Sodium atoms and clusters on graphite by density functional theory

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Sodium atoms and clusters (N≤5) on graphite (0001) are studied using density functional theory, pseudopotentials and periodic boundary conditions. A single Na atom is observed to bind at a hollow site 2.45 Å above the surface with an adsorption energy of 0.51 eV. The small diffusion barrier of 0.06 eV indicates a flat potential energy surface. Increased Na coverage results in a weak adsorbate-substrate interaction, which is evident in the larger separation from the surface in the cases of Na3, Na4, Na5, and the (2×2) Na overlayer. The binding is weak for Na2, which has a full valence electron shell. The presence of substrate modifies the structures of Na3, Na4, and Na5 significantly, and both Na4 and Na5 are distorted from planarity. The calculated formation energies suggest that clustering of atoms is energetically favorable, and that the open shell clusters (e.g., Na3 and Na5) can be more abundant on graphite than in the gas phase. Analysis of the lateral charge density distributions of Na and Na3 shows a charge transfer of ~0.5 electrons in both cases.

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I. INTRODUCTION

Graphite is a semimetal that is widely utilized in experimental surface physics. The planar geometry and weak van der Waals–type interlayer interaction make it possible to split flat, chemically inert, and clean graphite (0001) surfaces [highly oriented pyrolytic graphite (HOPG)], which are ideal for studying adsorption layers and clusters. The electronic 2D semimetal properties of graphite are well-known both experimentally and theoretically. For instance, density functional theory (DFT) has provided information about the valence charge density, electronic density of states, band structure, elastic constants, and equation of state.1–4

An interesting research field considers alkali metal atoms and clusters on graphite. Reactivity and metallic properties make alkali metals exciting both for nanotechnological applications and basic science, and the properties of adsorbed alkali metal atoms on HOPG evolve as a function of coverage. Initially, a dispersed and highly polarized phase (“correlated liquid”) is found where alkali atoms maintain a maximum distance between each other. After a critical density of adatoms is reached, a nucleation to more closely packed configurations (islands) occurs.5 Alkali metals seem to have a higher charge transfer to HOPG with lower coverage, and an increase in adatom density tends to reorganize the charge into the alkali metal layer forming a two-dimensional metallic state that has a small surface corrugation and is almost decoupled from the substrate.5–9 It has been proposed that alkali-metal-plated graphite could have practical applications as a substrate in studying normal and superfluid He films.6,10

Despite the similar electronic structure of alkali metals, deviations in island formation and interaction with HOPG are observed as the atomic number increases. While lithium atoms either intercalate between the graphene layers5 or form a planar inconsummrate hcp superstructure on HOPG,11 it has been suggested that sodium nucleates only in buckled (110) bcc overlayers.12,13 The larger alkali atoms (K,Ru,Cs) are found to intercalate via surface defects6,9 or to adsorb in a (2×2) phase occupying hollow sites of the hexagonal substrate.6,9,14,15 In addition, cesium can exist in an incomensurate hexagonal or a more sparse (√7×√7)R19.11° phase,6,14,15 and a dense (√3×√3)R30° structure has been proposed for potassium.16,17 Obviously, the above observations are related to the atomic radius and ionization potential of the alkali atom in question, which affect both the adatom-adatom and adatom-surface interactions.

The first experimental studies considered metal islands and metal layers on graphite. A more controlled treatment of adsorbates is challenging, and it is difficult to study separated atoms and size-selected small clusters. Contemporary experimental techniques are able to deal with the practical difficulties such as the substrate temperature, surface defects, kinetic energy of cluster deposition (“soft-landing”), and cluster aggregation.18 Theoretical studies concern mostly single atoms19–22 or atomic layers on graphite formed by periodic boundary conditions.7,8,23 However, research on supported clusters is needed because they form a bridge between isolated atoms and ordered nanolayers, and they may have nanotechnological importance (quantum dots, catalysis). Several attempts to model small clusters and molecules on HOPG have been made,8,20,21,24–29 but the large number of substrate atoms and the semimetallic nature of graphite (k points) make reliable calculations very demanding.

