Electronic and Structural Properties of Carbon Nano-Horns

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We use parametrized linear combination of atomic orbitals calculations to determine the stability, optimum geometry and electronic properties of nanometer-sized capped graphitic cones, called "nano-horns". Different nano-horn morphologies are considered, which differ in the relative location of the five terminating pentagons. Simulated scanning tunneling microscopy images of the various structures at different bias voltages reflect a net electron transfer towards the pentagon vertex sites. We find that the local density of states at the tip, observable by scanning tunneling spectroscopy, can be used to discriminate between different tip structures. Our molecular dynamics simulations indicate that disintegration of nano-horns at high temperatures starts in the highest-strain region near the tip.

61.48.+c, 61.50.Ah, 68.70.+w, 73.61.Wp

Since their first discovery, carbon nanotubes have drawn the attention of both scientists and engineers due to the large number of interesting new phenomena they exhibit, and due to their potential use in nanoscale devices: quantum wires, nonlinear electronic elements, transistors, molecular memory devices, and electron field emitters. Even though nanotubes have not yet found commercially viable applications, projections indicate that this should occur in the very near future, with the advent of molecular electronics and further miniaturization of micro-electromechanical devices (MEMS). Among the most unique features of nanotubes are their electronic properties. It has been predicted that single-wall carbon nanotubes can be either metallic or semiconducting, depending on their diameter and chirality. Recently, the correlation between the chirality and conducting behavior of nanotubes has been confirmed by high-resolution scanning tunneling microscopy (STM) studies.

Even though these studies have demonstrated that atomic resolution can be achieved, the precise determination of the atomic configuration, characterized by the chiral vector, diameter, distortion, and position of atomic defects, is still a very difficult task to achieve in nanotubes. Much of the difficulty arises from the fact that the electronic states at the Fermi level are only indirectly related to the atomic positions. Theoretical modeling of STM images has been found crucial to correctly interpret experimental data for graphite and has been recently applied to carbon nanotubes. As an alternative technique, scanning tunneling spectroscopy combined with modeling has been used to investigate the effect of the terminating cap on the electronic structure of nanotubes.

Among the more unusual systems that have been synthesized in the past few years are cone-shaped graphitic carbon structures. Whereas similar structures have been observed previously near the end of multi-wall nanotubes, it is only recently that an unusually high production rate of up to 10 g/h has been achieved for single-walled cone-shaped structures, called "nano-horns", using the CO2 laser ablation technique at room temperature in absence of a metal catalyst. These conical nano-horns have the unique opening angle of ≈20°.

We consider a microscopic understanding of the electronic and structural properties of nano-horns a crucial prerequisite for understanding the role of terminating caps in the physical behavior of contacts between nanotube-based nano-devices. So far, neither nano-horns nor other cone-shaped structures have been investigated theoretically. In the following, we study the structural stability of the various tip morphologies, and the inter-relationship between the atomic arrangement and the electronic structure at the terminating cap, as well as the disintegration behavior of nano-horns at high temperatures.

Cones can be formed by cutting a wedge from planar graphite and connecting the exposed edges in a seamless manner. The opening angle of the wedge, called the disclination angle, is n(π/3), with 0 ≤ n ≤ 6. This disclination angle is related to the opening angle of the cone by θ = 2 sin−1(1 − n/6). Two-dimensional planar structures (e.g. a graphene sheet) are associated with n = 0, and one-dimensional cylindrical structures, such as the nanotubes, are described by n = 6. All other possible graphitic cone structures with 0 < n < 6 have been observed in a sample generated by pyrolysis of hydrocarbons. According to Euler’s rule, the terminating cap of a cone with the disclination angle n(π/3) contains n pentagon(s) that substitute for the hexagonal rings of planar graphite.

The observed cone opening angle of ≈20°, corresponding to a 5π/3 disclination, implies that all nano-horns contain exactly five pentagons near the tip. We classify the structure of nano-horns by distinguishing the relative positions of the carbon pentagons at the apex which determine the morphology of the terminating cap. Our study will focus on the influence of the relative position...
We associate the tip region of a hypothetically infinite nano-horn with all the sites excluding the edge. Structural details and the results of our stability calculations are presented in Table I. These results indicate that atoms in nano-horns are only ≈0.1 eV less stable than in graphite. The relative differences in $<E_{\text{coh, tot}}>$ reflect the strain energy changes induced by the different pentagon arrangements. To minimize the effect of under-coordinated atoms at the edge on the relative stabilities, we excluded the edge atoms from the average when calculating $<E_{\text{coh, tip}}>$. Since our results for $<E_{\text{coh, tip}}>$ and $<E_{\text{coh, tot}}>$ follow the same trends, we believe that the effect of edge atoms on the physical properties can be neglected for structures containing hundreds of atoms. Even though the energy differences may appear minute on a per-atom basis, they translate into few electron-volts when related to the entire structure. Our results suggest that the under-coordinated edge atoms are all less stable than the cone mantle atoms by ≈0.5 eV. Also atoms in pentagons are less stable than those in hexagons by ≈0.1 eV, resulting in an energy penalty of ≈0.5 eV to create a tetragone if the strain energy induced by bending the lattice could be ignored.

