Gate-tunable magnetism of C adatoms on graphene

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We have performed density functional theory calculations of graphene decorated with carbon adatoms, which bind at the bridge site of a C–C bond. Earlier studies have shown that the C adatoms have magnetic moments and have suggested the possibility of ferromagnetism with high Curie temperature. Here we propose to use a gate voltage to fine tune the magnetic moments from zero to 1 µB while changing the magnetic coupling from antiferromagnetism to ferromagnetism and again to antiferromagnetism. These results are rationalized within the Stoner and RKKY models. When the SCAN meta-GGA correction is used, the magnetic moments for zero gate voltage are reduced and the Stoner band ferromagnetism is slightly weakened in the ferromagnetic region.

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I. INTRODUCTION

A functionalization of semiconductor devices with ferromagnetic properties is one of the greatest challenges of modern spintronics.1–3 Much research is nowadays focused on incorporating magnetic properties into semiconductor system and, in particular, into graphene.4–18 Graphene modified by defects is very promising for such purpose. Because of the reduced coordination number in the two-dimensional system, the defect states are naturally expected to have weak coupling to graphene, thus broadening of their energy levels is expected to be small and with high density of states (DOS). If the Fermi level (EF) further falls inside the broadened energy level, the Stoner instability19 removes the spin degeneracy of the impurity states, leading to the onset of magnetization. Thus, graphene can magnetize even first row element adatoms20 producing two-dimensional d8 semiconductor magnetism. This kind of scenario becomes particularly interesting for external gate voltage (Vg) control of the Fermi energy. In fact, the impurity state can be easily occupied/drained using Vg.21–23 Consequently, a rich phenomenology emerges from the interplay between magnetic impurities and the gate-voltage control of the interactions. As we shall see in the next sections, the description of this interplay is rather complex and it has not been carefully examined in earlier studies. Experimentally gate-voltage control has been used for producing magnetism on graphene oxide,11 N-doped graphene oxide16 and graphene grafted with Pt-porphyrins,14 revealing the existence of ferromagnetic phases with significant magnetic moments for particular values of Vg. The magnetic impurities develop a narrow band,19,24 which has been proposed to give rise to Stoner band ferromagnetism with high Curie temperatures even for sp electron systems.25 Moreover, the indirect exchange interaction of impurities on graphene has been intensively studied within the Ruderman-Kittel-Kasuya-Yosida (RKKY) framework26–32 and the recently developed extension of this approach, which takes into account a resonant hybridization between the adatoms and graphene.33

Most of the literature considers impurities interacting with one of the two sublattices A or B of graphene. The Lieb’s theorem34 for bipartite lattice applies for these cases. As a corollary of this theorem, two impurities connected to the same sublattice interact ferromagnetically, while the interaction is antiferromagnetic for impurities connected to different sublattices. This sublattice-dependence has been verified by the various computational and experimental studies.13,18,35–37 For adatoms, the situation depicted above corresponds to the top-site binding. However, the bridge site is in general energetically more favorable than the top site,20,38 despite the fact that it has not been studied as intensively. Interestingly, the magnetic interactions for bridge site adatoms is different since coupling with both sublattices occurs, as pointed out by Gerber et al. for carbon adatoms.39 In this situation, the Lieb’s theorem does not apply anymore because the lattice is no more bipartite. These conditions are therefore more favorable for ferromagnetism within the C adatom network.

The carbon adatom39–46 can be considered as an exemplar case for the bridge-site binding. It has only one partially filled magnetic orbital at about 0.3 eV below graphene Dirac point (ED). Here, we name this orbital as ψp⊥ since it has p symmetry with its symmetry axis perpendicular to the C–C bond of the bridge site, as shown in Fig. 1. The ψp⊥ orbital hybridizes only weakly with the graphene backbone, thus it has high partial DOS and it preserves the Dirac cone shape. Therefore, the ideal model considering a localized state and a Dirac cone remains valid. Taking all these considerations into account, one can conclude that C adatoms are interesting for both conceptual and applied purposes. Nevertheless, we must bear in mind that the controlled production of C adatom networks on graphene remains a major challenge. To address these difficulties, a recent paper by Kim et al.47 has suggested a facile pathway for the realization of this system.47 In this study the C adatoms have been monitored using a powerful Raman technique.48 Another ob-
FIG. 1. (Color online). Magnetization density isosurface plot for a C adatom on graphene. The isosurface is at 0.004 e/Å³, while the corresponding negative isosurface is not visible. The $\psi_p,\perp$ state (with $p_x/p_y$-symmetry) is visible at the adatom. Its $p_z$ character on the graphene sheet can be seen as well. Left inset: another view of the geometry. Right inset: schematic electronic structure, the occupied portion of the graphene/$\psi_p,\perp$ states are in grey/red, respectively.

obstacle resides in the stability of the adatoms, which are mobile even at room temperature. The C adatoms stick in one place only for few seconds due to the migration barrier ranging from 0.35 eV to 0.60 eV. However, the intrinsic magnetism reported in graphite suggests that C adatoms can be stabilized at least under some conditions since it is highly plausible that part of this magnetism originates from C adatoms. Also for instance curvature of the graphene sheet has been reported to increase the migration barrier.

The present paper provides a comprehensive theoretical study of graphene decorated with C adatoms and shows that the magnetic properties can be controlled with a gate voltage. The paper is organized as follows: Sec. II contains the methodology and part A discusses the used computational supercells, part B gives the other computational details and part C discusses the theoretical framework. Sec. III reports the results of the study. Part A focuses on effects of bias to a single adatom while part B presents the most important results of this study, which is the behavior of the magnetic interactions as a function of gate voltage for remote interaction distances and its interpretation. Part C contains results for close interaction distances. Part D illustrates the spatial spin polarization patterns induced in graphene by the $\psi_p,\perp$ state. Sec. IV contains the conclusions and outlook of the present work.