Various theoretical methods are capable of studying metal atoms and clusters on graphite (0001). In addition to deciding which theoretical tools to use, a crucial question is how to model a graphite surface, i.e., how many graphene layers are needed, how large should the substrate be, and does the adsorbate change the surface geometry? One approach is to place the metal cluster under study onto an isolated hydrogen-terminated piece of graphite (“cluster”) that mimics a continuous surface.19–22,29 The question then is how large should the graphite cluster be in order to get realistic results? On the other hand, there is a problem in optimizing the geometry of the substrate if several graphene layers are involved. This is due to the fact that the layers are interlocked, and the system (Bernal graphite, stacking ABAB) is not fully symmetric at the substrate edges. With periodic...
boundary conditions one can describe, in principle, a continuous infinite system ("slab") in the lateral dimensions. In this case, the problem is the distance between adsorbate replicas, which should be large enough to exclude charge density overlap. The large substrate that must be used increases the computational cost greatly.

In the present work, a DFT method with periodic boundary conditions has been used to model Na atoms and clusters (N ≤ 5) on HOPG. The substrate consisted of three graphene layers with 32 (60) carbon atoms each. It was found that the HOPG potential energy surface (PES) is very flat with the hollow site of the carbon hexagonal structure being preferred. Although alkali metal atoms tend to be more weakly bound to the surface when the coverage increases, this tendency is not so clear in small Na clusters. The calculated cluster energetics favor clustering processes on HOPG, and the stability of open shell clusters (Na_3 and Na_5) is increased.

II. SIMULATION METHODS

The calculations have been performed using the Car-Parrinello molecular dynamics (CPMD) package, which is based on density functional theory. The electron-ion interaction is described by ionic pseudopotentials having the non-local, norm-conserving, and separable form suggested by Troullier and Martins. Periodic boundary conditions are employed, and the plane wave basis has a kinetic energy cutoff of 70 Ry. The generalized gradient-corrected approximation of Perdew, Burke, and Ernzerhof is adopted for the exchange-correlation energy functional. The electronic Hamiltonian is rediagonalized after each geometry optimization step, and a finite temperature functional (T = 1000 K) by Alavi et al. is used for the Kohn-Sham (KS) orbital occupancies due to the small energy gap between the occupied and unoccupied states (band gap). The ionic positions are optimized using a conjugate gradient method until all the components of nuclear gradient are below 10⁻⁴ a.u.

We model two periodic substrates of Bernal graphite which consist of three graphene layers (stacking ABA) in orthorhombic supershells of 9.84×8.53×16.70 (96 C atoms) and 12.30×12.79×16.70 Å³ (180 C atoms). The smaller substrate with Na_3 is shown in Fig. 1 from two perspectives. Our tests for different numbers of graphene layers have shown that at least three layers are needed in order to reach a convergence in Na adsorption. The spacing between the layers is fixed to the experimental value 3.35 Å, since the PBE functional used has problems in describing weak van der Waals–type interactions. The choice of z dimension keeps the slab replica 10 Å apart, which is sufficient for most applications. However, a weak binding of Na_2 (and large separation from the surface) forced us to use 2 Å larger spacing in this case. For Na_4 and Na_5, the interaction between cluster replicas becomes significant in the smaller box, and a larger substrate in x and y dimensions is needed, where the minimum distance between the clusters is now 7.62 and 6.53 Å, respectively.

