing parameter of 136 meV, which corresponds to an electronic temperature of 1578 K. The Brillouin zone integration is performed with a uniform k-points grid of (15 $\times$ 15 $\times$ 1). The convergence of the electronic and spin properties with the electronic temperature and k-points grid has been tested. Our choice of 1578 K with (15 $\times$ 15 $\times$ 1) k-points grid is within the range of typical electronic temperatures used to describe graphitic systems, and it is motivated by the fact that such values ensure converged results with reasonable computational effort. Structures are relaxed until forces are below 0.02 eV/Å. We used a supercell composed by (6 $\times$ 6) repetition of a graphene unitary cell in order to ensure a negligible interaction between periodic images of the adsorbed molecule. Spin-polarized calculations are also performed, and we consider polarization along the z-axis (i.e. perpendicular to the graphene plane).

Modeling FeTPA$_4$ through a smaller molecule. The FeTPA$_4$ molecule is difficult to simulate due to its large number of atoms (33 atoms) and, especially, of the size of the graphene supercell that should contain it. As mentioned above, however, the Fe(OH)$_4$ molecule mimics well the Fe(TPA)$_4$ molecule. This smaller, case-study system contains the same FeO$_4$ core as the bigger one, but the four large pyrenic arms are simply replaced by hydrogen atoms. As shown in the top panels of fig. 1, the two molecules have a similar planar structure. The charge distribution around the iron core is computed projecting all the electronic wave functions $\varphi_{i,\sigma}$ on the atomic orbitals centered in each given atom through the equation

$$n^\beta = \sum_{i,\ell,m,\sigma=\uparrow,\downarrow} \left| \left\langle \varphi_{i,\ell,m}^{\beta} | \varphi_{i,\sigma} \right\rangle \right|^2,$$

(1)

where $\varphi_{i,\ell,m}^{\beta}$ is the atomic-like orbital for atom $\beta$, labelled with its energy and angular momentum quantum numbers $m, \ell$.

We observe in table 1 that the charge associated to the core iron atom (Lowdin charge) remains very similar in both cases. The only slight charge difference between the two cases concerns the O atoms in Fe(OH)$_4$ having around 0.2 electrons more than in Fe(TPA)$_4$. We also computed the spin polarization along the z-direction for Fe and O atoms, as the difference between the spin-up and -down charges. As shown in table 1, the spin polarization of Fe atoms is very similar, being 3.20 and 3.10 $\mu_B$ in Fe(OH)$_4$ and Fe(TPA)$_4$ molecules, respectively. In both cases this polarization comes from unpaired electron in 3d iron orbitals. The polarization of the O atoms, which comes from 2p orbitals, is slightly higher in the Fe(OH)$_4$ molecule. The total polarization of the core is 3.94 and 3.59 $\mu_B$ in Fe(OH)$_4$ and Fe(TPA)$_4$, respectively, which

| Molecule | Atom | Lowdin charge | Polarization |
|----------|------|---------------|--------------|
| Fe(OH)$_4$ | Fe | 15.628 | 3.196 |
| | s | 2.503 |
| | p | 6.722 |
| | d | 6.402 |
| | O | 6.450 | 0.186 |
| | s | 1.686 |
| | p | 4.764 |
| Fe(TPA)$_4$ | Fe | 15.611 | 3.102 |
| | s | 2.473 |
| | p | 6.741 |
| | d | 6.395 |
| | O* | 6.246 | 0.122 |
| | s | 1.623 |
| | p | 4.623 |
corresponds to a variation of around 10%. We can thus safely consider these two molecules equivalent from the spin magnetic point of view in the void; given the relative inertness of graphene sheets with respect to full metallic surfaces, we can reasonably extrapolate a very similar behavior when grafted on a graphene sheet.

**Calculation details of the hybrid system.** We use, to study the hybrid system composed by the Fe(OH)$_4$ molecule adsorbed on graphene, an hexagonal supercell composed of 36 unit cells (fig. 1(c)) with a lattice parameter in the $xy$-plane of $a_0 = 14.76$ Å, and of 11.10 Å in the $z$-direction. Due to the Brillouin zone refolding, the Dirac ($\mathbf{K}$-) point$^2$ of graphene refolds on the $\Gamma$-point. The three high-symmetry adsorption sites of Fe(OH)$_4$ on graphene are shown in fig. 1(d). We find that the most energetically favorable adsorption site is the top site, and only this one will then be considered in the following.

