Site-selective adsorption of phthalocyanine on h-BN/Rh(111) nanomesh

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I. CALCULATIONS STRUCTURAL AND ELECTRONIC PROPERTIES OF NANOMESH

A. Structural parameters

The modulation of the height of N atoms, of the height of the Rh atoms of the topmost layer of the slab, and of the BN bond length have been calculated along the [21] diagonal of the nanomesh (NM) unit cell. The three different line colors indicate the results for the three tested models, revPBE-D3 (black), PBE-rVV10 (blue), and vdW-DF (red).

![Graph showing modulation of height](image)

FIG. 1. Height profile over the diagonal crossing the nanomesh unit cell along the [21] direction: (a) N sublattice and o (b) Rh atoms of the topmost layer. (c) Modulation of the BN bond length of the atoms laying along the same line. The height of the atoms is measured with respect to the average height of the Rh atoms in the topmost layer.

B. Projected density of states

The (normalized) density of state (DOS) of the h-BN/Rh(111) nanomesh is shown in Figs. 2(a) and 2(b) for the pore and wire regions, respectively. It can be seen that the two functionals lead to DOSs which are very similar, in particular for the pore region. It has already been shown in Refs. 1–3 that the \(\sigma\)-band splitting of about 1 eV measured experimentally\(^4\) could be well reproduced by the present model of nanomesh (i.e., \(13 \times 13\) h-BN monolayer on top of \(12 \times 12\) Rh(111) surface). This shift, coming from the shift between the N-\(p\) DOSs of the pore and wire regions can also be observed in the present work.
FIG. 2. N-p partial density of states for the N atoms in (a) the pore and (b) the wire regions obtained from the revPBE-D3 and PBE-rVV10 functionals. The curves have been divided by the number of N atoms in their respective regions. The Fermi energy is set at zero.

C. Simulated STM topography
II. CALCULATIONS OF H$_2$PC IN GAS PHASE

The structure of the molecule optimized in gas phase is planar, since all C-C and C-N have sp2 character. Bond lengths and bond angles do not vary significantly by changing among the three models revPBE-D3, PBE-rVV10, and vdW-DF.
FIG. 4. H$_2$Pc structure optimized in gas phase: H cyan, C yellow, N grey.

III. CALCULATIONS OF H$_2$PC ADSORBED ON THE NANOMESH

The calculations have been restricted to the case of H2Pc, because CuPc requires spin polarized electronic structure optimizations, which makes the convergence slower and increases significantly the computational costs. Every structure optimization involves a simulation cell containing almost 1000 atoms. The system is metallic, which makes the wavefunction optimization slowly converging; typically about 50 iterations of the self consistent cycle are required. The shallow potential energy surface for the adsorption of the molecule makes it difficult to locate the minimum. A typical structure optimization procedure requires the generation of more than 100 configurations.

For all optimized adsorption configurations of H$_2$Pc on the NM, there are in principle two possible locations of the innermost H atoms. One is with the N-H bonds along the [21] direction, the other is with them perpendicular to the [21] direction. The influence of the H position to the adsorption is first evaluated by structure optimization with the innermost H atoms. One is with the N-H bonds along the [21] direction, the other is with them perpendicular to the [21] direction. The STM images were generated for the optimized structures (Fig. 2(a)). When it belongs to the benzene ring, respectively.

A. revPBE-D3 adsorption energies

The most stable configuration for the revPBE-D3 model is with H$_2$Pc centered in the pore. In order to understand

<table>
<thead>
<tr>
<th>Adsorption site</th>
<th>poreA</th>
<th>wire</th>
<th>offA1</th>
<th>offA2</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E_{ads}$</td>
<td>-3.03 (-2.75)</td>
<td>-2.31</td>
<td>-2.96</td>
<td>-2.93 (-2.93)</td>
</tr>
<tr>
<td>$E_{DFT}$</td>
<td>2.35</td>
<td>2.52</td>
<td>2.72</td>
<td>2.36</td>
</tr>
<tr>
<td>$E_{disp}$</td>
<td>-5.37</td>
<td>-4.83</td>
<td>-5.68</td>
<td>-5.29</td>
</tr>
<tr>
<td>$E_{nm}$</td>
<td>0.077</td>
<td>0.051</td>
<td>0.224</td>
<td>0.053</td>
</tr>
<tr>
<td>$E_{pc}$</td>
<td>0.015</td>
<td>0.017</td>
<td>0.021</td>
<td>0.028</td>
</tr>
</tbody>
</table>