II. METHODOLOGY

A. Studied configurations

The magnetic interactions between C adatoms were studied by placing two adatoms $C_{1}^{ad}$ and $C_{2}^{ad}$ on a graphene supercell (SC) and by computing the energies of the parallel and antiparallel spin configurations, which we also refer to ferromagnetic (FM) and antiferromagnetic (AFM) solutions throughout the present paper. The energy difference $\Delta E = E_{FM} - E_{AFM}$ contains the information about magnetic interactions. A comprehensive set of adatom orientations on different graphene supercells (SCs) were considered for both one-sided and two-sided adsorptions, where the adatoms are adsorbed on the same side and opposite sides of the graphene sheet, respectively. Two configuration types were used, as explained by the following paragraphs.

In adatom pair configurations, the adatoms were placed close to each other. These configurations were used to model short range pairwise interactions as in the study by Gerber et al. The results for these pairs are discussed in Sec. III C. Eight different orientations of the adatoms were considered, labeled #1 to #8, and they are illustrated in Fig. 2. These configurations contain all the pairs that are separated at most 5.8 Å (Conf. #8) and are...
stable for one-sided adsorption. For ranges shorter than \( \sim 3.8 \text{Å} \) (Conf. #1) the adatoms dimerize as shown by our simulations with relaxation. The pair configurations were studied in two supercell sizes. The first is the \( 7 \times 7 \) SC containing 98 graphene atoms plus the two adatoms. In this case, the separations between periodic images are at least 11.7 Å (Conf. #8), therefore the pairs are reasonably well isolated. The SC size \( 7 \times 7 \) is also used by Gerber et al., and their Conf. C, D and E correspond to our Conf. #3, #2 and #6, respectively. Another size is \( 6 \times 3 \), these SCs have 36 graphene atoms plus the two adatoms. The interactions between adatoms and their periodic images are strong. Thus the adatom pairs interact with their surroundings. These cells were intended for simulating random adatom distributions with a coverage of one adatom per 18 graphene atoms.

In adatom array configurations, the adatoms and their periodic images form regular and infinite arrays, which were employed in simulations involving long-range interactions. These arrays replace the isolated adatom pairs, which would have prohibitively large SCs in the case of remote interactions. The adatom array results are discussed in Sec. III B. The studied array types were labeled \( \alpha, \beta \) and \( \gamma \). For \( \alpha \) arrays a \( N \times N \) graphene SC with one adatom was repeated twice to yield two adatoms in a \( 2N \times N \) SC. This operation leads to equilateral triangular array as illustrated in Fig. 2 (a) for the case of \( 8 \times 4 \) SC. In this study, two SC sizes (\( 6 \times 3 \) and \( 8 \times 4 \)) were used and they correspond to \( \sim 7.5 \text{Å} \) and \( \sim 10 \text{Å} \) adatom separations, respectively. For the \( \beta \) arrays, \( C_3^d \) was moved to the neighboring bridge site, breaking the \( D_{3h} \) symmetry of the \( \alpha \) arrays. The \( \gamma \) arrays were obtained by further moving \( C_3^d \). In these arrays six nearest-neighbour adatom-atom interactions per unit cell are present. However, in interactions between an adatom and its own periodic images the spins are parallel in both FM and AFM solutions, hence their contribution to \( \Delta E \) cancels. There are two interactions per unit cell leading to this cancellation (blue dashed lines in Fig. 2 (a)), hence only four nearest-neighbor interactions (red dashed lines in Fig. 2 (a)) contribute to \( \Delta E \).

### B. Computational details

The calculations were performed within the density functional theory (DFT) as implemented in the Vienna ab initio simulation package (VASP). DFT is exact apart from the exchange-correlation (XC) functional taking into account the many-body quantum effects. To describe the XC functional, we used the generalized gradient approximation (GGA) in the form proposed by Perdew, Burke and Ernzerhof (PBE). GGA is a correction to the old local density approximation, which in iron improves the stability of FM phase. Moreover, a set of calculations were repeated with the strongly constrained and appropriately normed (SCAN) meta-GGA functional, which is a more precise XC scheme obeying the 17 known exact constraints. SCAN typically produces superior results compared to most GGA functionals. The Kohn-Sham orbitals were expanded in a plane wave basis set with an energy cutoff of 600 eV. The electron-ion interactions were taken into account using the projector augmented wave (PAW) method and the electronic energy minimization was performed with a tolerance of \( 10^{-5} \) eV. Each structure was optimized until the residual forces became smaller than 0.01 eV/Å. In these relaxation runs we used first order Methfessel-Paxton smearing with width of 0.1 eV. The Brillouin zone was sampled with dense \( \Gamma \)-centered meshes with \( k \)-point separations lower than \( 0.017 \cdot 2\pi/\text{Å} \). For example, a \( 4 \times 4 \times 1 \) mesh was used for \( 7 \times 7 \) SCs and a \( 4 \times 8 \times 1 \) mesh was used for \( 8 \times 4 \) SCs. We found that the high \( k \)-point density is essential in describing the magnetic \( \psi_{p,\uparrow} \) state of the adatom correctly. Some of these parameters were refined for the SCAN simulations (see Sec. 1.1 of the supplemental material). To obtain more accurate total energies and DOS, the Methfessel-Paxton method was upgraded to the tetrahedron method with Blöchl corrections. Moreover, the \( k \)-point grids were increased to \( 10 \times 10 \times 1 \) mesh for \( N \times N \) SCs and \( 8 \times 16 \times 1 \) mesh for \( 2N \times N \) SCs. As in earlier work, the gate voltage was modeled by adding \( \Delta Q/e \) electrons. In order to avoid divergence in the Ewald summation, the unit cell was kept charge neutral by adding a compensating helium background charge. These computational schemes may produce errors as discussed in detail in Sec. 1.2 of the supplemental material. Moreover, when \( \Delta Q \gtrsim 1.5 \text{e} \) and when the SCs are small, some charge spilling to the vacuum occurs. This problem was overcome by decreasing the the unit cell height to \( 15 \text{Å} \) or even \( 13 \text{Å} \) from the standard value of \( 20 \text{Å} \). The aforementioned errors are increased by the decreasing of the unit cell height, but our results are not significantly affected by all these errors as explained in the Sec. 1.2 of the supplemental material.