When comparing the stabilities of the tip regions, described by $<E_{\text{coh, tip}}>$, we found no large difference between blunt tips that have all the pentagons distributed along the cylinder mantle and pointed tips containing a pentagon at the apex. We found the structure shown in Fig. (c) to be more stable than the other blunt structures with no pentagon at the apex. Similarly, the structure shown in Fig. (e) is most stable among the pointed tips containing a pentagon at the apex. Equilibrium carbon-carbon bond lengths in the cap region are $d_{\text{CC}} = 1.43 - 1.44$ Å at the pentagonal sites and $d_{\text{CC}} = 1.39$ Å at the hexagonal sites, as compared to $d_{\text{CC}} = 1.41 - 1.42$ Å in the mantle. This implies that the “single bonds” found in pentagons should be weaker than the “double bonds” connecting hexagonal sites, thus con-
the apex and a relatively blunt tip.

The most marked differences are noted near the Fermi level, at $E \approx 0$ eV.

Figure 2 shows the charge density associated with occupied states near the Fermi level, corresponding to the bias voltage of $V_b = 0.2$ V, are suggestive of a net electron transfer from the hexagonal to the pentagonal sites. The charge density contour displayed corresponds to the value of $\rho = 1.35 \times 10^{-3}$ electrons/Å$^3$. Dark lines depict the atomic bonds to guide the eye.

In Fig. 2, we present such simulated STM images for the nano-horns represented in Figs. (c) and (d). We show the charge density associated with occupied states within a narrow energy interval of 0.2 eV below the Fermi level as three-dimensional charge density contours, for the density value of $\rho = 1.35 \times 10^{-3}$ electrons/Å$^3$. Very similar results to those presented in Fig. 2 were obtained at a higher bias voltage of 0.4 eV. As seen in Fig. 2, $pp\pi$ interactions dominate the spectrum near $E_F$. These images also show a net excess charge on the pentagonal sites as compared to the hexagonal sites. This extra negative charge at the apex should make pointed nano-horn structures with a pentagon at the apex better candidates for field emitters than structures with no pentagon at the apex and a relatively blunt tip.

It has been shown previously that theoretical modeling of STM images is essential for the correct interpretation of experimental data. Atomically resolved STM images, however, are very hard to obtain especially near the terminating caps of tubes and cones, due to the large surface curvature that can not be probed efficiently using current cone-shaped STM tips. A better way to identify the tip structure may consist of scanning tunneling spectroscopy (STS) measurements in the vicinity of the tip. This approach is based on the fact that in STS experiments, the normalized conductance $(dI/dV)/(V/I)$ is proportional to the local density of states which, in turn, is structure sensitive. We have calculated the local density of states at the terminating cap of the tip for the different nano-horn structures shown in Fig. 1. Our results are shown in Fig. 3 convoluted using a Gaussian with a full-width at half-maximum of 0.3 eV.

To investigate the effect of pentagonal sites on the electronic structure at the tip, we first calculated the local density of states only at the 25 atoms contained in the five terminating pentagons. The corresponding densities of states, shown by the dashed curve in Fig. 3, are found to vary significantly from structure to structure near the Fermi level. Thus, a comparison between the densities of states at $E \approx E_F$ should offer a new way to discriminate between the various tip morphologies. For an easy comparison with experiments, we also calculated the local density of states in the entire terminating cap, including all five pentagons and consisting of $N \approx 40 \rightarrow 50$ atoms,
depending on the structure. The corresponding density of states, given by the solid line in Fig. 3 is vertically displaced for easier comparison. Our results show that the densities of states, both normalized per atom, are very similar. Thus, we conclude that the pentagonal sites determine all essential features of the electronic structure near the Fermi level at the tip.

Next, we have studied the heat resilience as well as the decay mechanism of nano-horns at extremely high temperatures using molecular dynamics simulations.

In our canonical molecular dynamics simulations, we keep the structure at a constant temperature using a Nosé-Hoover thermostat and use a fifth-order Runge–Kutta interpolation scheme to integrate the equations of motion, with a time step of $\Delta t = 5 \times 10^{-16}$ s. We found the system to remain structurally intact within the temperature range from $T = 2,000 - 4,000$ K. Then, we heated up the system gradually from $T = 4,000$ K to $5,000$ K within 4,000 time steps, corresponding to a time interval of 2 ps. Our molecular simulations show that nano-horn structures are extremely heat resilient up to $T \leq 4,500$ K. At higher temperatures, we find these structures to disintegrate preferentially in the vicinity of the pentagon sites. A simultaneous disintegration of the nano-horn structures at the exposed edge, which also occurs in our simulations, is ignored as an artifact of finite-size systems. The preferential disintegration in the higher strain region near the pentagon sites, associated with a large local curvature, is one reason for the observation that nano-horn tips are opened easily at high temperatures, in presence of oxygen.

In summary, we used parametrized linear combination of atomic orbitals calculations to determine the stability, optimum geometry and electronic properties of nanometer-sized capped graphitic cones, called nano-horns. We considered different nano-horn morphologies that differ in the relative location of the five terminating pentagons. We found a net electron transfer to the pentagonal sites of the cap. This extra charge is seen in simulated scanning tunneling microscopy images of the various structures at different bias voltages. We found that the local density of states at the tip, observable by scanning tunneling spectroscopy, can be used to discriminate between different tip structures. Our molecular dynamics simulations indicate that disintegration of nano-horns at high temperatures starts in the highest-strain region near the tip.

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40 The nano-horn structures considered in our calculation are clusters containing edge atoms. We found that due to the charge transfer towards the under-coordinated atoms at the edge, the “Fermi level” of these clusters may drop by as much as ≈1.0 – 1.2 eV with respect to that of a graphene monolayer. To compensate for this artifact, we excluded the edge region from the calculated density of states and found the $E_F$ value to lie very close to that of a graphene monolayer.
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