**Results and discussion.**

**Structural properties.** We performed the structural relaxation of the hybrid system by setting a fixed distance $z_{\text{Fe-G}}$ between the Fe atom in the FeOH$_4$ molecule and the graphene plane. According to our calculations, the top adsorption site is found to be the most stable from an energetic point of view and we defined $z_{\text{Fe-G}}$ as the distance along $z$ between the Fe atom and the first C atom below, which we indicate as $C_{\text{top}}$. We calculated the binding energy as a function of this distance (see fig. 2), as $E_{\text{binding}} = E_{\text{graphene+FeOH}_4} - E_{\text{graphene}} - E_{\text{FeOH}_4}$. The obtained equilibrium distance for the system is $z_{\text{Fe-G}} = 4.1$ Å, and the corresponding binding energy is $E_{\text{binding}} = -0.36$ eV. However, it is well known that *ab initio* DFT calculations do not take into account Van der Waals interactions, which are of course important in weakly interacting systems. As a result, the distance we obtain is likely to be overestimated. A properly *ab initio* correction should explicitly take into account these contributions in the DFT functional. However, this procedure is rather cumbersome and computationally costly, and a common approximation consists in carrying out the calculations at the experimental equilibrium distance. The most flagrant case is graphite, a 3D crystal formed by layers of graphene, whose experimental interplane equilibrium distance is about 3.35 Å, while the DFT equilibrium value is more than 5 Å. In an analogous way, in the following we always compare results obtained for the DFT equilibrium distance $z_{\text{Fe-G}}^0 = 4.1$ Å with results obtained at a shorter distance, $z_{\text{Fe-G}} = 3.1$ Å which is comparable to the characteristic interaction distance observed in $sp^2$ carbon materials [24,25].

In both cases, the flatness of the graphene plane is slightly perturbed by the presence of the molecule. While C-C bond distances are practically unaffected ($\leq 0.1\%$), the $z$-coordinate of C atoms in the relaxed hybrid system changes by as much as 1% as shown in fig. 1. This surface modulation extends up to a distance of around 5.5 Å whether $z_{\text{Fe-G}}$ equals 4.1 or 3.1 Å.

**Electronic properties.** The charge transfer between graphene layer and the molecule can be quantified by computing the difference between the total charge of the graphene sheet in the hybrid system and of isolated graphene. We obtain a charge difference of $-0.63$ electrons, revealing an induced hole doping on graphene. This

---

$^2$Note that the $\mathbf{K}$-point in the first Brillouin zone corresponds to $k = 0$ since we are using a super-cell which is $6 \times 6$ graphene unit cell.

Fig. 2: (a) Binding energy of the hybrid system as a function of the distance along $z$ between the graphene surface and the iron atom of the Fe(OH)$_4$ molecule. (b) Displacements ($\Delta z_{C\#}$) of graphene C atoms positions along $z$ with respect to the isolated case, as a function of distance, within the $xy$-plane, to the adsorption (top site) $C_{\text{top}}$ atom. The horizontal arrow represents the planar extension of the Fe(OH)$_4$ molecule. Two fixed values of $z_{\text{Fe-G}}$ are considered: the equilibrium distance 4.1 Å (filled circles) and the characteristic distance 3.1 Å (empty triangles).
charge transfer is equal to a surface charge density of about \(-3.4 \times 10^{13} \text{ cm}^{-2}\), which is coherent with common predictions and experiments involving graphene [26–28].

We also evaluate the local electronic charge transfer for each C atoms in the graphene plane and extract contributions from the different orbitals by projecting the electronic eigenstates on the atomic wave functions as in eq. (1). At a molecule-graphene distance of 4.1 Å we observe that the charge transfer from the graphene sheet originates mostly (98%) from the 2$p$ orbitals of C atoms. A maximum charge transfer is observed for 2$p$ sheet originates mostly (98%) from the 2$p$ orbitals of C atoms. A maximum charge transfer is observed for 2$p$ orbitals of C atoms. A maximum charge transfer is observed for 2$p$ orbitals of C atoms.

For distances greater than 6 Å the spatial charge transfer states of C atoms at 2$p$ atoms. A maximum charge transfer is observed for 2$p$ atoms at 2.5 Å from the Fe center, while for distances greater than 6 Å the spatial charge transfer remains constant. For $z_{Fe-G} = 3.1$ Å, a similar behavior is observed. From this result we determine an interaction length of about 6 Å, which is in agreement with the one estimated by the analysis of the surface corrugation (see fig. 1). Moreover, the comparison between the electronic band structure of isolated graphene and of the hybrid system (not reported) indicates that, despite a shift of the Fermi level, the perturbation induced by the molecule is negligible, as also shown by the non-dispersive character of the molecular states.

This analysis of the charge transfer and of the electronic band structure of the hybrid system suggests that the quasi-metallic character of graphene is not perturbed by its interaction with the grafted magnetic molecule. This result is promising in view of the use of graphene as a sensitive detector.