### C. Theoretical analysis

In conventional RKKY theory, the indirect exchange interaction between two adatoms mediated by delocalized electrons is described by the following Hamiltonian:

\[
\hat{H}_{\text{RKKY}} = J \delta(\mathbf{r} - \mathbf{R}_1) \hat{\mathbf{S}}_1 + J \delta(\mathbf{r} - \mathbf{R}_2) \hat{\mathbf{S}}_2,
\]

where \( \hat{\mathbf{S}} \) is the electron spin operator, \( \mathbf{I}_{1,2} \) are the adatoms spins, \( J \) is the exchange constant, and \( \mathbf{R}_{1,2} \) denote positions of the two adatoms. The energy of the system is obtained by treating Eq. (1) as a perturbation, the second order correction depends on the configuration of the adatoms spins. This interaction oscillates as a function of the distance between the magnetic centers, in the case of graphene its sign also depends on the positions of the adatoms in relation to the graphene sublattices \( A \) and \( B \). For small adatoms separations the alignment is FM for \( AA \) and AFM for \( AB \) configuration. This well-known result is a consequence of Lieb’s theorem for bipartite
lattice.\textsuperscript{34} A completely different situation occurs if the bound adatom states at energy $\varepsilon_0$ are resonantly coupled to the graphene 2D continuum, i.e., $\varepsilon_0$ falls in the range of occupied states of graphene ($\varepsilon_0 \ll E_F$).\textsuperscript{33} In this case an effective resonant hybridization occurs between the bound states and the graphene states lying within a small energy range near $\varepsilon_0$. The resonant type of hybridization makes the perturbation approach used by conventional RKKY theory inapplicable. However, the problem can be solved using a different approach.\textsuperscript{33,73,74} This generalized theory of indirect exchange considers the following Hamiltonian:

$$\hat{H} = \hat{H}_0 + \hat{H}_T + \hat{H}_{\text{RKKY}},$$

(2)

where $\hat{H}_0$ describes the non-interacting adatoms and graphene, $\hat{H}_T$ describes the coupling of the magnetic centers to the graphene, in particular it incorporates the details of the coupling to the sublattices $A$ and $B$, $\hat{H}_{\text{RKKY}}$ is the exchange term of Eq. (1). The general expression for the indirect exchange energy reads:\textsuperscript{33}

$$E_{\text{RKKY}}^{\text{ex}} = \frac{E_F}{\pi} \int_{-\infty}^{E_F} \frac{dE}{\arctan j^2 E^2} g(E, \mathbf{R}) \text{sign} E \left(\varepsilon_0 - E\right)^2 - j^2.\right.$$  

(3)

Here $j$ is the exchange energy constant describing direct exchange interaction between the adatom and graphene. The function $g(E, \mathbf{R})$ encapsulates all the details of the coupling of the two interacting adatoms and graphene and $\mathbf{R}$ connects the adatoms. In the case of the resonant coupling ($\varepsilon_0 \ll E_F$), only the energy range close to $\varepsilon_0$ contributes to the interaction due to the poles of the integrand in Eq. (3). In other words, the resonant indirect exchange ($E_{\text{RKKY}}^{\text{ex}}$) is effectively mediated only by the electrons with approximately resonant energy $\varepsilon_0$ and the adsorption geometry does not play any role. Consequently, the sign of $E_{\text{RKKY}}^{\text{ex}}$ only depends on the position of the resonant state relative to the Dirac point:

$$\text{sign} E_{\text{RKKY}}^{\text{ex}} = -\text{sign} \left(\varepsilon_0 - E_D\right).$$  

(4)

If the bound impurity state lies above $E_D$, the indirect exchange is mediated by electron-like states of graphene and the indirect exchange is FM at a small distance between the adatoms, whereas if $\varepsilon_0 < E_D$, it is AFM at a small distance as it is mediated by graphene hole-like states.\textsuperscript{33} In the case of carbon adatoms, the $\psi_{p, \perp}$ state is located below $E_D$, therefore, we expect the C adatoms to interact antiferromagnetically as when $E_P$ is tuned above $\varepsilon_0$ by gate voltage, which we refer to as resonant region.

The opposite case ($\varepsilon_0 \gg E_F$) is the familiar conventional RKKY. Mathematically the difference stems from the poles being outside the integration limits in Eq. (3), thus the whole range of the occupied states contributes to the integral. Eq. (3) simplifies for the $AA$, $AB$ and bridge-bridge configurations into

$$E_{\text{RKKY}}^{\text{ex,AA}} (\mathbf{R}) = -\frac{\tau j^2}{8\pi \varepsilon_0} \left(\frac{\hbar v_F}{R}\right)^3 \cos^2 \left(\frac{x(\mathbf{R})}{2}\right),$$

$$E_{\text{RKKY}}^{\text{ex,AB}} (\mathbf{R}) = \frac{\tau j^2}{8\pi \varepsilon_0} \left(\frac{\hbar v_F}{R}\right)^3 \cos^2 \left(\frac{x(\mathbf{R})}{2} - \theta_R\right),$$

$$E_{\text{RKKY}}^{\text{ex,AA}} (\mathbf{R}_{AB}) \approx E_{\text{RKKY}}^{\text{ex,AA}} (\mathbf{R}_{AA}) + E_{\text{RKKY}}^{\text{ex,AA}} (\mathbf{R}_{BB}) + E_{\text{RKKY}}^{\text{ex,AB}} (\mathbf{R}_{AB}) + E_{\text{RKKY}}^{\text{ex,AB}} (\mathbf{R}_{BA}),$$