Extensive tests for different numbers of k points have shown that the simple Γ-point approximation is not reliable for the systems studied. This is manifested by an artificial planar elongation of the graphite hexagons during geometry optimization, and is probably related to a strong downward dispersion of the upper σ bands at the Γ point. The problem does not occur with a 2×2×1 Monkhorst-Pack k-point mesh, and a variation of lateral dimension results in a value 1.421 Å for the C-C nearest-neighbor distance (experimental value 1.420 Å). The Na adsorption energies obtained show that a 5×5×1 mesh is adequate (see also Table I), whereas the forces are already well converged for the 2×2×1 mesh. We have also tested whether it is possible to use a smaller kinetic energy cutoff: In comparison with 70 Ry, a calculation with 50 Ry yielded 0.17 eV (33%) weaker binding of Na, and the related perpendicular distance from the surface increased by 0.19 Å (7.8%). This is surprising since it is generally accepted that 50 Ry is appropriate for C using Troullier-Martins pseudopotentials, and a proper treatment of the smooth Na pseudopotential requires even fewer plane waves. Presumably, the underlying reason is the relatively weak interaction between the Na atom and the substrate that involves charge transfer. The situation is analogous to hydrogen-bonded systems where a high kinetic energy cutoff is required.

The effect of substrate relaxation has been studied by re-

<table>
<thead>
<tr>
<th>Point</th>
<th>ΔE (eV)</th>
<th>d (Å)</th>
<th>Na-C (Å)</th>
<th>N_C</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.12/0.51</td>
<td>2.45</td>
<td>2.83</td>
<td>6</td>
</tr>
<tr>
<td>1</td>
<td>0.08/0.47</td>
<td>2.49</td>
<td>2.59, 2.78</td>
<td>3</td>
</tr>
<tr>
<td>2</td>
<td>0.06/0.44</td>
<td>2.53</td>
<td>2.53</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>0.07/0.45</td>
<td>2.53</td>
<td>2.63</td>
<td>2</td>
</tr>
<tr>
<td>4</td>
<td>0.06/0.44</td>
<td>2.54</td>
<td>2.54</td>
<td>1</td>
</tr>
<tr>
<td>5</td>
<td>0.09/0.47</td>
<td>2.50</td>
<td>2.60, 2.78</td>
<td>3</td>
</tr>
</tbody>
</table>

FIG. 1. (Color) Optimized Na_3-HOPG system shown from two perspectives. The superslab size is 9.84×8.53×16.70 Å³. Each graphene layer consists of 32 atoms.
which causes increased symmetry in the PES. These findings
II, where Na
3

commensurate periodic structure with twice as many Na at-

6

D
3

estimate of 0.51 eV for the energy minimum
points enhances binding in a systematic way yielding to an
d
(2

C
points 2 and 4 above
energy surface with a maximum variation of 0.07 eV . The
D

3

different locations along the surface
HOPG system geometry. The changes are small (e.g., the
C-C distance 1.424 Å) which validates the use of fixed sub-
strate in real applications. A bench mark calculation for Na-
HOPG shows that the local spin density (LSD) approximation
does not improve the results because of the large number of
KS states involved and the nonmagnetic nature of the
system. The calculations below are done with spin-
degenerate KS orbitals except for isolated Na atom and Na
clusters.

3

II. RESULTS

In order to map the potential energy surface of a Na-
HOPG system, we have optimized the Na atom position for
different locations along the surface (see Fig. 2). The results
for adsorption energy (ΔE⊥), separation from the surface
(d⊥), nearest carbon atom distances (Na-C), and carbon co-
ordination numbers (Nc) are presented in Table I. Here, we
do not approach the real zero-density limit of Na, but the
atoms are distributed in 9.84×8.53 Å intervals due to the
periodic boundary conditions applied.16 Inclusion of more k
points enhances binding in a systematic way yielding to an
estimate of 0.51 eV for the energy minimum (point 0, 5×5
×1 mesh). Comparison with other locations shows only
small deviations in ΔE⊥ and d⊥, indicating a flat potential
energy surface with a maximum variation of 0.07 eV. The
points 2 and 4 above Cα and Cβ (Fig. 2) give similar results,
which causes increased symmetry in the PES. These findings
resemble the results by Lamoen and Persson8 who obtained
ΔE⊥ = 0.52 eV for a K-HOPG system and a small diffusion
barrier (variation 0.05 eV). No k points were used in these
calculations, but we expect a systematic shift in ΔE⊥ similar
to the one we found.

