The nature and strength of inter-layer binding in graphite

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We computed the inter-layer bonding properties of graphite using an ab-initio many body theory. We carried out variational and diffusion quantum Monte Carlo calculations and found an equilibrium inter-layer binding energy in good agreement with most recent experiments. We also analyzed the behavior of the total energy as a function of interlayer separation at large distances comparing the results with the predictions of the random phase approximation.

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The excitement generated by the ability to fabricate graphene layers and possibly to tune their electronic properties has renewed great interest in weak interactions in graphitic systems \cite{1}. The hope of using graphene as a component for next generation electronics relies, among other things, on a detailed understanding and control of how it interacts with its surrounding (e.g. with supporting substrates)\cite{2}.

Unfortunately the nature and strength of binding in graphitic materials are poorly understood. For example, large uncertainties are associated with a fundamental physical quantity such as the strength of inter-layer binding in graphite. In addition, the way the interaction between planes decays as a function of distance is controversial\cite{3}\cite{4}, casting doubts on our current understanding of weak binding in carbon based system. Both binding strength and power law behavior of interlayer interactions in graphite are relevant to the comprehension of a multitude of materials, including graphite intercalation compounds, novel nano-electronic components, and carbon based systems for hydrogen storage.

From a theoretical standpoint, unravelling binding in graphitic systems is intimately related to understanding the role played by dispersion forces, and to acquiring the ability to describe these purely quantum mechanical interactions at a high level of accuracy. Local (LDA)\cite{5} or semi-local\cite{6} approximations to Density Functional Theory (DFT) do not correctly describe long range correlations, due to the local character of the exchange and correlation potential. Progress\cite{7} has been recently made in including dispersive interactions within a DFT formalism, in a self consistent, non empirical manner and binding between graphene layer has been predicted. In Ref.\cite{7} the authors report an interlayer distance overestimated by 7% with respect to experiment, and a binding energy of 45.5 meV/atom consistent with most extrapolated measurements of exfoliation energy\cite{12}. (Note that the calculations of Ref.\cite{7} are for two isolated graphene layers, not for a graphite solid). Semi empirical methods\cite{8}\cite{9} have been often used to treat dispersion correlations where DFT energies are corrected with a contribution coming from attractive $C_6/R^6$ potentials between pair of nuclei. However the power law behavior used in empirical and semi-empirical approaches to describe non retarded dispersion forces has been recently questioned\cite{8}.

At present, no direct measurement of graphite binding energy is available. However estimates based on theoretical models have been reported in the literature, using experimental data for exfoliation energies (EE, the energy required to remove one graphene plane from the surface of a graphite solid). Three experiments have reported data for EE\cite{10}\cite{11}\cite{12}, and a common aspect to the experimental analyses is the use of simple, fitted force fields to model either C-C or C-H interactions. The early work of Girifalco and Lad\cite{12} gave a EE value of 43(5) meV/atom. Using a Lennard-Jones potential, the difference between exfoliation and cleavage energy (that is the interaction between two semi-infinite crystals) was estimated to be 18% but the exact difference remains unknown. The work by Benedict et al.\cite{11} extrapolated the interaction energy between graphite layers (33 meV/atom) from measurements on collapsed nanotubes, using a force field to model the tubes elastic properties. More recently Zacharia et al. \cite{10} performed detailed desorption experiments of aromatic molecules from a graphite surface. The graphite EE was derived by extrapolating the molecules EE as a function of the number of carbon atoms, thus obtaining a value of 52(5) meV/atom. This yields an estimate of the cleavage energy (62 meV/atom) which is twice as much as that reported in Ref.\cite{11}.

Given the state of experiment and theory in determining the binding in graphite, there is a clear need for accurate calculations, eliminating as much as possible all approximations used so far in the literature, and possibly providing guidance to future experiments. Here we report the binding curve of graphite in AB stacking as a function of the number of carbon atoms, thus obtaining a value of 52.5 meV/atom. This yields an estimate of the cleavage energy (62 meV/atom) which is twice as much as that reported in Ref.\cite{11}.

We computed the inter-layer bonding properties of graphite using an ab-initio many body theory. We carried out variational and diffusion quantum Monte Carlo calculations and found an equilibrium inter-layer binding energy in good agreement with most recent experiments. We also analyzed the behavior of the total energy as a function of interlayer separation at large distances comparing the results with the predictions of the random phase approximation.