(5)

where $R = ||\mathbf{R}||$, $\theta_R$ is the polar angle of $\mathbf{R}$, $x(\mathbf{R}) = (\mathbf{K} - \mathbf{K}') \cdot \mathbf{R}$, $\mathbf{K}$ and $\mathbf{K}'$ are the graphene Dirac points, $\tau$ is the energy parameter defining the strength of the adatoms coupling to graphene and $\mathbf{R}_{AB}$ is the vector connecting the A sublattice basal site of the first adatom to the B sublattice basal site of the second adatom. Eqs. (5) are in agreement with the conventional RKKY theory results for graphene.\textsuperscript{26–32} The difference in the pre-factor is due to the details of the model where the bound state level exists also in a non-resonant case.\textsuperscript{33} As a consequence of Eqs. (5), the $AA$ interaction is FM while $AB$ is AFM. However, the sign of $E_{\text{RKKY}}^{\text{ex,AA}}$ depends on the relative position of the adatoms. It can be approximated by summing the four $AA/AB$ type interaction pairs between the two basal sites of the first adatom and the two basal sites of the second adatom, as in Ref. \textsuperscript{30} for selected configurations. As the amplitude of the AFM terms is three times larger than the FM contributions, AFM configurations are more frequent. For example, if $\theta_R = \pi/2$, $E_{\text{RKKY}}^{\text{ex,AA}}$ is FM only if the separation between adatoms is $3n$ graphene unit cells ($n \in \mathbb{N}$), otherwise it is AFM. The $n = 2$ case corresponds to Conf. \#5 defined in Sec. II A. For the adatom arrays $\Delta E$ can be approximated by summing the nearest neighbor interactions. For example, in the $\alpha(8 \times 4)$ array shown in Fig. 2, there are two different adatom pairs: (i) $\theta_n = \pi/2$, $n = 4$, (ii) $\theta_R = \pi/6$, $n = 4$. The sum of their contributions is AFM. The results on both conventional and resonant RKKY are summarized in Table I.

| Geometry | Conventional RKKY ($E_F < \varepsilon_0$) | Resonant RKKY ($E_F > \varepsilon_0$) |
|----------|---------------------------------|----------------------------------|
| $AA$     | FM                             | AFM                             |
| $AB$     | AFM                            | AFM                             |
| Bri      | AFM/FM                         | AFM/FM                          |

TABLE I. Dependency of the sign of $E_{\text{RKKY}}^{\text{ex}}$ to geometry of the system and to location of $\varepsilon_0$ with respect to $E_F$ and $E_D$. |
However, in practice the impurity states have some spatial spillover on the graphene backbone and therefore a hybridization occurs and an impurity band develops with a typical width proportional to the coupling strength.\textsuperscript{19} A weak coupling between the impurity states leads to a narrow band with a large DOS. When the Fermi level being adjusted by $V_g$ falls inside such an impurity band, ferromagnetic configuration becomes favorable. We refer to this region as the intermediate region. In order to study the onset of ferromagnetism, we consider the Hubbard Hamiltonian to account for the on-site electron-electron interaction:

$$H_U = \sum_i U n_{i\uparrow} n_{i\downarrow},$$  \hspace{1cm} (6)

where the summation is over all graphene and adatom sites, $n_{i\sigma}$ is the occupation number operator for electron with spin projection $\sigma$ at site $i$. Here $U$ is the on-site Coulomb interaction energy for the two electrons occupying carbon $p_z$ atomic orbital. As the result of the mean-field approximation the Hubbard model leads to the Stoner model,\textsuperscript{19} which typically yields ferromagnetism when the Stoner criterion is satisfied for the onset of magnetization:

$$U \cdot \text{DOS}(E_F) > 1.$$  \hspace{1cm} (7)

While DOS($E_F$) remains at about $1\;\text{eV}^{-1}$ for a moderate gate voltage applied to graphene, an impurity band induced by hybridization with the adatoms would produce a large DOS. The Stoner criterion is fulfilled if the DOS in the impurity band approaches the magnitude $\sim 10\;\text{eV}^{-1}$ and if $E_F$ adjusted by the gate voltage falls in this band. As for now, there are no direct experiments which allow to extract a value for the model parameter $U$ to adequately reflect the Coulomb-electron-electron interactions in graphene. In some works, it is reported to be $U \sim t$, where $t \approx 2.7\;\text{eV}$ is the tight-binding hopping parameter of graphene.\textsuperscript{75} Other experiments indicate that the strength of the Coulomb interaction in graphene is about an order of magnitude smaller.\textsuperscript{76} We will use this lower estimate of $U \sim 0.1\;\text{eV}$ which still justifies the existence of FM solutions based on the Stoner criterion.

### III. RESULTS

#### A. One C adatom per unit cell

We first focus on effects of gate voltage to the magnetic and geometrical properties of a single C adatom on $N \times N$ supercells, which leads to a periodic adatom coverage due to the periodic boundary conditions. Figure 3 illustrates the DOS of such system with a $4 \times 4$ SC, as well as $l, m$ quantum number decomposed PDOS (PDOS) projected to the adatom for different $\Delta Q$ values. In this PDOS one can see contributions from the magnetic $\psi_{p,\downarrow}$ orbitals and also from strongly hybridized lower-lying adatom states denoted as $\psi_{p,\uparrow}$. When 0.5 electrons have been removed from the unit cell, both $\psi_{p,\downarrow}$ and $\psi_{p,\downarrow}$ orbitals are unoccupied and degenerate in energy (Figs. 3(a) and 3(b)). When $V_g = 0$ (Figs. 3(c) and 3(d)), the $\psi_{p,\downarrow}$ orbital becomes partly occupied and the $\psi_{p,\downarrow}$ orbital is lifted in energy by Stoner splitting. Therefore, the C adatom network magnetizes and the magnetic moment per adatom is $M = 0.38\;\mu_B$ for PBE and $M = 0.24\;\mu_B$ for SCAN. The application of a positive $V_g$ fills the $\psi_{p,\downarrow}$ impurity state completely as illustrated in Figs. 3(e) and 3(f) for the case $\Delta Q = 1\;e$. Interestingly, at $\Delta Q = 1\;e$ the Fermi level is exactly at $E_F$ since the extra electron is completely absorbed by the $\psi_{p,\downarrow}$ orbital. Therefore, only electrons of the graphene matrix are present at the Fermi level.