Table II shows the formation energetics of Na atoms and
clusters. The formation energy ΔE is divided into two com-
ponents: the binding energy of a free cluster or a separated
monolayer (ΔEb), and a term (ΔE⊥) describing the adsorp-
tion energy. Three phases of Na-HOPG are included in Table
II, where Na(I) refers to the initial sparse system, Na(II) is a
commensurate periodic structure with twice as many Na at-
oms per unit shell as Na(I) (Na-Na separation 6.51 Å), and
Na(III) corresponds to the (2×2) Na monolayer with four
times the coverage of Na(I) and hexagonal symmetry. The
phase Na(II) was encountered as a by-product of Na3 stretch-
ing, and it corresponds to the maximal separation of Na at-
oms allowed by the smaller supershell. The effect of
nearby Na atoms becomes clear in Na
3

dimer separation from the surface i s 1 Å larger than for Na
3

and Na
4
. The same effect is apparent in the

7

3

3

Nearest-neighbor binding is compensated by the interaction with other Na at-
oms, and the resulting ΔE per atom is slightly larger than for
Na(I). A similar (2×2) structure is stable for potassium,9
and theoretical studies have shown that K forms a metallic
state on HOPG.6–8 Our results corroborate this finding,
but—in the case of Na monolayer—the spacing between Na
atoms (4.92 Å) does not match typical Na-Na distances (see
Na clusters in Table III), and the energy difference with the
lower coverage phase Na(I) is relatively small.

The cluster formation energies in Table II reveal signifi-
cant differences between individual clusters. Na2 binds only
very weakly due to its closed valence electron shell, and the
dimer separation from the surface is 1 Å larger than for Na3
and Na4. The same effect is apparent in the ΔE⊥ values. It is

TABLE II. Na atoms and clusters on graphite (0001). ΔE and
ΔE⊥ are calculated for both the 2×2×1 and 5×5×1 k-point
meshes, but the geometries are optimized using the 2×2×1 mesh
alone.

<table>
<thead>
<tr>
<th></th>
<th>ΔE/atom (eV)</th>
<th>ΔE∥/atom</th>
<th>ΔE⊥/atom</th>
<th>d⊥ (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na(I)</td>
<td>0.12/0.51</td>
<td>0.12/0.51</td>
<td>2.45</td>
<td></td>
</tr>
<tr>
<td>Na(II)</td>
<td>0.15/0.33</td>
<td>0.09</td>
<td>0.07/0.24</td>
<td>2.81</td>
</tr>
<tr>
<td>Na(III)</td>
<td>0.46/0.57</td>
<td>0.19</td>
<td>0.27/0.38</td>
<td>3.21</td>
</tr>
<tr>
<td>Na3</td>
<td>0.40/0.48</td>
<td>0.34</td>
<td>0.06/0.14</td>
<td>3.95</td>
</tr>
<tr>
<td>Na4</td>
<td>0.56/0.68</td>
<td>0.35</td>
<td>0.21/0.34</td>
<td>2.95, 2.98</td>
</tr>
<tr>
<td>Na5</td>
<td>0.65/0.64</td>
<td>0.45</td>
<td>0.20/0.19</td>
<td>2.88, 3.12</td>
</tr>
<tr>
<td></td>
<td>0.68/0.71</td>
<td>0.46</td>
<td>0.22/0.25</td>
<td>3.08, 3.77</td>
</tr>
</tbody>
</table>

*Larger substrate of 60 atoms per layer and simulation box of
12.30×12.79×16.70 Å

TABLE III. Optimized structures of adsorbed Na clusters. Distances in Å and angles in degrees. The values in parentheses refer to
the gas phase structures.

