Nanotube-Substrate Interactions: Distinguishing Carbon Nanotubes by the Helical Angle

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We investigate the interaction of a carbon nanotube with a graphite substrate, using an interlayer potential that explicitly treats the registry dependence of the interaction. The carbon-carbon bond lengths in nanotubes differ slightly from those in flat graphite, so that the naively commensurate angular orientations for the tube with respect to the substrate lattice are destroyed. The interaction of a one-dimensional tube with a two-dimensional substrate then leads to an unusual registry phenomenon not visible in standard layer-on-layer growth: the system develops favorable orientations which clearly are incommensurate.

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Carbon nanotubes [1] are gaining prominence as potential components of future nanoelectronic and nanomechanical systems [2]. The tubes’ exceptional electronic [3] and mechanical [4] properties can be very sensitive functions of the wrapping indices \((n, m)\), which define the tube’s circumference in graphitic lattice coordinates \([an (n,m) tube wraps at a helical angle of \(\theta_0 = \tan^{-1}(\sqrt{3}m/(2n + m))\) to the hexagonal rows of graphite [5]). Postsynthesis separation into metal and semiconducting components already has been demonstrated [6,7] to complement the reasonably good control of the average diameter for as-grown tubes [8–11]. Separations based on electronic response are attractive for electronics applications, since they target the relevant materials property. However, a wider range of applications might be opened up if tubes could be separated more precisely, e.g., by their helical pitch. This structural property, which reaches the fundamental \((n,m)\) distinction, is finer grained than the metal/semiconductor distinction. Here we describe the physical basis of one potential means for separating nanotubes according to their helical pitch.

Experimental observations of spontaneous nanotube alignment on graphite suggest that nanotubes have a registry-dependent interaction with an underlying graphitic lattice [12]. Preliminary calculations [13] show sharp peaks every \(60^\circ\) in the tube/graphite binding as a function of angle \(\theta\) measured about an axis perpendicular to the substrate. Here we refine these important preliminary results and demonstrate that the energy selectivity is 5 to 10 times stronger than previously suspected. In addition, we demonstrate that the preferred tube-graphite orientations are not limited to commensurations. Surprisingly, many incommensurate orientations are also favorable, even in the realistic case of slightly different C-C bond lengths for the tube and substrate. These unexpected incommensurate lock-in angles arise from finite-size effects perpendicular to the tube axis, and are unique to one-dimensional objects interacting with two-dimensional substrates.

Nanotube alignment on a graphite substrate is sensitive to the variation of the tube-substrate binding energy under shifts in registry between graphene layers. Thus, a careful choice of the interaction potential is required. A standard Lennard-Jones potential is valuable for averaged treatments of the overall binding [14] and for qualitative indications of registration-dependent effects. However, this potential underestimates the corrugation in the interlayer interaction. An improved interlayer potential that treats explicitly the short-ranged (e.g., orbital overlap) corrugation effects has been proposed previously [15]. This potential was trained on data for sliding between two flat and parallel graphene layers. Recent first-principles calculations suggest that this preliminary potential tends to overestimate the corrugation of nanotube/nanotube and graphite/nanotube sliding by a factor of roughly 2. We have since refined the potential and reduced the overestimation to about 50% [16].

For simplicity, we consider first a rigid nanotube on a flat graphitic surface. The tube center of mass has three translational degrees of freedom. The tube can also spin about its axis. We assume that the tube axis is parallel to the substrate, which eliminates one rotational degree of freedom. The remaining angular degree of freedom, namely, the angle about an axis perpendicular to the substrate, is the main variable of interest. We hold this angle fixed at a succession of values (every \(0.1^\circ\)) while optimizing the other parameters. For a fully incommensurate system, shifts of the center of mass parallel to the plane of the substrate do not change the energy; otherwise a careful minimization is required. To reduce finite-size fluctuations, the tubes are very long (about 300 nm). We treat the substrate as a single rigid graphene layer. The...
deeper layers are too far away to substantially affect the registry dependence. However, by pulling the nanotube closer to the surface they could indirectly increase most of the corrugation effects described here by about 10%. Subtle long-ranged effects (e.g., due to scattering by the tube of graphene electrons in low energy states) also might affect the orientation dependence of the tube-substrate binding energy. Such effects, which are not treated in the short-ranged potential used here, might be suppressed by using a substrate such as hexagonal B-N.