Comprehensive magnetization results are given by $M(\Delta Q)$ curves presented in Fig. 4. For PBE, the curve is rather linear with a slope of about $0.65\;\mu_B/e$ until the magnetization saturates at $\Delta Q = 1\;e$. This slope is rather steep because the PDOS of the $\psi_{p,\downarrow}$ state is high compared to the graphene DOS. However, in the
region below $M \lesssim 0.2 \mu_B$ the curve presents a nonlinear behavior related to a breakdown of the Stoner model. For SCAN significant differences with respect to the PBE results can be noticed. In particular, the growth of the magnetization curve is steeper. Below $M \approx 0.5 \mu_B$ the slope is $1.4 \, e/\mu_B$, but it gradually decreases to $0.8 \, e/\mu_B$ before saturation. Interestingly, the magnetization slope is larger than $1 \, \mu_B/e$ for most of the curve, which implies that $\psi_{p,\perp}$ moves downwards in energy while being filled from the voltage bias, and therefore also attracts electrons from the graphene lattice. This feature of SCAN (which contains corrections to PBE) indicates a stronger response of magnetic moment to $V_G$ in comparison to PBE. Another experimentally verifiable SCAN result is that at $\Delta Q = 0 \, e$, the magnetization (0.24 $\mu_B$) is significantly reduced in comparison to the PBE result (0.38 $\mu_B$). We have observed similar trend in simulations with other SCs. This observation can explain the fact that intrinsic carbon magnetism is observed only in rare occurrences\textsuperscript{52–54,57,58} instead of being a common phenomenon. Moreover, our results indicate that gate voltage can be used as a control tool to enhance magnetism in various graphitic samples that are either weakly magnetic or even nonmagnetic. In fact, the same principle applies to various graphene-based magnetism, as recently experimentally demonstrated.\textsuperscript{11,14,16}

Our results with other SC sizes show that the slopes of $M(\Delta Q)$ curves become smaller when the adatom is placed on bigger SCs. This observation can be explained as follows. For bigger SCs there is a larger number of graphene states. Therefore, a larger $\Delta Q$ contribution occupies the graphene states instead of the $\psi_{p,\perp}$ band. However, smaller $M(\Delta Q)$ slopes do not necessarily mean weaker magnetic response to the gate voltage since this response (i.e., slope of $M(V_G)$) is related to the DOS peak width of the impurity state. In fact, the peak width clearly decreases with increasing SC size because the hybridization between the localized $\psi_{p,\perp}$ state with its periodic images is reduced. The peak widths at $\Delta Q = 0$ have the following trend for different SCs: \(3 \times 3: 0.23 \, eV; \quad 4 \times 4: 0.10 \, eV; \quad 5 \times 5: 0.10 \, eV; \quad 6 \times 6: 0.05 \, eV; \quad 9 \times 9: 0.03 \, eV\). Nevertheless, more dilute adatom coverage produces also smaller magnetization per surface area and weaker magnetic coupling between adatoms.

The gate voltage affects also geometrical properties according to Fig. 4. The inset of Fig. 4 defines $h(ad)$ as the height of the adatom $C$ with respect to the basal atoms, $h(bas)$ as the height of the basal atoms with respect to the nearest neighbor graphene atoms and $\Delta z$ as buckling of graphene. The total height of the structure is $\Delta z(\text{tot}) = h(ad) + h(bas) + \Delta z$. Without gate voltage, these values are for SCAN $h(ad) = 1.30 \, \text{Å}, h(bas) = 0.27 \, \text{Å}, \Delta z = 0.15 \, \text{Å}$ and $\Delta z(\text{tot}) = 1.72 \, \text{Å}$. $V_G$ does not have much effect on $h(ad)$ nor in overall on the shape of the nearly equilateral triangle formed by the adatom and the basal atoms. More noticeable is the upwards movement of the triangle when $\Delta Q$ is filled. This motion can be tracked in the increase of $h(bas)$ by 0.08 $\text{Å}$ and in the increase of $\Delta z$ by 0.04 $\text{Å}$. The PBE values are nearly identical for $h(ad)$, about 0.03 $\text{Å}$ larger for $h(bas)$ and 0.04 $\text{Å}$ to 0.06 $\text{Å}$ larger for $\Delta z$. The upwards motion of the triangle and increase in the buckling of graphene, produced by the filling of $\psi_{p,\perp}$ state, could be explained by the repulsive Coulomb interaction of the $\psi_{p,\perp}$ state with the $p_z$ orbitals of graphene. Nevertheless, $h(bas)$ and $\Delta z$ are significant even when the $\psi_{p,\perp}$ state is emptied.

We have also verified that our binding energies are consistent with previously calculated values of $-1.46 \, eV$\textsuperscript{40} and $-1.63 \, eV$\textsuperscript{42}. Our results yield $-1.46 \, eV$, $-1.52 \, eV$, $-1.51 \, eV$ and $-1.52 \, eV$ with $3 \times 3, 4 \times 4, 5 \times 5$ and $9 \times 9$ SCs, respectively. Therefore, the binding energies are not sensitive to the SC size until at high concentrations. For SCAN a slight decrease in binding energy was observed as for $4 \times 4$ and $5 \times 5$ SCs our results gave values of $-1.44 \, eV$ and $-1.46 \, eV$, respectively.

B. Remote interaction adatom array configurations

This subsection focuses on adatom arrays $\alpha$, $\beta$ and $\gamma$ (see Sec. II A on their definitions). These structures have two adatoms in the unit cell and they are suitable for description of remote interactions. Both one-sided adsorption and two-sided adsorption are considered. The long range interactions between the adatoms depend on their respective lattice positions and affect the magnetizations and energies of the FM and AFM solutions. Figures 5 (a) and 5 (b) illustrate the average magnetiza-
tion per adatom of the FM solutions ($\overline{M} = M/(FM)/2$) for $6 \times 3$ and $8 \times 4$ graphene SCs, respectively. These quantities are functions of the average added charge per adatom ($\Delta Q = \Delta Q/2$). As an example, the $\overline{M}$ curves of $\alpha(8 \times 4)$ are identical (but with fewer data points) to the $M$ curves in Fig. 4 because the FM $\alpha(8 \times 4)$ solution is equal to doubling the solution of the $4 \times 4$ SC with one adatom. The remaining $\overline{M}$ results are in overall similar. At $V_g = 0$ the SCAN magnetizations are smaller than for PBE, in fact for $\alpha(6 \times 3)$ it is even zero. The slopes of the $\overline{M}$ curves depend on SC size as discussed in Sec. III A, but they do not depend significantly on the array type. However, there is a dependency on the offset magnetization so that at $V_g = 0$ the PBE magnetizations vary from $\overline{M}(\alpha) = 0.17 \mu B$ to $\overline{M}(\gamma) = 0.44 \mu B$ for $6 \times 3$ SCs and from $\overline{M}(\beta) = 0.35 \mu B$ to $\overline{M}(\gamma) = 0.43 \mu B$ for $8 \times 4$ SCs. The corresponding SCAN magnetizations vary from $\overline{M}(\alpha) = 0.00 \mu B$ to $\overline{M}(\gamma) = 0.40 \mu B$ for $6 \times 3$ SCs. This variation of magnetizations can be connected to steep magnetization slopes and large DOS associated to the $\psi_{p, \perp}$ states. In these conditions, tiny energy shift produced by interactions of adatoms change $\psi_{p, \perp}$ occupations significantly. For this reason, the magnetization values are also sensitive to the computational details, particularly on the choice of XC functional and density of the $k$-point mesh. One can now understand why calculated magnetic moments vary significantly in the literature. However, with positive gate voltage the differences for the magnetization values become smaller. Moreover, $\overline{M}(\beta)$ and $\overline{M}(\gamma)$ robustly reach the saturation magnetism at around $\Delta Q = 1 e$.