<table>
<thead>
<tr>
<th></th>
<th>Na2</th>
<th>Na3</th>
<th>Na4</th>
<th>Na5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na-Na</td>
<td>3.07 (3.05)</td>
<td>3.35 (3.17)</td>
<td>3.53 (3.43)</td>
<td>3.24 (3.33)</td>
</tr>
<tr>
<td></td>
<td>3.26 (3.97)</td>
<td>3.27 (3.02)</td>
<td>3.34 (3.42)</td>
<td>3.62 (3.36)</td>
</tr>
<tr>
<td></td>
<td>4.11 (3.46)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>d⊥</td>
<td>3.95</td>
<td>2.95, 2.98</td>
<td>2.88, 3.12</td>
<td>3.08, 3.77</td>
</tr>
<tr>
<td>Na-C</td>
<td>4.52</td>
<td>3.10</td>
<td>2.92, 3.30</td>
<td>3.09-4.12</td>
</tr>
<tr>
<td>Angle</td>
<td>58.2 (77.7)</td>
<td>55.2 (52.2)</td>
<td>58.1 (61.8)</td>
<td>62.4 (63.7)</td>
</tr>
<tr>
<td></td>
<td>66.7 (59.5)</td>
<td>124.2 (127.8)</td>
<td>55.3 (58.7)</td>
<td>75.9 (60.5)</td>
</tr>
<tr>
<td></td>
<td>155.2 (177.7)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Torsion</td>
<td>8.7</td>
<td></td>
<td>14.9, 30.7</td>
<td></td>
</tr>
</tbody>
</table>
interesting that the deviation of $\Delta E$ for $2 \times 2 \times 1$ and $5 \times 5 \times 1 \mathbf{k}$-point meshes becomes smaller as the distance between Na atoms and surface increases (see, e.g., Na$_3$). This implies changes in charge transfer. The larger substrate used for Na$_4$ and Na$_5$ requires fewer $\mathbf{k}$ points to converge the formation energy, which can be seen as nearly identical $\Delta E$ values. The $\Delta E$ values of Na$_3$, Na$_4$, and Na$_5$ are larger than for the $(2 \times 2)$ Na monolayer, which shows that the clustering of Na atoms is preferred.

The optimized cluster structures are related to the ground state geometries of free Na clusters. For Na$_3$, Na$_4$, and Na$_5$ the corresponding isomers are an isosceles triangle, a rhombus, and a planar $C_{2v}$ isomer. The clusters are placed on HOPG in a way that assumes that the hollow site (point 0) is energetically favorable for each Na atom. The related bond distances, angles, torsional angles, and distances from the surface are given in Table III. As mentioned above, Na$_2$ binds weakly, and this can also be seen in the very small change in dimer bondlength (0.02 Å). For the other clusters changes are more obvious: Na$_3$ adopts a geometry close to an equilateral triangle with significant changes in bondlengths and angles. Na$_4$ bends away from planarity (torsional angle 8.7°), and the bondlengths increase systematically, but the angles remain close to the initial values. For Na$_5$ all atoms occupy similar sites on top of point 5 (see Figs. 1, 2), not far from hexagon centers. The atoms of Na$_4$ are coordinated with HOPG in two ways: the two corners of the rhombus are above C atoms and bent towards the substrate [a result contradictory to the PES of Na(I)], whereas the other two Na atoms are close to the hexagon centers. The geometry and position of Na$_4$ is shown in an electron density isosurface plot in Fig. 3.

The adsorption of Na$_4$ leads to a significant distortion from planarity, with the central Na atom being much farther from the surface (0.69 Å) than the other atoms. Simultaneously, the longest Na-Na bond is broken (4.11 Å, see Table III), and the resulting $C_s$ structure (Fig. 4) comprises two identical triangles connected via their apices. As for the other clusters, changes in bond lengths are considerable, and there are also changes in bond angles. The Na atoms are coordinated with the surface in three ways: the central atom is on a hollow site, the two atoms that initially comprised the broken Na-Na bond sit on top of C-C bonds, and the two corner atoms are directly above C atoms. Here, we have optimized the cluster geometry with respect to a substrate consisting of two graphene layers alone (120 C atoms). The obvious changes are an increased separation of the middle Na atom from the surface (0.20 Å) and a further elongation of the longest (broken) Na-Na bond (0.11 Å). The other bond distances and the formation energy $\Delta E$ are unchanged.

