Carbon cage-like materials as potential low work function metallic compounds: Case of clathrates.
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We present an ab-initio calculation of the electronic affinity of the hypothetical C-46 clathrate by studying its bare and hydrogenated (100) surfaces. We show that such a system shares with the diamond phase a small electronic affinity. Further, contrary to the diamond phase, the possibility of doping endohedrally these cage-like systems allows to significantly raise the position of the Fermi level, resulting in a true metal with a small work function. This is illustrated in the case of the Li$_8$@C-46 doped compound. Such a class of materials might be of much interest for the design of electron-emitting devices.

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There is a strong incentive in trying to synthesize materials which could be used as efficient electron-emitters for vacuum electronic devices. While alkali metals display the lowest work function (W $\sim$ 2-3 eV) of all elemental metals, their poor mechanical and chemical properties (melting temperature, resistance to ionic bombardment, etc) forbid their use in actual devices, and therefore metals with much higher work function such as tungsten or molybdenum (W $\sim$ 4.5 eV) are preferred for their superior mechanical and chemical properties.

A promising class of materials which have been extensively studied are carbon diamond based systems. Due to their large gap, carbon diamond based systems have been shown to display a small, or even negative, electronic affinity [1, 2]. Further, their mechanical properties are superior to the one of any existing metal. However, the difficulty of doping them "n-type" in order to obtain a significant density of emitting electrons limits their efficiency as good emitters.

In the last few years, a novel structure has attracted much attention for its specific structural and electronic properties. Column-IV clathrates are cage-like materials composed of face-sharing $X_{20}$, $X_{24}$ and $X_{28}$ clusters (where X=Si,Ge). All atoms are in the same sp$^3$ environment as in the diamond phase, but the network is composed of 87% of pentagonal rings. As a result, it has been shown in particular that Si clathrates present a band gap which is $\sim$ 0.7 eV larger than the one of the diamond phase [3, 4]. Furthermore, the possibility of filling each cage by a dopant atom allows to significantly raise the position of the Fermi level to yield a metallic system with a large density of states (eDOS) at the Fermi level ($E_F$) [5, 6].

The possibility of efficiently doping such systems has incited us to study the case of the hypothetical carbon clathrates. Even though such phases have not been synthesized so far, the reports on the synthesis of connected C$_{36}$ clusters [7] of diamond-like thin films synthesized by deposition of small carbon clusters [8] or even of polymerized C$_{60}$ suggests that the synthesis of such phases might be at hand or that the results presented below may apply to the above-mentioned existing phases.

We present in this Letter an ab-initio study of the electronic properties of C-46 and Li$_8$@C-46 within the density functional theory (DFT) [9]. We study in particular the C-46 (100) bare and hydrogenated surfaces and analyze the effect of doping on the position of the Fermi level, the affinity of the system and the density of states at $E_F$. We show that under doping such a class of materials are true metals with a low work function, thus combining the advantages of diamond-like systems with a large carrier concentration.

The local-density approximation (LDA) [11] was used for all calculations and a pseudopotential approach was...
adopted \[12\]. We used the Siesta package \[13\], which is a self-consistent DFT code, employing numerical atomic-like orbitals (NAO) as a basis set. A well-converged basis set, consisting of doubled \(\{s,p_x,p_y,p_z\}\) orbitals plus polarization \(d\)-orbitals was applied. Some of the calculations were also performed using a standard plane-wave (PW) basis set. All systems studied below were fully relaxed, both with respect to cell size and atomic positions. To perform the electronic affinity analysis, we calculated the plane-averaged self-consistent potential in the direction perpendicular to the surface for each of the slabs. The positions of the valence-band maximum (VBM) and conduction-band minimum (CBM) were found by adding the energy difference between the average self-consistent potential in the bulk and the VBM (CBM) in the bulk to the average self-consistent potential inside the slab.

Before studying the C-46 clathrate phase, we have tested the method by exploring the electronic and structural properties for the well-known bare and hydrogenated diamond (100) surfaces. Following Ref.\[1\], we have adopted a slab geometry with 10-layers but with 2-atoms per layer. The k-point sampling was thus increased to a \(4\times4\times1\) special grid \[14\]. A vacuum of 10 Å was kept between neighboring slabs.

The results for PW and NAO-basis are reported in Table I. As in Ref.\[1\], we observe that the hydrogenation, by building a surface dipole, leads to a negative affinity system. Our results are in good agreement with those previously published \[1\], and the NAO-basis calculations reproduce within \(\sim 0.15\) eV the results of the PW calculations.

As the DFT-LDA is well known to underestimate the band gap of insulators, we have performed further a quasiparticle study within the GW approximation \[15\] for the bulk diamond and C-46 phases. Such an approach is known to yield quasiparticle energies to within 0.1 eV as compared to experimental photoemission data. More details about the formalism and results will be provided elsewhere \[15\]. The main outcome of such calculations is that the quasiparticle correction to the DFT-LDA eigenstates are very similar in both systems and that the CBM is pushed to higher energy by \(\sim 0.7\) eV in both phases (while the VBM is corrected down by \(\sim 0.8\) eV). This allows to correct the calculated affinity as reported in the right column of Table I.