Magnetic properties. As mentioned above, we also performed spin-polarized calculations on isolated molecule and on the hybrid system. The obtained magnetic moment for the isolated molecule Fe(OH)$_4$ is about 3.94 $\mu_B$, whereas in the hybrid system it is about 3.34 $\mu_B$. The magnetization difference between the two systems is about 0.6 $\mu_B$ indicating that the interaction with the graphene substrate generates 0.6 “paired” electrons in this hybrid system which is coherent with the estimated charge transfer of 0.63 electrons found in the previous section. We then analyze this variation of the molecular magnetization within the hybrid system spin states occupations. Thus, we calculated the projected density of states (PDOS) for spin-up and spin-down states within the hybrid system. More precisely, we focused on the projection into 2$p_z$ states of carbon atoms, which are the most sensitive to their environment.

In fig. 3(a) we report the difference between the PDOS for spin-up and -down states for different C atoms, labelled with their distance in the $xy$-plane with respect to Fe atom in the Fe(OH)$_4$ molecule. This figure shows an energy range $[-3.2; -3.1] \text{ eV}$ where the PDOS for spin-up and spin-down differs and, in particular, for carbon atoms located up to the 5th neighbour (i.e. 5 Å from $C_{top}$ atom). This spin-interaction distance is consistent with the previously discussed structural and electronic interaction lengths of the hybrid system.

![Fig. 3: (Color online) (a) Difference between PDOS (see text) for spin-up and -down states within the hybrid system. Projections are performed onto carbon 2$p_z$ atomic orbitals. Carbon atoms are labelled by their neighbouring rank, in the $xy$-plane, with respect to $C_{top}$. The sketch represents the C atoms of the graphene. Fe(OH)$_4$ is located at 3.1 Å vertically above the black $C_{top}$ atom and the different colors indicates the different neighboring rank. PDOS$_{2p_z}$↑ − PDOS$_{2p_z}$↓ is averaged for every C atom at the same distance from $C_{top}$. (b) Spatial distribution of the magnetization $\Delta n(\vec{r}) = n(\vec{r})_↑ − n(\vec{r})_↓$ within hybrid system for two distances $z_{Fe-G} = 4.1$ Å (top) and $z_{Fe-G} = 3.1$ Å (bottom).]

To go further, we analyze the mixing between atomic orbitals of the Fe(OH)$_4$ molecule and the ones of graphene in the hybrid system. In order to quantify the hybridization level of the occupied states, we define the quantity $c_{g/M}$:

$$c_{g/M} = \sum_{\text{AO}} c_{g,M}^{\text{AO}} ,$$

where $c_{g,M}^{\text{AO}}$ represents the projection of an electronic states into atomic orbitals (AO) belonging to graphene atoms (g) or to the molecule atoms (M). These states in the range $[-3.2; -2.0] \text{ eV}$, with respect to the Fermi level, correspond to occupied electronic states where $c_{g,M} \in [10\%; 90\%]$. Within that interval, both systems (i.e. graphene and molecule) have a significant contribution to the global electronic wave functions which suggest a remarkable hybridization. This is consistent with findings shown in fig. 3(a), where we observe strongly polarized states in a region within $[-3.0; -1.5] \text{ eV}$ and also with the existence of additional mixed states. In particular, at $k = 0$ and at $-2.4 \text{ eV}$ there is an electronic state which is particularly important since it represents the overlap of Fe 3$d_{yz}$ (13%) and top C atom 2$p_z$ orbital.
Moreover, the contribution of this atomic orbital from the closest carbon atom is not present in any other mixed wave function. This indicates, for instance, the negligible interaction between the C atom and the oxygen atoms. All other Fe states are only present in pure molecular electronic states and not in mixed states. Conversely, we do not observe strongly hybridized spin-down states. The hybridized states in the range [−3.2; −2.0] eV show mainly overlap between oxygen 2p_{x,y} states and 2p_{z} states of carbon atoms located at distances $\geq 5.5 \, \text{Å}$ from the Fe atom. On the basis of our findings, we expect a spin-up charge excess around the carbon atoms closer to the Fe atom and an excess of spin-down charges within a crown of carbon atoms around 5.5 Å from the Fe atom. This local magnetization within the graphene plane is shown in fig. 3(b). The induced magnetization in the graphene plane is suggested by concentric crowns of spin-up/-down electrons in 2p states of the carbon atoms.