The electron density isosurface plot of Na$_5$-HOPG in Fig. 4 illustrates how the density is distributed within the Na$_5$ cluster. The largest values are obtained inside the two remaining triangles, but there is a component also in the interstitial region next to the broken (or elongated) Na-Na bond. The Na atom in the middle has a pronounced hole in the density, but an atom-centered integration of charge density within a small spherical volume ($R = 1.5$ Å) gives similar results (charges) for each Na atom. This is explained by the fact that the most coordinated Na atom has density contributions from both triangles. Presumably, this atom prefers a larger distance from the surface and a hollow site because of its higher Na coordination, whereas the lower coordinated Na atoms tend to acquire positions closer to carbons and C-C bonds. The same applies to Na$_4$, but on a smaller scale (see Fig. 3).

We have listed on Table IV formation energies for different cluster/atom products on top of the HOPG substrate, calculated assuming an initial state of $N$ free Na atoms in the gas phase. The most obvious feature is the small formation energy of Na$_2$ containing products. This is caused by the full valence electron shell of Na$_2$ that reduces binding with the substrate. Even two separated Na atoms are more stable on graphite than a dimer. For larger systems Na$_3$, Na$_4$, and Na$_5$ are favored, indicating clustering processes, and the product Na$_3$+Na is slightly higher in formation energy than Na$_4$. The unpaired electron on the outermost shell of Na$_3$ (and Na) increases binding with HOPG as seen in Table II. A similar conclusion can be made about the high stability of Na$_5$. This indicates that open shell clusters can be more abundant on graphite than in the gas phase.

The weak binding of Na$_2$ compared to two separated atoms has suggested to us to investigate the breaking of this
TABLE IV. Formation energies of Na products on graphite (0001). A large separation of end products is assumed.

<table>
<thead>
<tr>
<th>Reactants (free)</th>
<th>Products (graphite)</th>
<th>$\Delta E$ (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na</td>
<td>Na</td>
<td>0.51 eV</td>
</tr>
<tr>
<td>2×Na</td>
<td>Na</td>
<td>0.96 eV</td>
</tr>
<tr>
<td>2×Na</td>
<td>Na$_3$</td>
<td>1.02 eV</td>
</tr>
<tr>
<td>3×Na</td>
<td>Na$_3$</td>
<td>2.05 eV</td>
</tr>
<tr>
<td>3×Na</td>
<td>Na$_2$ + Na</td>
<td>1.47 eV</td>
</tr>
<tr>
<td>4×Na</td>
<td>Na$_3$ + Na</td>
<td>1.53 eV</td>
</tr>
<tr>
<td>4×Na</td>
<td>Na$_3$ + Na$_2$</td>
<td>1.92 eV</td>
</tr>
<tr>
<td>4×Na</td>
<td>Na$_3$ + 2×Na</td>
<td>1.98 eV</td>
</tr>
<tr>
<td>5×Na</td>
<td>Na$_3$</td>
<td>2.04 eV</td>
</tr>
<tr>
<td>5×Na</td>
<td>Na$_4$</td>
<td>2.55 eV</td>
</tr>
</tbody>
</table>

bond. For this purpose the Na$_2$ bond distance has been increased gradually up to a point where the periodic Na(II) phase is obtained. Each configuration has been optimized with respect to the surface, and the total energy is calculated with the 5×5×1 k-point mesh. Our results show a monotonic increase up to Na(II), which is the upper limit of Na-Na distance (6.51 Å) in the supershell chosen. At this point, the energy is 0.30 eV higher, which should be considered as the lower bound of the Na$_2$ dissociation energy on HOPG. This is still significantly less than the gas phase value 0.68 eV, but the substrate now causes the interaction between Na atoms to be long ranged. On the other hand, Na$_2$ stretches readily; as the Na-Na separation is increased to 4.26 Å where both atoms sit on a hollow site (second nearest hexagons) the total energy change is only 0.09 eV, but the distance $d_1$ = 3.09 Å is 0.86 Å less. This suggests that Na$_2$ on graphite has very low frequency vibrational modes in both lateral and perpendicular directions.