### Table I: Calculated electronic affinity for the diamond and clathrate carbon surfaces (eV).

| surface          | LDA-PWs | LDA-NAOs | GW     |
|------------------|---------|----------|--------|
| C (100)          | 2.17    | 1.99     | 1.3    |
| C (100):H        | -0.45   | -0.59    | -1.3   |
| C-46 (100)       | 3.47    | 2.8      |        |
| C-46 (100):H     | 1.47    | 0.8      |        |
| Li\(_8\)@C-46 (100):H | 2.18 | \(\sim 1.5\) |        |

The stability and electronic properties of C-34 and C-46 clathrates have already been discussed in some details at the DFT-LDA level \[4, 17, 18, 19\]. In particular, it has been shown that for the C-34 and C-46 clathrates, the cohesive energy, bulk modulus and band gap are respectively \(\sim 0.1\) eV/atom, \(\sim 10\) % and \(\sim 0.2\) eV smaller than those of the diamond phase. As such, the stability and mechanical properties of carbon clathrates are close to those of the diamond phase. In the present work, we obtained for the lattice parameter, bulk modulus and band gap the values of 6.64 Å, 376 GPa and 3.75 eV respectively with both NAO and PW basis sets, which is in good agreement with previously reported results. Within the GW approximation, the gaps increase to 5.55 and 5.35 eV for the diamond and clathrate phases respectively.

We now study the C-46 (100) surface represented in Fig. 2. There are several non-equivalent (100) surfaces and we adopt a surface consisting of ”half emerging” C\(_{24}\) clusters. After structural relaxation, we find that the emerging semi-clusters hardly reconstruct besides a slight contraction inward of the three-fold coordinated surface atoms, leading to a minimum C-C distance of 1.40 Å on the surface. Hydrogen coverage is performed by ”passivating” these surface three-coordinated atoms. We note at this point that the results presented below may certainly change quantitatively with the chosen surface but the purpose of the present paper is to illustrate on this specific example the potentialities of such novel systems. Fig. 2 shows the calculated plane-averaged, self-consistent potential for bare and hydrogenated C-46 (100) surfaces.

**FIG. 2:** Plot of the plane-averaged self-consistent potential for the C-46(100) bare and hydrogenated surfaces, and for Li\(_8\)@C-46(100):H surface. The good agreement between the bulk and slab potential on the innermost atoms of Li\(_8\)@C-46(100):H indicates that the slab is large enough.
FIG. 3: Electronic density of states (eDOS) for bulk C-46 and Li$_8$@C-46. A 0.1 eV broadening has been used. The eDOS is normalized per carbon atom.

The electronic affinity is reported in Table I. The results obtained for the undoped compounds suggest that carbon clathrates are a priori less favorable for emission than, say, the (100) diamond surface, taking the electronic affinity as the only relevant criteria. However, we emphasize again that clathrates, contrary to the diamond phase, can be efficiently doped [19, 20] by intercalating one atom in each cage in order to obtain a true metallic system with a large density of electrons available for emission. This is a crucial advantage as compared to the diamond phase.

By analogy with the existing doped Si clathrates and intercalated carbon graphitic systems, we study the $n$-doped Li$_8$@C-46 phase. The corresponding bulk eDOS is represented in Fig. 3, where we have arbitrarily aligned the eDOS to the top of the C-46 valence bands for sake of comparison. As can be readily seen, intercalation of Li atoms leads to the filling of the conduction bands of the host carbon matrix by Li 2s electrons. The system becomes metallic with the Fermi level located near a peak in the density of states, which is in excellent agreement with previous work [19].

As in the case of diamond surfaces, hydrogenation leads to a significant decrease in the electronic affinity of the empty phase. We now explore the effect of doping on the electronic affinity (Table I) of the hydrogenated surface. The important finding is that Li doping leads to the formation of a true metallic system with a quite low work function ($\sim 1.5$ eV GW value). This work function is significantly lower than the one of alkali metals and much lower than the one of e.g. tungsten. The eDOS for states located around $E_F$ can be further shown to have a strong weight on both bulk and surface atoms, a factor which is crucial for emission. One can conclude that doped carbon clathrates are potentially true metals with a work function lower than the lowest affinity of all elemental metallic systems but with much superior mechanical properties.

It is interesting to note that the work function of the Li-doped clathrate is actually larger than the electronic affinity of the empty phase, despite the raise of the Fermi level under doping. This means that the entire band structure is shifted to lower energy with respect to the vacuum level under doping. This shift can originate in two effects: a decrease of the average potential inside the slab (bulk effect) and an increase of the surface dipole barrier (surface effect). An analysis of both effects show that in the present case it is mainly a bulk effect.

In conclusion, we have shown that doped carbon clathrates are potentially true metallic systems with a work function significantly lower than the one of alkali metals but with a stability typical of the carbon diamond phase. This opens new perspectives for the making of efficient electron-emitters. An alternative choice of dopant atom and/or surface treatment (e.g. cesium coverage) may lead to even smaller work function.

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