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in accord with the measurements of Zacharia et al. \[10\].
We find that at distances between 4 Å and 8Å the total energy curve exhibits a \( D^{-4.2} \) behavior as a function of inter-layer spacing \( D \), which is very similar to that predicted by the Random Phase Approximation (RPA) applied to two-dimensional (2D) semi-conducting layers \[2\].

In our investigation we have carried out variational Monte Carlo (VMC) and Lattice Regularized Diffusion Monte Carlo (LRDMC) calculations \[10\] with the TurboRVB code \[17\]. Our many body wavefunction is the product of a Slater Determinant and a Jastrow many body factor. The determinant is obtained with \( N/2 \) molecular orbitals \( \psi_j(\vec{r}) \), each doubly occupied by opposite spin electrons ( \( N \) is the total number of electrons). The orbitals \( \psi_j(\vec{r}) \) are expanded in a Gaussian single-particle basis set \( \{ \phi_i \} \), centered on atomic nuclei, i.e.
\[
\psi_j(\vec{r}) = \sum_i \lambda_{i,j} \phi_i(\vec{r})
\]

Electron correlation effects are included in our wave function (WF) through the Jastrow factor \( J(\vec{r}_1, \cdots, \vec{r}_N) = \prod_{i<j} \exp(f(\vec{r}_i, \vec{r}_j)) \), where \( f(\vec{r}, \vec{r}') \) depends only upon two-electron coordinates. The function \( f \) is expanded in a basis of Gaussian atomic orbitals \( \phi_i \):
\[
f(\vec{r}, \vec{r}') = \sum_{i,j} g_{i,j} \phi_i(\vec{r}) \phi_j(\vec{r}')
\]
The convergence of this expansion is improved by adding an homogeneous term and a one body contribution, thus satisfying the electron-electron and the electron-ion cusp conditions, respectively \[15\] [18]. The basis set used for the Jastrow includes 2s2p Gaussian orbitals. By optimizing the coefficients \( g_{i,j} \), we can treat in a non perturbative way the dynamical transitions to high angular momentum atomic states for pairs of electrons localized around each atoms. As shown for the benzene \[19\] and water dimer \[19\], these transitions are responsible, at the first order of perturbation theory, for the weak attractive dispersive forces between atoms at large separation.

In the following, the molecular orbitals \( \psi_j \) in the Slater determinant are obtained from a self-consistent DFT-LDA calculation. One may then optimize only the Jastrow factor, by keeping fixed the determinant built from LDA orbitals (hereafter referred to as J-DFT WF approach); alternatively one may simultaneously optimize both \( J \) and the determinant using the method described in Ref.\[20\]. The minimal Gaussian basis set required to build an accurate determinant was chosen by comparing graphite binding energies (BE) \[20\] as obtained using plane waves (PW) and Gaussian basis sets (see Fig.\[1\]). We note that at each atomic positions, PW calculations are free of basis set superposition errors and they can be converged by controlling one parameter, the kinetic energy cutoff. In the case of Gaussian, we used an even tempered local basis where the parameters \( \alpha_l \) and \( \beta_l \) of the Gaussian exponent \( Z_l = \alpha_l/2 \beta_l^2 \) of each angular momentum \( l \), were optimized by performing a series of total energy LDA calculations. We considered two basis sets: 4s4p2d and 8s8p4d. For both of them we computed the binding energy (BE) of graphite at the LDA level for a system of 32 atoms using only the \( \Gamma \) point. The 8s8p4d basis set reproduces the same BE curve obtained with PW converged with respect to the kinetic energy cutoff (90Ry).

In our QMC calculations we simulated a 2 \( \times \) 2 \( \times \) 1 and 2 \( \times \) 2 \( \times \) 2 super-cell with periodic boundary condition, containing 32 (128 electrons) and 64 atoms (256 electrons) respectively. The carbon valence-core interaction was described by a energy-consistent pseudopotential \[21\]. In our calculations we fixed the in-plane geometry to the one determined experimentally (C-C distance = 1.42 Å). We verified, at the LDA level, that a change as large as 10% in the in-plane carbon-carbon length affects the inter-plane equilibrium distance by less than 3%, while the BE changes by 1 \( \rightarrow \) 2meV/atom.