We turn now to discuss an important result of this study concerning the energy differences $\Delta E = E(FM) - E(AFM)$. This quantity is plotted as a function of $\Delta Q$ in Figs. 5 (c) and 5 (d) for $6 \times 3$ and $8 \times 4$ SCs, respectively. The differences in $\Delta E$ between the respective one-sided and two-sided configurations are significant, but differences in the magnetization are not. We also observe very different energies in the different $\Delta Q$ regions, which have been discussed in Sec. III A. In the resonant region the $\psi_{p, \perp}$ orbitals are fully occupied by the definition of this region, and this condition occurs at $\Delta Q = 1 e$ for PBE and at smaller values for SCAN. Antiferromagnetism dominates in this region, as predicted by the resonant RKKY theory for impurity states below $E_D$. The representative values of $\Delta E$ are around $5 \text{ meV}$ and $2 \text{ meV}$ (both within PBE and SCAN) for $6 \times 3$ and $8 \times 4$ SCs, respectively.

In the intermediate region (with only partially filled $\psi_{p, \perp}$ orbitals), PBE and SCAN behave differently, but
in both cases a clear ferromagnetic valley is formed in the middle of the region for each array. Representative $\Delta E$ maximum values are about $-30$ meV and $-5$ meV for $6 \times 3$ and $8 \times 4$ SCs, respectively. In the case of structures with two-sided adsorption the energy difference magnitudes are larger for $\alpha$ but smaller for $\gamma$. For $\beta$ arrays, the ferromagnetism for two-sided adsorption is clearly weaker for the $8 \times 4$ SCs but for the $6 \times 3$ SCs there is no clear difference in the maximum $\Delta E$ magnitude. In general, the $\gamma$ arrays yield the most robust ferromagnetism, especially for the $8 \times 4$ SCs. The most stable FM solution depends on the occupation of the $\psi_{p,\perp}$ state. Regardless of the SC size or the XC functional, this solution corresponds to $\mathcal{M} = (0.40 \pm 0.05) \mu_B$, $(0.35 \pm 0.10) \mu_B$ and $(0.55 \pm 0.05) \mu_B$ for $\alpha$, $\beta$ and $\gamma$ arrays, respectively. These results can be understood within the Stoner model. In the AFM solution there is no hybridization between $\psi_{p,\perp}^1$ and $\psi_{p,\perp}^2$ impurity states, which are associated to $C_{1d}$ and $C_{2d}$ adatoms, respectively. On the contrary, in the FM solution $\psi_{p,\perp}^1$ and $\psi_{p,\perp}^2$ hybridize, which reduces the magnetization of the FM solution. The hybridization leads to a trade for potential energy by the expense of kinetic energy. The Stoner’s criterion reveals whether or not the trade was beneficial. Because of the peak in the DOS, the FM solution is stabilized. This phenomenon explains also why the maximum of the FM interaction is given at $\mathcal{M} \approx 0.5 \mu_B$, since the DOS reaches its maximum around half-filling. Incidentally, the Stoner model becomes irrelevant in the resonant region, where the impurity band is completely filled. One can also visualize the present phenomenology as follows. The hybridization between $\psi_{p,\perp}^1$ and $\psi_{p,\perp}^2$ creates bonding and antibonding states. The filling of the bonding state ($\mathcal{M} \rightarrow 0.5 \mu_B$) stabilizes the FM solution, while the filling of the antibonding state ($\mathcal{M} \rightarrow 1 \mu_B$) weakens it until the two contributions cancel. When SCAN and PBE $\Delta E$ are compared, the results in the most strongly FM region look similar, suggesting that PBE captures the Stoner phenomenology rather well. However, the SCAN FM stability is systematically slightly weaker. As a matter of fact, SCAN promotes orbital localization, since SCAN is more sensitive to chemical bonds than PBE by taking into account the kinetic energy density of the electrons.