* The geometry of the nanomesh was kept fixed at the PBE-rVV10 one. The H$_2$Pc molecule was allowed to relax.

why the preferred adsorption site with revPBE-D3 is the center of the pore while with PBE-rVV10 is off-center, we compare how the DFT and dispersion contributions change. With revPBE-D3, $E_{DFT}$ is 0.37 eV less repulsive for poreA than for offA1, while $E_{disp}$ is only 0.31 eV less attractive. The different contributions almost compensate each other, which makes a 0.07 eV lower adsorption energy for poreA. On the other hand, with PBE-rVV10, the change in the repulsive DFT term between poreA and offA1 is only 0.046 eV in favor of the former, while the gain in dispersion of offA1 is of 0.227 eV. This can be explained with the fact that at the center of the larger PBE-rVV10...
NM pore, the molecule minimize the interaction with the atoms of the rim, and all its atoms are equally distant from the closest BN pairs. Moving the molecule towards the rim there is enough space to let the attractive dispersion interaction to increase, between one lobe of the molecule and the rim, still keeping the repulsive term small. With the revPBE-D3, the molecule is almost as large as the NM pore, and already when centered in the pore, some of its atoms are already quite close to the rim. Pushing the molecule further towards the rim brings one lobe almost beyond the rim, thus increasing the attraction term between C atoms and BN pairs, but also inducing an equal increase of the repulsive contribution. In order to confirm the implications of the pore shape, H$_2$Pc has been optimized with revPBE-D3 on the fixed NM geometry as obtained with PBE-rVV10. Similarly, we have optimized H$_2$Pc with PBE-rVV10 on the fixed revPBE-D3 NM. It turns out that with the PBE-rVV10 NM shape the off center adsorption is always favorite, irrespective of the functional used to calculate the interaction.

![Diagram](image)

**FIG. 5.** Top view of Pc adsorbed on the NM in poreA and offA1 configurations as obtained with the revPBE-D3 functional (top) and with the PBE-rVV10 functional (bottom). The atoms of the h-BN lattice are colored according to their height over the Rh surface: Blue $d_z < 2.4$ (pore), yellow $d_z > 2.4$ red $d_z > 3.7$. The atoms of H$_2$Pc are all magenta.

With the vdW-DF functional the adsorption energy of H$_2$Pc in the does not change significantly depending on the specific site. It is always about 3.06 eV. Due to the choice of the functional and the less corrugated h-BN layer, the distance between BN and molecule is larger and does not change significantly by moving the molecule over the pore. Hence, the two terms of the interaction energy change also little.

**B. Electronic structure of adsorbed H$_2$Pc**

The charge density difference $\Delta \rho(r)$ is calculated as

$$\Delta \rho(r) = \rho_{pc/NM}(r) - \rho_{pc}(r) - \rho_{NM}(r),$$  \hspace{1cm} (1)

where $\rho_{pc}(r)$ and $\rho_{NM}(r)$ are the charge densities of the two subsystems taken alone, but in the same geometry of the complex. These volumetric data structures represent the changes in the electronic charge distribution due to the interaction between molecule and substrate. The rather small $\Delta \rho(r)$ computed for all pc/NM configurations confirm that no chemical bonding is formed. However, some polarization effects are present, as it can be observed in the
FIG. 6. Density difference plots: side view (top) and top view (bottom) of the Δρ isosurface at +0.007 el/Å³ (red) and -0.007 el/Å³ (green) calculated for three optimized structures, poreA, offA1, and offB1. The Pc molecule and the h-BN layer are represented by lines along the bonds. C are brown, H white, N blue and B orange. The Rh substrate is represented by grey balls. Each panel displays only a portion of the NM unit cell.

plots of Δρ(r) isosurfaces displayed in figure 6 for three configurations, poreA, offA1 and offB1. The red isosurface corresponds to accumulation of charge, and it is mainly located in the area below the pyrrole-like subunits, indicating polarization of the molecular charge towards the substrate. The corresponding charge depletion is illustrated by the green isosurface and it is localized close to atomic centers of the molecule (mainly H and N) and at some N and B of the NM pore underneath the molecule. The electron distribution at the metal, instead, is barely affected by the presence of the molecule.