In Table I we compare the results obtained by full wave function optimization with those of the J-DFT WF approach for graphite at inter-layer distance \( D = 3.7 \) Å. The latter provides a reasonably accurate variational guess. However, full optimization of the WF parameters (including the exponent of the basis set) yields a decrease of energy per atom of 34(3) meV/atom, which is of a relevant magnitude to the energy scale we wish to investigate in this work. The LRDMC energy is much less sensitive to the initial state (guiding function), used in this ground state projection technique. In fact the optimization of Jastrow and Determinant in the guiding function, leads to a consistent LRDMC energy. The quantitative agree-
ment between the VMC and LRDMC calculations confirms that the key ingredients of the electron correlations are already included in our Jastrow factor. In the following the results for the $2 \times 2 \times 1$ super-cell are obtained by fully optimizing the wavefunction, while, due to the computational cost of the optimization, we will present only LRDMC calculations for the 64 atom system.

After assessing the accuracy of the guiding wave function, we considered finite-size (FS) effects. We expect the errors due to FS effects in the in-plane directions (i.e. $x$ and $y$ directions) to cancel out to a large extent, as we compute energy differences between systems (graphite and graphene) with rather similar bonding and electronic properties. In-plane FS errors arising from the kinetic and Hartree terms (one-body corrections) can be treated within a standard DFT approach with appropriate k-point sampling. Other FS errors come from the artificial periodicity of the exchange-correlation hole due to the periodic Coulomb potential. Kwee, Zhang and Krakauer (KZK) proposed to calculate the two-body corrections within LDA where the exchange and correlation functional is replaced by the LDA energy parametrized for a finite system. We applied KZK corrections as implemented in the PWSCF code. We cannot rely on any error cancellation in the $z$ direction, i.e. F.S. errors due to a finite number of graphene layers in the simulation cells. In addition, the KZK method cannot provide a robust correction scheme in this case due to the lack of long range effects in the LDA exchange and correlation functionals. We estimate the long range behavior of the interaction between planes by fitting the results of calculations performed on the $2 \times 2 \times 1$ super-cell at distances $D > 4 \AA$. These VMC (LRDMC) results are reported in Fig. 2 and show a behavior $E(D) \sim D^{-\alpha}$ with $\alpha = 4.2(1)/4.2(3)$. Although the LRDMC data are affected by larger error bars, we can safely conclude that $\alpha \geq 4$. We note that using the RPA applied to 2D systems, Dobson et al. found a power law behavior $\sim D^{-3} \log(D/D_0)$ for infinite $\pi$-conjugated layers. One does not expect to see the asymptotic form of Ref. in our work because the unusual interaction, arising from coupling between long-wavelength fluctuations in the plane, it is expected to arise at much larger distances than those studied here.

Using the power law determined in our calculation we can derive a scaling relation between the graphite BE and the number of layers $n$. Integrating over the super-cell volume we find that the total energy scales as $\sim 1/D_{max}^3$ where $D_{max}$ is the linear size of the super-cell in the $z$ direction, i.e as $\sim 1/n^3$. In Fig. 2(b) we report the BE obtained within the LRDMC method as a function of $\sim 1/n^3$. The BE curves close to the minimum for the 32 and 64 atoms cells are reported in Fig. 3. Extrapolating the results reported in Fig. 2(b) to an infinite number of layers, we obtain a value of the BE of 60(5) meV/atom. This is reduced to 56(5), after adding zero point motion ($\sim 2$ meV/atom) and lattice vibrational contributions at 300 K ($\sim 2$ meV/atom), as computed from vibrational free energies using the data of Ref. 28 for phonon frequencies.