Charge transfer between the adsorbate and the substrate is studied in detail in the case of Na-HOPG and Na$_3$-HOPG, and the laterally averaged charge density differences ($\Delta \rho$) are presented in Fig. 5. In both cases, the oscillating profile of $\Delta \rho$ shows that the presence of adsorbate affects the whole system including the lowermost (third) graphene layer. The negative node close to the Na/Na$_3$ indicates a charge transfer to the substrate that is partially counterbalanced by the strong positive peak next to the first graphene layer (GR1, see Table V). The location and shape of the negative node is different for Na and Na$_3$: for a single atom the charge is depleted throughout the whole atomic volume causing a broad minimum in $\Delta \rho$, whereas for Na$_3$ the minimum is deeper and biased to the lower side of the cluster. Integration over this area gives values $\Delta q = -0.47$ e and $\Delta q = -0.48$ e for Na and Na$_3$, respectively. The similar $\Delta q$ values indicate that the substrate does not support more excess charge, and it explains the increased $d_\perp$ for Na$_3$, Na$_4$, Na$_5$, and (2×2) Na monolayer.

A layer-by-layer analysis of the graphite substrate in Table V shows that the charge transferred is distributed over the three layers. In comparison with the middle layer (GR2), $\Delta q$ is slightly larger for the lowermost layer (GR3). This is probably a finite-size effect, a conclusion that is supported by the $\Delta \rho$ profile. The inclusion of k points leads to more pronounced oscillations near GR2 and GR3, whereas GR1 has more accumulated charge in the $\Gamma$-point approximation. This shows that the charge transferred becomes more delocalized as k points are introduced in the lateral dimension. The $\Gamma$-point approximation underestimates the amount of charge transfer also for Na, whereas for Na$_3$ the values are similar. Lamoen and Persson found $\Delta q = -0.40$ e for a (4×4) K monolayer, which agrees with our result $\Delta q = -0.39$ for a corresponding density of Na using a single $\Gamma$ point (Table V).

| TABLE V. Charge transfer in Na-HOPG and Na$_3$-HOPG (in electrons). The values in parenthesis are for the $\Gamma$-point approximation. |
|-----------------|-----------------|-----------------|-----------------|
| Na-HOPG         | Na               | GR1             | GR2             | GR3             |
| Na              | -0.47 (−0.39)   | 0.25 (0.28)     | 0.09 (0.03)     | 0.13 (0.09)     |
| Na$_3$-HOPG     | -0.48 (−0.49)   | 0.26 (0.37)     | 0.10 (0.03)     | 0.12 (0.09)     |
The conduction band is now being filled by the Fermi level, and the characteristic features of graphite substrate are visible. The DOS of graphite substrate shows typical properties of graphite, although the system is finite in the perpendicular direction.

A very small effect is observed due to the Na₅ adsorption, and the electronic densities of valence states (DOS) of Na₅-HOPG and HOPG are plotted in Fig. 6. The calculations were done using a 5 × 5 × 1 k-point mesh, and the KS eigenvalues obtained are interpolated to correspond 9 × 9 × 1 mesh, and Gaussians of 0.10 eV width were used for each data point. The dashed line marks the Fermi level.

The electronic densities of valence states (DOS) of Na₅-HOPG and HOPG are plotted in Fig. 6. The calculations were done using a 5 × 5 × 1 Monkhorst-Pack k-point mesh, and the KS eigenvalues obtained are interpolated to correspond a 9 × 9 × 1 mesh (this resembles the common tetrahedron method). The DOS of graphite substrate shows typical features, including a steep rise at −20 eV due to the 2D character of graphite, a dip at −13 eV after the first two σ bands, a large peak at −6.5 eV followed by a shoulder in the decreasing profile with zero weight and zero gap at the Fermi energy. Our substrate model then captures the relevant properties of graphite, although the system is finite in the perpendicular direction.