We begin with the case where all nanotube C-C bonds have the same length as in bulk graphite; this provides a helpful starting point for the later inclusion of more subtle effects. Figure 1 shows the angular dependence of the tube-substrate interaction energy for three nanotubes with similar diameters but different helical angles. As expected, all tubes have a sharp minimum precisely at \( \theta_0 \), repeating every 60° [13]. The gain in the binding energy at the preferred orientation is substantial, about 8% of the total binding energy and tens of electron volts for tubes of a few hundred nanometers in length.

The curve between the principal minima is not quite flat. Some of the features are known accidental commensurations for graphite [17]. For example, two graphitic layers become commensurate at 38.2° \( \cos^{-1}(11/14) \). However, the highly symmetric (10, 0) and (5, 5) tubes have additional minima at incommensurate angles, e.g., 30°. These unexpected features can be explained by looking at patterns in the distribution of local interlayer registries.

In commensurate systems, the points cluster into a finite set whose energetics can be optimized through center of mass motion of the nanotube relative to the substrate or spinning about the tube axis. In normal incommensurate systems, the points cover the entire unit cell uniformly, yielding a smooth interaction. However, the finite transverse extent of the nanotube allows a third possibility, which is seen here: the points can spread uniformly, but only over a discrete set of parallel lines that slice through the substrate unit cell.

In this third case, even though the system is axially incommensurate, the energy can be lowered by spinning the tube about its axis and/or shifting its center of mass parallel to the substrate and perpendicular to the tube axis. The effect is largest for the high-symmetry zigzag nanotubes with short axial unit cells, at angles \( \theta = 30.0° \) and 41.4° \( \cos^{-1}(3/4) \). This case is absent in normal two-dimensional thin-film growth, since it is a nanoscale finite-size effect. The system is partway between the commensurate and incommensurate states, so we term it a “pseudocommensuration.” The effect is purely geometric. It would appear also in a Lennard-Jones treatment or for a hexagonal nongraphitic (or doped graphite) substrate.

The separation of the primary minima for the (10, 0) tube of Fig. 1 into a broad valley and a sharp spike results in matching bond lengths in the nanotube and graphene substrate. The distributions of interlayer registries for the minima marked “[2a]” and “[2b]” (at 30.0° and 38.2°, respectively) are shown in Fig. 2.
from similar physics. The sharp minimum occurs as the set of projected registries condenses from lines into isolated points. The broader valley follows the extension of these lines across the unit cell and their subsequent smearing into a uniform and fully incommensurate coverage. The angular widths $\Delta \theta$ of the primary minima depend on the tube length as $1/\sqrt{L}$. For zigzag tubes $\Delta \theta = 1/\sqrt{N}$, where $N$ is the number of axial unit cells.

Careful ab initio studies [18] show that the C-C bonds, lengthen slightly when graphene is rolled into a nanotube. For tubes of the diameter studied here, the mismatch is about 0.40%, meaning that even for the perfect angular alignment at $\theta_0$, registry is destroyed over about 40 nm (barring axial relaxation of the tube, to be discussed later). Do any substantial commensurate valleys survive this bond-length mismatch in long tubes? To answer this question, we treat next the case of mismatched bonds. Bonds parallel to the tube axis ($a_\parallel$) are extended by 0.35%. Bonds perpendicular ($a_\perp$) are extended by 0.45% [18]. The other bonds follow a linear interpolation. Strikingly, rather strong valleys survive, as shown in Fig. 3. The primary peaks shrink by less than 20% (except for the armchair tube) compared to the ideal case of Fig. 1. Several new minima appear, none of which is a standard commensuration: for mismatched bonds, essentially all preferred angles result from pseudocommensurate sampling. For example, in the (10,0) tube minima are now at 0.0°, 0.16°, and 4.8°. The second angle solves the equation $\sqrt{3} = (a_\parallel \cos \theta - a_{\text{graphene}})/(a_\parallel \sin \theta)$, which yields $\theta = \sin^{-1}[(a_\parallel - a_{\text{graphene}})/\sqrt{3}a_\parallel]$ for small $\theta$s. The 4.8° valley occurs when the projected parallel bond in the nanotube matches the substrate: $\theta = \cos^{-1}(a_{\text{graphene}}/a_\parallel)$. For chiral tubes this happens when the angle $\theta$ satisfies $a_\parallel \cos \theta + \theta_0 = a_{\text{graphene}} \cos \theta_0$.