Absorption experiments of aromatic molecules on graphite provide a measurement of the EE, while the cleavage energy is estimated to be 18% larger than exfoliation, on the basis of force field calculations. The cleavage energy is close, although not identical, to the BE defined here. Therefore our comparison with experiment can only be indirect, as we computed BE, while experiment reports EE. Nevertheless it appears that our computed BE ($\sim 56$ meV/atom) is in good agreement with the estimate for the cleavage energy from the most recent experiments: 62 meV/atom. We note that in the analysis of experimental results, one makes use of fitted force fields to evaluate the contribution to EE of carbon-

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**TABLE I: Variational (VMC) and Diffusion (LRDMC) total energy and energy variance ($\sigma$) obtained using two different guiding wave functions for the $2 \times 2 \times 1$ super-cell at a separation distance $D = 3.7 \AA$. $J-DFT$ denotes a wave function with optimized Jastrow and a determinant part from LDA calculations. “Opt.” refers to the guiding wavefunction where only LRDMC calculations for the 64 atom system.**

| Basis Set | Method       | $E$ (eV/atom)   | $\sigma$ (eV/atom) |
|----------|--------------|-----------------|--------------------|
| 4s4p2d   | Opt. VMC     | $-154.428(3)$   | $1.79(5)$          |
| 8s8p4d   | J-DFT VMC    | $-154.505(1)$   | $1.66(1)$          |
| 8s8p4d   | Opt. VMC     | $-154.540(1)$   | $1.60(1)$          |
| 4s4p2d   | Opt. LRDMC   | $-154.787(8)$   | ...                |
| 8s8p4d   | J-DFT LRDMC  | $-154.891(5)$   | ...                |
| 8s8p4d   | Opt. LRDMC   | $-154.899(5)$   | ...                |

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**FIG. 2:** Binding energy (BE) of graphite as a function of separation between planes (a), and of the number of layers $n$ included in the periodic cell used in our calculations (b). Note the logarithmic scales. In (a) results obtained with VMC and LRDMC are reported. In (b) only LRDMC results are shown.
hydrogen bonds. This is needed because an extrapolation is made on hydrocarbon adsorption energies, as a function of the number of C atoms. Although the force field parameters were adjusted to experimental data, it is unclear whether the assumption of additivity of forces close to the minimum is fully justified. In addition, the ratio between EE and BE is at present unknown and could only be estimated.

While the value of the BE can be extrapolated for an infinite number of planes (as in Fig. 2), the value of the equilibrium separation cannot. From the minima of the curves reported in Fig. 3, we obtain 3.350(24) and 3.243(26) Å at the VMC and LRDMC level, respectively, for the 2 × 2 × 1 cell, and 3.426(36) Å at the LRDMC level, for the 2 × 2 × 2 cell. Difficulties arising from very flat BE curves and, most importantly, from the lack of an extrapolation procedure as a function of the number of layers prevent us to find a fully converged equilibrium bond-length. The value found for the 2 × 2 × 2 cell is in good agreement with experiment (2% overestimate).

In conclusion, we have investigated the bonding properties of graphite in AB stacking, providing for the first time an estimate of the binding energy and long range behavior of the total energy based on ab-initio, many body theory. Our calculated binding energy is in good agreement with most recent experiments, providing a benchmark result for future calculations and further experimental measurements. The interaction energy between planes varies as $D^{-4.2}$, i.e. with a power law close to that found for two semiconducting planes.

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The BE is defined as $E_{n\text{-layer}} - n \times E_{1\text{-layer}}$, where $E_{n\text{-layer}}$ is the total energy of a graphite periodic cell containing $n$ layers at the optimized, equilibrium position and $E_{1\text{-layer}}$ is the total energy of a graphene layer.

Corrections to the two-body term may also be treated using the method proposed in Ref. [25], which requires the knowledge of the structure factor $S(q)$ for $q \to 0$ vectors. In principle $S(q)$ can be directly evaluated using QMC calculations. In practice, for semi-metallic graphite and graphene small error bars are difficult to achieve. In Ref. [26] the FS correction schemes proposed in Ref. [25] and [23] were shown to perform equally well.

We note that sums of pair-wise forces proportional to $1/R^6$ yield a behavior of the type $E \sim D^{-4}$. Of course our results showing a $\sim D^{-4}$ behavior are not to be taken as confirming that description of binding in graphite with pair-wise $1/R^6$ forces is correct.

For the $2 \times 2 \times 2$ super-cell the position of the minimum was estimated considering LRDMC calculations at lattice space of $a = 0.4$ Å. In the case of the $2 \times 2 \times 1$ cell we observed that the extrapolation procedure for $a \to 0$ does not affect the equilibrium position.