A very small effect is observed due to the Na₅ adsorption, and the characteristic features of graphite substrate are clearly visible. The conduction band is now being filled by the Fermi level. The band structure of the three (spin-degenerate) Na₅-HOPG conduction states reveals that the dispersion of the first does not correspond to its graphite counterparts (π* bands), but resembles the valence states more (π bands). This is not true for the two other conduction states, where the lower one shows only minor variation as a function of k, and the higher one resembles closely the graphite conduction bands. The two uppermost valence states are also affected by the presence of Na₅, which can be seen as smaller dispersion. Together with the lowest conduction state, this results in a small hump in the DOS next to the minimum separating conduction and valence bands.

FIG. 6. Normalized DOS of Na₅-HOPG and HOPG systems calculated with the 5 × 5 × 1 k-point mesh. The data is interpolated to correspond a 9 × 9 × 1 mesh, and Gaussians of 0.10 eV width were used for each data point. The dashed line marks the Fermi level.

IV. CONCLUSION

We have studied Na atoms and small Na clusters (N = 5) on graphite using a DFT method that uses pseudopotentials and periodic boundary conditions. In order to obtain reliable results the simulated slab of graphite consists of three graphene layers, and is sufficiently large to yield an appropriate separation between the adsorbate replica in lateral dimension. In addition, a high kinetic energy cutoff (70 Ry) for the plane wave basis set and k points make the calculations very demanding in terms of CPU time and memory.

For a dispersed phase, an Na atom has an adsorption energy of 0.51 eV at the hollow site 2.45 Å above the surface. The small diffusion barrier of 0.06 eV shows that the potential energy surface of the Na atom is flat. These results are similar to the recent results for NaC₆₀ compounds, where an adsorption energy of 0.65 eV and a diffusion barrier of 0.07 eV were observed at the hexagonal site. A higher Na coverage leads to a decreased interaction with the substrate as shown for the (2 × 2) monolayer (d₃ = 3.21 Å). The dispersed phase and (2 × 2) monolayer differ little energetically, and neither is found to be stable experimentally. Instead, the calculated cluster formation energies favor clustering processes (island formation) in agreement with experiment. The formation energies of the open shell systems Na, Na₃, and Na₅ are larger than those of closed shell cases Na₂ and Na₄. This is related to the spin-degeneracy of the highest molecular orbital (odd-even staggering) and, in contrast to free metal clusters, gives rise to increased stability of odd cluster sizes on HOPG.

A charge density analysis for Na and Na₃ shows that approximately 0.5 electrons are transferred to the substrate in both cases, indicating that HOPG does not support much excess charge, and that polarization effects weaken as the Na coverage is increased. As shown before for K, this leads to decoupling between the adsorbate and substrate, and a two-dimensional metallic film on HOPG results. In the case of Na clusters, the partial loss in electron density is evident in significant changes in cluster geometries. For example, Na₃ is more similar to a closed shell Na₃ ion, and consequently, the geometry is closer to an equilateral triangle than that of a free Na. An interesting observation is that the planarity of Na₃ and Na₅ is broken as the atoms having more Na-Na bonds move farther from the surface. Whether this is related to the experimentally observed buckling of Na over-layers remains an open question.

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31 N. Troullier and J.L. Martins, Phys. Rev. B 43, 1993 (1991). For the carbon 2s22p2 valence electrons, we use s-nonlocal and p-local components (rcut = 1.23 a.u.). The sodium pseudopotential is constructed for the 3s valence electron only (rcut = 2.3 a.u.) without nonlinear core corrections.
33 A. Alavi, J. Kohanoff, M. Parrinello, and D. Frenkel, Phys. Rev. Lett. 73, 2599 (1994). The finite temperature functional used can cause small changes in the adsorbate geometries found with the zero-temperature functional.
35 Especially the distance from the surface is sensitive to the number of graphene layers. For two graphene layers the distance is 0.16 Å (6.6%) larger than for three and four layers.
36 This system corresponds to a (4×4) Na monolayer in terms of coverage.
37 The corresponding box size 12.30×12.79×13.35 Å3.
38 The value of charge transfer can depend on the method used for its determination. We use the same method as the previous studies on alkali-HOPG systems (Refs. 6–8).