The dipole moment of the H₂Pc molecule in gas phase and adsorbed on the surface has been calculated using the method of the maximally localized Wannier function (MLWF)\(^2\). Once the localization procedure has converged, the MLWF are associated either to the molecule or to the substrate. The center of the MLWF, i.e. the Wannier center, represents the position of the localized electron pairs, charge -2. The atomic positions are insted the location of the positive core charges, i.e. +5 for N, +4 for C, and +1 for H. The dipole moment is then obtained from the sum over all the particles, ionic cores and Wannier centers, of the product of the position vector times the corresponding charge

\[ \mu = \sum_{i=1}^{N_{\text{atom}}} R_i q_i - 2 \sum_{x=1}^{N_{\text{wc}}} R_x \]  

(2)

In gas phase, all the three Cartesian components of the dipole are null. The interaction with the substrate, instead, induces a molecular dipole, with the largest component in the direction perpendicular to the surface, \( \mu_z \).

We have computed the projected density of states (PDOS) for H₂Pc in gas phase and adsorbed in the offA1 configuration. In figure 7 (a) and (b), the black curves are the PDOS on C atoms and N atoms, respectively, where the energies are reported with respect to the center of the homo-lumo gap. The isosurfaces correspond to the wavefunctions of the homo-1, homo, lumo, and lumo+1 states are displayed in panel (c). The same PDOS computed for the molecule adsorbed in the offA1 configuration are reported in red. In this case the energies are given with respect to the Fermi energy. It is noticed that the lumo of H₂Pc is almost aligned with the Fermi energy of the

<table>
<thead>
<tr>
<th>Adsorption site</th>
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<th>wire</th>
<th>offA1</th>
<th>offB1</th>
</tr>
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<tbody>
<tr>
<td>( \mu_x )</td>
<td>1.526</td>
<td>2.151</td>
<td>0.944</td>
<td>0.044</td>
</tr>
<tr>
<td>( \mu_y )</td>
<td>-2.71</td>
<td>-1.039</td>
<td>1.339</td>
<td>3.340</td>
</tr>
<tr>
<td>( \mu_z )</td>
<td>5.105</td>
<td>1.298</td>
<td>5.365</td>
<td>5.387</td>
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<tr>
<td>( \mu )</td>
<td>5.336</td>
<td>2.719</td>
<td>5.610</td>
<td>6.338</td>
</tr>
</tbody>
</table>

TABLE II. Dipole moment components and norm in Debye
complex. Otherwise, the structure of the molecular orbitals seems not to be significantly perturbed by the interaction with the substrate.

FIG. 7. a) and b) projected density of states on C and N atoms of H$_2$Pc, respectively. The black curve is the PDOS computed for the molecule optimized in gas phase, the red curve for the molecule adsorbed in the offA1 configuration.
FIG. 8. Sequence of constant current STM images at different biases showing the energy variations in the HOMO-LUMO positions of H$_2$PC on the h-BN/Rh(111) NM. All images have been measured at 26 pA set point current. The yellow circle denotes the same position in all images.
C. Mobility within the pore

At higher temperature the molecules cannot be imaged stably anymore, but that they are pushed around in the pore during the STM scanning. The “double cross” configuration seen in the figure originates from the bi-stable position of the molecule in the pore and not from two molecules sitting in the same pore. This later situation is realized at higher molecule coverage as can be seen in the figure 8, where the two molecules in the pore are well resolved. Depending on the tip condition and tunneling parameters the images at 77 K can become very blurred due to the mobility of the molecules in the pore. Despite the mobility the molecules are not pushed out of the pores.

![Figure 9](image)

FIG. 9. a) Constant current STM image of CuPc on the h-BN/Rh(111) NM measured at 77 K where a single molecule per pore is imaged switching between two off-center positions. The image is rotated 45° with regard to the scan direction, i.e. the direction of the molecule displacement is not in the direction of the scan. b) DFT minimum energy configurations of H$_2$Pc on opposite off-center positions.