For low magnetizations, SCAN yields strong AFM excursions especially for $\gamma(6 \times 3)$ and $\beta(6 \times 3)$ but also for $\alpha(6 \times 3)$, meanwhile this phenomenon is completely absent in the PBE results. These AFM anomalies can be explained within conventional RKKY. At low magnetizations, the $\psi_{p,\perp}$ orbitals become more localized for SCAN, but PBE cannot capture this localization. As discussed in Sec. II C, the Stoner model can break down in this dilute limit. In these conditions, the indirect exchange mediated by the conduction graphene electrons dominates. For low magnetizations, the integration of Eq. (3) has not yet reached the poles, thus $E_{\text{ex}}^{\text{RKKY}} \approx E_{\text{ex, conv}}$. Now, Eq. (5) explains that $E_{\text{ex, conv}}$ is slightly AFM for $\alpha(6 \times 3)$ and $\alpha(8 \times 4)$, strongly AFM for $\beta(6 \times 3)$ and even more strongly AFM for $\gamma(6 \times 3)$. This scenario qualitatively agrees with the SCAN results since strong AFM excursions appear for $\gamma(6 \times 3)$ and $\beta(6 \times 3)$ and a weaker one for $\alpha(6 \times 3)$. There is a small inconsistency with the conventional RKKY picture in the $\alpha(8 \times 4)$ curve, where the expected small AFM peak is absent. However, as discussed in Sec. 1.3 of the supplemental material, in this conventional RKKY region there is substantial numerical instability in the AFM solution, manifesting the delicateness of the underlying physics. Therefore, $\Delta E$ is sensitive to computational details and the SCAN results for low $\mathcal{M}$ may contain error in the meV range. Moreover, it should be noted that the Eq. (5) is only approximative for the bridge site and that $\alpha(6 \times 3)$ and $\alpha(8 \times 4)$ arrays contain competing AFM and FM interactions. Therefore, the conventional RKKY ferromagnetism for $\alpha(8 \times 4)$ is not conclusive, but the AFM peaks for $\gamma(6 \times 3)$ and $\beta(6 \times 3)$ are robust and thus confirm the conventional RKKY model.

### C. Close interaction adatom pair configurations

We now consider that the adatoms are in the neighborhood of each other by using close interaction pair Conf. #1–#8 in $7 \times 7$ and $6 \times 3$ SCs. A striking difference with respect to the remote interaction cases are nonlinear growths of the magnetizations curves displayed in Figs. 6(a) and (b). The $\Delta E$ values shown in Figs. 6(c) and (d) are also substantially larger than the corresponding remote interaction energies.

Let us first focus on the $7 \times 7$ SC structures, which represent well-isolated adatom pairs. In the region $\Delta Q \lesssim 0.5 \pi$, the magnetizations stay roughly constant or even decrease when more electrons are added. This behavior is produced by minority spin $\psi_{p,\parallel}^1$ orbitals. Normally, these states lie deep in energy (as shown in Fig. 3), but strong interactions between adatoms and large structural distortions raise these states to $E_F$ (see the DOS plots in Sec. 2 of the supplemental material). Upon increasing $\Delta Q$ both $\psi_{p,\parallel}^1$ and $\psi_{p,\parallel}^2$ are being occupied. These fillings yield opposing contributions to the magnetization. Thus increasing $\Delta Q$ often results in zero or even negative magnetization slope. Moreover, the $\psi_{p,\parallel}$ states may have different occupations at $C_{1d}$ and $C_{2d}$ at low $\Delta Q$ values, leading to adatom magnetic moments with different magnitudes. Thus in many cases the AFM solution becomes a ferrimagnetic solution with nonzero total magnetization (see Sec. 2 in the supplemental material for details about these cases). Furthermore, the $\psi_{p,\parallel}$ states bring complication in $\Delta E$ because the number of interacting states becomes higher and PDOS($E_F$) can have both significant spin up and spin down components. Besides, the graphene Dirac cone is severely deformed. Nevertheless, the values for $\Delta E$ reveal a preference for strong ferromagnetism with values as high as $-104$ meV (Conf. #6) but Conf. #7 and #8 display antiferromagnetism (or ferrimagnetism) with maximum magnitude of $40$ eV. These
results suggest high Curie temperatures in many of the studied configurations.

We now discuss higher gate voltages. Beyond the limit $\Delta Q \gtrsim 0.5 e$, the $\psi_{p,\parallel}$ states become fully occupied. In this regime, the magnetization and $\Delta E$ recover a behavior similar to the remote interaction cases discussed in the previous section. In the intermediate region ($0.5 e \lesssim \Delta Q \lesssim 1 e$), the magnetizations are linear and their values for the respective one-sided and two-sided adsorption models are almost equal. Moreover, each one stabilizes the FM solution (though some cases only weakly). In the resonant region, the magnetizations saturate and the antiferromagnetism is stabilized. However, a couple of exceptions with zero or even FM $\Delta E$ can be noticed. These cases do not contradict the resonant RKKY model since the $\psi_{p,\perp}$ states are not anymore strictly below $E_D$. For example, there may be splitting of the bonding and antibonding $\psi_{p,\perp}$ bands and the bands can be located exactly at $E_D$ or even above $E_D$.

The $6 \times 3$ SC structures have significant next-nearest neighbor interactions between $C_6^d$ and the periodic images of $C_6^d$, and vice versa. All these interaction distances are in the range $5 \AA$–$10 \AA$, which corresponds to the remote interaction case with FM character in the region $\Delta Q < 1 e$. Thus, in comparison to the $7 \times 7$ SC results, one expects to observe stronger ferromagnetism in Confs. #1–#6 and weaker antiferromagnetism or even ferromagnetism in the Confs. #7 and #8. Actually, Conf. #8 becomes FM and #7 less AFM. On the contrary, the ferromagnetism in Confs. #1–#6 decreases as typical $\Delta E$ values drop to around $-30$ meV and are about $-55$ meV at maximum. The reason is that the Stoner criterion is weakened because the tighter packing of the adatons spreads the $\psi_{p,\perp}$ energy band. The typical widths of the $\psi_{p,\perp}$ states in PDOS are $\sim 0.4$ eV but for the corresponding $7 \times 7$ SC configurations they are $\sim 0.1$ eV. In our tests with even tighter packing the ferromagnetism is further reduced. This behavior is consistent with Ref. 25, which argues that in band ferromagnetism with $sp$ character the magnetization must be inhomogeneous with only a fraction of the sample ferromagnetically ordered. In the resonant region the results are nearly equal with respect to the $7 \times 7$ SC cases.

In summary, our close interaction results show robustness of the Stoner and RKKY paradigms for the C adatom system on graphene. Even when the $\psi_{p,\parallel}$ states are present at $E_F$ (below $\Delta Q \approx 0.5 e$), the general trend is that the interaction energies are FM in the intermediate region, which implies that the ferromagnetism could persist in more complicated and experimentally realizable carbon systems. The gate voltage can be used to further optimize FM interaction strengths. Our study also
reveals that in this regard $6 \times 3$ is roughly the optimal SC size for two adatoms, which corresponds to one adatom per 18 graphene atoms. However, our results indicate that within this concentration, it is beneficial if adatoms are scattered inhomogeneously, since stronger ferromagnetism is observed for close interaction adatom pairs than for adatom arrays within the $6 \times 3$ SCs. Concerning the resonant RKKY, the antiferromagnetism persists in the resonant region even when the $\psi_{p,\perp}$ states are split and raised in energy.

