

Accurate global potentials for the interaction between rare gases and graphene-based surfaces. A joint atom-bond and computational approach

Massimiliano Bartolomei¹, Estela Carmona-Novillo¹, Marta I. Hernández¹, José Campos-Martínez¹ and Fernando Pirani²

¹ Instituto de Física Fundamental, IFF-CSIC, c/ Serrano, 123, 28006 Madrid, Spain
maxbart@iff.csic.es

² Dipartimento di Chimica, Università di Perugia, I-06123, Perugia, Italy

The interaction between atoms and molecules with graphene type surfaces is a topic of great interest given the potential new applications of these materials. For example it is in principle possible to detect with high resolution the presence of a single atom adsorbed on these surfaces[1]; also its porous sheets synthesized with atomic precision[2] have been proposed as a nano-scale membrane which could be used as an atmospheric nanofilter[3]. Thus in this kind of studies it is necessary not only a reliable and accurate description of the interaction potential but it is also fundamental to obtain a convenient parametrization of the involved non-covalent interactions to support dynamical studies on the physisorption of atoms and molecules.

We report reliable global potentials for the physisorption of rare gases with graphene and graphite surfaces amenable for a variety of dynamics simulations.

An atom-bond pairwise additive form[4] of the potential is used, where the interaction pairs, represented by proper analytical functions, are constituted by the Rg atom (Rg= He, Ne, Ar, Kr) and the C-C bonds of the graphene sheet(s). The parameters of the atom-bond pair potential, derived from the polarizability of the interacting partners, are fine-tuned exploiting calculations of the prototypical Rg--coronene system(see Fig. 1) using high level electronic structure methods, such as DFT-SAPT[5] and MP2C[6], and large basis sets. The atom-graphene/graphite potential is further expanded in a Fourier series and it is found that, for an accurate representation of the interaction, only a small number of corrugation terms need to be added to the laterally averaged potential. Furthermore, this corrugation part of the potential is identical for both Rg--graphene and Rg--graphite; in other words, inner layers of graphite only play a role in the laterally averaged Rg-graphite potential. For all systems, the hollow at the center of the carbon ring is preferred over the bridge and top adsorption sites (see Fig. 1), although diffusion barriers are low. Present results[7] are found to compare well with previous data[8] regarding well depths and equilibrium distances at different adsorption sites and, for graphite, the long-range dispersion coefficient C_3 . Moreover for Rg-graphene an estimation of the long-range dispersion coefficient C_4 is provided for the first time.

In addition, the interaction between Rg and porous graphene is modeled and barriers for penetration through the pores have been obtained; a potential use for lighter gases separation from the heavier ones is proposed.