Figure 4 summarizes the typical (i.e., plateau) and maximal binding energies for several tubes. The displacement between the maximal and typical binding curves, even in the realistic mismatched-bond-length case, suggests that an ensemble of tubes could be distinguished, or perhaps even separated, based on their angle-dependent interactions with a substrate of hexagonal symmetry [19]. Such a separation would have to satisfy at least three requirements. First, the nanotube free energy within an ambient fluid [20] must be comparable to the average tube-substrate interaction to facilitate equilibration. Second, the tube sample must have a fairly narrow diameter distribution [8–11]. Third, a preferred orientation must be imposed on the nanotube population, for example, by prealigning them through extensional fluid flow or external fields [21] or by patterning the substrate [22]. Figure 4(b) demonstrates that, e.g., (8, 4) tubes could be separated out if a graphite substrate is oriented correctly to a heterogeneous mixture of aligned tubes.

Although the C-C bond is extremely stiff, deformations due to the tube-substrate interaction are possible. Thus, we investigate the role of tube deformations using the Tersoff-Brenner potential [23] to relax the tubes directly to a heterogeneous mixture of aligned tubes.

![FIG. 3. Same as Fig. 1, but the C-C bonds in the nanotube are extended by 0.35% along the axis and 0.45% perpendicular to it, as described in the text. The samplings for [2c] and [2d] are shown in Figs. 2(c) and 2(d).](image)

![FIG. 4. (a) Average (circles) and maximum (squares) interaction energy as a function of nanotube radius for nanotubes of different wrapping indices [from left to right: (5, 5), (6, 4), (7, 3), (8, 2), (9, 1), (10, 0), (6, 6), (7, 5), (8, 4), (9, 3), (10, 2), (11, 1), (12, 0)]. The hollow (solid) points are for matching (mismatched) bond lengths, as described in the text. (b) Tube-substrate interaction energy for nanotubes of various helicities and nearly constant radii (4.1 to 4.3 Å); the nanotube bonds are mismatched. From left to right: (11, 0), (10, 1), (9, 3), (8, 4), (7, 5), (6, 6), (5, 7), (4, 8), (3, 9), (1, 10). All tubes make a 19.7° angle with the substrate, which was chosen to favor the (8, 4) tube. The (4, 8) tube is not favored, since the combined system lacks inversion symmetry.](image)
systems described above. The attraction between tube and substrate flattens the tube slightly to increase the contact area. However, even for a relatively large (10, 10) tube, the tube-substrate binding energy increases by less than 10%, and the angular dependence of the binding energy shows no substantial changes. The tube also can relax along its axis, as in the Frenkel-Kontorowa model [24]. A sufficient axial compression could restore equal bond lengths in tube and substrate. Compressing an isolated small-diameter nanotube from equilibrium to a length corresponding to the C-C bond length of flat graphene costs about 0.3 meV per atom [18], which is comparable to the difference in the tube-substrate interaction energy for matched and unmatched bonds (0.1–0.2 meV per atom).

Since these energies are comparable, the nanotubes could deform along their axis and lock into commensurations with the substrate. To study axial relaxation, we simulated a 10 nm long nanotube with free hydrogenated ends on a graphene sheet. The nanotube typically does not escape the local minimum, meaning that (1) a slightly compressed bond-matched tube at the main commensurate angle remains compressed due to interactions with the substrate [25], and (2) a bond-mismatched (i.e., equilibrium length) tube at a pseudocommensurate angle retains mismatched bonds. The energies of states (1) and (2) for the same tube are nearly indistinguishable, except for the armchair tubes, where the commensurate state is more favorable.

To summarize, we describe a new type of registry-dependent interaction for a one-dimensional system on a two-dimensional substrate: certain tube/substrate orientations are preferred due to finite-size effects, even though they are manifestly incommensurate. We also identify a possible mechanism to distinguish carbon nanotubes by helical angle, which could improve our ability to produce samples with precisely defined structure and properties.

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[19] The (n, n) tubes have unexpectedly large binding energies for matching bond lengths. In a narrow (n, n) tube, a central axial row of atoms forming the line of contact with the substrate can be placed in the most energetically favorable position: above the hexagonal centers in the graphite substrate (as in AB stacked graphite). Neighboring rows of atoms, which are positioned in energetically less favorable positions directly above atoms in the substrate, curve away from the substrate, so they have less important interactions. If the bonds on the (n, n) nanotube are mismatched to the substrate, then this effect disappears, since atoms on the nanotube are uniformly spread along the substrate unit cell. This effect is therefore restricted to short and narrow armchair nanotubes.
[25] The (10, 0) tube does extend slightly, since the commensuration turns into a pseudocommensuration under axial extension for this special case.