D. The spin polarization maps

Experimentally, the adatom-induced spin polarization of the graphene sheet can be probed by using the scanning tunneling microscopy (STM), as shown by González-Herrero et al. for hydrogen adatoms. The DFT calculations also yield these spin patterns and they can be visualized as spatial spin polarization maps. Figure 7 illustrates these maps for selected C adatom configurations at $V_g = 0$.

Such map for a spin up C adatom fixed on the top site at $A$ sublattice ($M = 2.00 \mu_B$) is shown in Fig. 7(a). The magnetization pattern has triangular symmetry and the magnetizations at $A/B$ sublattice sites are always positive/negative, respectively. The polarization is caused by the impurity state which extends to the graphene lattice and by the polarized graphene electrons carrying the indirect interactions. The tail of the impurity state is composed of $p_z$ orbitals belonging to the $A$ sublattice sites. The $B$ sublattice $p_z$ orbitals are not involved due to the bipartite nature of the graphene lattice. The corresponding patterns are similar for many magnetic defects coupling only to one sublattice, e.g. substitutional transition metal atoms. The $B$ sublattice sites are negatively polarized because the spin up impurity state at the $A$ sublattice sites attracts spin up electrons from the $B$ sublattice sites, thus magnetizing them negatively. In addition another impurity state placed on the $B$ sublattice interacts with the spin down graphene states. The resulting indirect interaction with the original $A$ sublattice impurity is AFM in agreement with the Lieb’s theorem. However, in the case of the hydrogen adatom, the magnetization pattern is inverted (the $A/B$ sublattice sites have negative/positive magnetizations, respectively) as explained by Casolo et al.

The richness of the DFT simulations is also reflected in the case of bridge configurations. Figure 7(b) presents the magnetization density associated with a single adatom on a bridge site ($M = 0.43 \mu_B$). Our simulations indicate that the corresponding magnetic patterns are general for any magnetic impurities coupled equally to the both graphene sublattices. The amplitudes are mostly positive, in particular, one can notice four blobs between the basal atoms and their nearest neighbor graphene atoms and six smaller magnetization regions, two of which extend to two graphene sites. These regions are also visible in the spin polarization isosurface plot in Fig. 1. Further away positive and negative magnetizations alternate but positive amplitudes dominate similarly with the top-site situation. The magnitudes of the polarizations are strongly reduced when compared to the top site case for two possible reasons.
the $\psi_{p,\perp}$ state now spreads both to the A and B sublattices. Secondly, the magnetization of the C adatom is much higher for the top site configuration. In fact, by increasing gate voltage, we observe strengthening of the positive magnetizations and gradual disappearance of the negative magnetizations. The resulting pattern is highly asymmetric. Consequently, the direct coupling acquires a strong angular dependency to the vector connecting the adatoms. Moreover, the respective angle of the basal sites of $C_{12}$ and basal sites of $C_{21}$ (which can be $\theta_{bas} = 0^\circ$, $60^\circ$ or $120^\circ$) becomes relevant. The present results show an overall trend for stronger ferromagnetism for the $\theta_{bas} = 0^\circ$ angle, as in the case of $\gamma$ arrays and Confs. 2, 4, 5, and 6. One might achieve adsorption with $\theta_{bas} = 0^\circ$ by applying uniaxial strain, leading to longer bonds in one direction. The strain might also stabilize the adatom network. These properties of the strained system are of interest for future study.

Figure 7(c) visualizes the magnetization patterns for non-isolated Conf. 4 adatom pair ($\overline{M} = 0.70 \mu_B$), which corresponds to a FM interaction with $\Delta E = -30 \text{ meV}$. In this case, the positive magnetizations induced to graphene by the two adatoms mutually strengthen, thus a mainly positive magnetization develops on the whole unit cell. This effect can be also pictured as a stronger delocalization of the $\psi_{p,\perp}$ states. In this $\Delta Q = 0$ plot, negative polarization areas are still visible, however, they gradually disappear when more charge is added.

Figure 7(d) contains magnetization patterns for isolated Conf. 2 adatom pair ($\overline{M} = 0.50 \mu_B$), which corresponds to a FM interaction with $\Delta E = -40 \text{ meV}$. An interesting feature is the negative excursion of the magnetization in between the adatoms, which is much stronger than any positive magnetization in the contour map, even though the positive magnetizations typically dominate in the FM solutions, as shown in Fig. 7(c).

Although magnetization patterns of hydrogen have been recently observed by González-Herrero et al., at the moment no experiments have yet measured the polarization maps related to bridge configurations. Since our DFT calculations predict that these maps are completely different, STM experiments probing these cases would be useful to test our present models.

**IV. CONCLUSIONS**

By using DFT first principle simulations and various models of magnetism, we have proposed how to use gate voltage to control magnetism of carbon graphene adatoms. We have found that the voltage bias can be deployed to fine tune the magnetic moment of the adatoms from zero to $1 \mu_B$ by emptying or filling the localized magnetic state on the adatom. Moreover, the gate voltage influences the strength and sign (FM or AFM) of the magnetic interactions between the adatoms. We find AFM behavior at both low and high adatom magnetic moments, which can be explained within the conventional and resonant RKKY models, respectively. The low-magnetization antiferromagnetism is only found in the DFT simulations within the more accurate SCAN meta-GGA framework. Ferromagnetism is predicted for intermediate magnetic moments, which can be rationalized by introducing Stoner band ferromagnetism. The DFT calculations reveal that both GGA and SCAN predict strong FM configurations in the intermediate magnetic moment region, nevertheless there is a small reduction when SCAN is applied. Calculated spin polarization maps show highly non-trivial magnetic distributions in space, which could be probed with magnetic scanning microscopes. The present results can be generalized and can explain some experimental findings concerning magnetism in systems based on graphene, useful for future spintronics applications.

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