**Conclusions.** We report a case-study of an iron-based magnetic molecule grafted on a graphene sheet and, through ab initio DFT calculations, we predict an induced magnetization effect to the graphene substrate. We also describe a typical interaction length of 5–6 Å within the graphene plane with respect to structural deformations, localized charge transfer and induced magnetization. The latter originates from a significant coupling between 2p_{z} and Fe 3d_{z} orbital for the closest carbon atom. On the other hand, we observe that the presence of an iron-based magnetic molecule in the close vicinity of a graphene sheet does not strongly perturb the electronic properties of the latter, thus providing a promising perspective to the design of non-destructive sensor devices.

***

We acknowledge useful discussions with X. Blase, V. Bouchiat, A. Candini, M. Urdampilleta and W. Wernsdorfer. Figures 1 and 3 have been performed with XCRYSDEN package [29]. Calculations were done at IDRIS (Orsay, France), Project No. CP9-91387.

REFERENCES

[1] Schedin F., Geim A. K., Morozov S. V., Hill E. W., Blake P. et al., Nat. Mater., 6 (2007) 652.
[2] Neto A. C., Guinea F., Peres N., Novoselov K. and Geim A., Rev. mod. Phys., 81 (2009) 109.
[3] Tombrinos N., Joza C., Popinciuc M., Jonkman H. and van Wees B., Nature, 448 (2007).
[4] Candini A., Klyatskaya S., Ruben M., Wernsdorfer W. and Affronte M., Nano Lett., 11 (2011) 2634.
[5] Bunch J., van der Zand A., Verbridge S., Frank I., Tanenbaum D. et al., Science, 315 (2007) 490.
[6] Eichler A. et al., Nat. Nanotechnol., 6 (2011) 339.
[7] Zhang H. et al., Nanoeletters, 11 (2011) 4047, DOI: 10.1021/nl200803q.
[8] Kessler, GRIT, ZETTL and BOUCHIAT, Phys. Rev. Lett., 104 (2010) 047001.
[9] Heersche H., Jarillo-Herrero P., Oostinga J., Vandersypen L. and Morpurgo A., Nature, 446 (2006) 5659.
[10] Krasheninnikov A. V., Lehtinen P. O., Foster A. S., Pyynkäl P. and Nieminen R. M., Phys. Rev. Lett., 102 (2009) 126807.
[11] Ling Xi, Xie Liming, Fang Yuan, Xu Hua, Zhang Haoli et al., Nanoeletters, 10 (2010) 553.
[12] Lopes Manuel, Candini Andrea, Urdampilleta Matias, Reserbat-Plantey Antoine, Bellini Valerio et al., ACS Nano, 4 (2010) 7531.
[13] Urdampilleta M. et al., Nat. Mater., 10 (2011) 502.
[14] Wernsdorfer W., C. R Chimi., 11 (2008) 1086.
[15] Xiao Ruijuan, Frisch Daniel, Kuzmin Michael D., Koepernik Klaus, Eschrig Helmut et al., Phys. Rev. Lett., 103 (2009) 187201.
[16] Leenaerts O., Partoens B. and Peeters F. M., Phys. Rev. B., 77 (2008) 125416.
[17] Johll H., Kang H. C. and Tok E. S., Phys. Rev. B, 79 (2009) 245416.
[18] Mao Y., Yang J. and Zhong J., J. Phys.: Condens. Matter, 20 (2009) 115209.
[19] Lehtinen P. O. et al., Phys. Rev. Lett., 91 (2003) 017202.
[20] Gambaradella Pietro, Štepanov Sebastian, Dmitriev Alexandre, Honolka Jan, de Groot Frank M. F., Lingenfelder Magal et al., Nat. Mater., 8 (2009) 189.
[21] Baroni S., Dal Corso A., de Gironcoli S., Giannozzi P., Cavazzoni C. et al., Quantum espresso: open-source package for research in electronic structure, simulation, and optimization, www.quantum-espresso.org (2005).
[22] Baroni S., Dal Corso A., de Gironcoli S. and Giannozzi P., Pwscf and phonon: plane-wave pseudopotential codes, http://www.pwscf.org/ (2005).
[23] Peredew J. P., Burke K. and Ernzerhof M., Phys. Rev. Lett., 77 (1996) 3865.
[24] Cardona M. and Merlin R., Light Scattering in Solids: Novel Materials and Techniques (Springer) 2007.
[25] Saito R., Dresselhaus G. and Dresselhaus M. S., Physical Properties of Carbon Nanotubes (Imperial College Press) 1998.
[26] Novoselov K. S. et al., Science, 306 (2004) 666.
[27] Novoselov K. S. et al., Nature (London), 438 (2005) 197.
[28] Ferrari A. C. et al., Phys. Rev. Lett., 97 (2006) 187401.
[29] Koval A., Comput. Mater. Sci., 28 (2003) 155. Code available from http://www.xcrysden.org/.