Persistence Length Control of the Polyelectrolyte Layer-by-Layer Self-Assembly on Carbon Nanotubes

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July 29, 2005

Journal of the American Chemical Society
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One-dimensional inorganic materials such as carbon nanotubes\(^1\) and semiconductor nanowires\(^2,3\) have been central to important advances in materials science in the last decade. Unique mechanical and electronic properties of these molecular-scale wires enabled a variety of applications ranging from novel composite materials,\(^4\) to electronic circuits,\(^5\) to new sensors.\(^6\) Often, these applications require non-covalent modification of carbon nanotubes with organic compounds,\(^7\) DNA and biomolecules,\(^8\) and polymers\(^9\) to change nanotube properties or to add new functionality. We recently demonstrated a versatile and flexible strategy for non-covalent modification of carbon nanotubes using layer-by-layer self-assembly of polyelectrolytes.\(^10\) Researchers used this technique extensively for modification of flat surfaces,\(^11\) micro-,\(^12\) and nano-particles;\(^13-15\) however, little is known about the mechanism and the factors influencing layer-by-layer self-assembly in one-dimensional nanostructures.

The exact conformation of polyelectrolyte chains deposited on single-walled carbon nanotubes (SWNT) is still unknown. There are two possible configurations: flexible polymers wrapping around the nanotube and stretched, rigid chains stacked parallel to the nanotube axis.\(^7\) Several factors, such as polymer rigidity, surface curvature, and polymer-stretching interactions, can determine the nature of assembly.\(^16-19\) Persistence length of the polymer chain should be one of the critical parameters, since it determines the chain’s ability to wrap around the nanotube. Indeed, computer simulations for spherical substrates show that polymer rigidity and substrate surface curvature can influence the deposition process.\(^16,17\) Computational models also show that the persistence length of the polymer must fall below the threshold values determined by target surface curvature in order to initiate polyelectrolyte deposition process.\(^20\) Although these models described the effects of salt concentration and target surface curvature,\(^16,17\) they considered only nano-particles with radius 5 nanometer and larger.\(^18,19\) One-dimensional materials, such as carbon nanotubes, provide an even more interesting template for studying self-assembly mechanisms, since they give us access to even smaller surface curvatures down to 1 nm.

We have examined the role of the polymer persistence length in layer-by-layer self-assembly process on carbon nanotubes by observing formation of multilayer polyelectrolyte shells around carbon nanotubes at different ionic strength. Persistence length of polyelectrolytes varies with solution ionic strength,\(^21\) due to screening of the electrostatic repulsion between the polymer backbone charges,\(^22-25\) therefore changing ionic strength is a convenient way to alter the configuration of the polymer molecule systematically.

We have used the layer-by-layer self-assembly technique to form 5-layer thick coating of the alternating polyallylamine hydrochloride (PAH) and sodium poly(styrenesulfonate) (PSS) layers on the surfaces of the pristine single-wall carbon nanotubes. For our experiments, we grew the nanotubes across copper TEM grid openings using catalytic chemical vapor deposition. The deposition solutions contained different amounts of NaCl to vary the ionic strength. After polymer multilayer formation we examined the resulting coating in high-resolution TEM.

Figure 1 shows representative TEM images of coated nanotubes. Notably, we observed no continuous coatings on carbon nanotubes at low ionic strength (Figure 1, A-C). For ion concentrations of 0.05 M to 0.2 M, our samples contained more than 90% of bare uncoated single-walled carbon nanotubes; no continuous coating could be found on the randomly selected SWCNTs on the TEM grid. Once the ionic strength became greater than 0.2 M, the polyelectrolytes formed a coating on about 85% of the nanotubes with the thickness steadily increasing with the ion concentration (Figure 2). Researchers have previously...
Figure 2. Average measured polymer thickness per layer of polyelectrolyte (■) on carbon nanotube surfaces as a function of electrolyte concentration. Experimental data for the same polymer pair (PAH/PSS) on flat quartz surfaces (○) are shown for comparison. Inset: Experimental data for PSS persistence length (L_p) as a function of electrolyte concentration (△,▽). Dashed lines indicate the upper and lower boundary for the half-circumference of carbon nanotubes used in our experiments. Shaded area corresponds to the electrolyte concentration region where persistence length crosses these boundaries.

described an increase of the polyelectrolyte layer thickness with the ionic strength for layer-by-layer assembly on flat surfaces, however, they have not reported an ionic strength threshold for the deposition to initiate.

Our results can be easily rationalized if we assume that the self-assembly mechanism involves wrapping of the polymer chains around the nanotubes. At low ionic strength the polymer chains are too rigid to bend around the nanotube and the deposition cannot occur. Once the ionic strength is high enough to diminish the persistence length below the nanotube curvature, the self-assembly can proceed and the polymer layers begin to form. In contrast to the individual single-wall nanotube behavior, larger diameter single-walled nanotube bundles were usually coated by the polymers at all ionic strengths used in our studies (images not shown), further supporting our hypothesis.

We also note that we have previously reported polymer multilayer formation on carbon nanotubes at ionic strength below the deposition threshold. Unlike the current work those experiments utilized nanotube surface pretreatment with ionic pyrene derivatives to create surface charge on the nanotube. In addition, in the present experiments we have always found a small number of coated nanotubes even at low ionic strengths. These nanotubes, as well as small bundles or two or three SWCNTs, which are coated at any ionic strength, could have also contributed to the previously reported result. To examine the link between chain conformation and polyelectrolyte layer-by-layer self-assembly on carbon nanotubes we have compared the average coating thickness obtained in our experiments with the literature data for identical PAH/PSS polymer multilayers deposited on the flat surface. This comparison (Figure 2) shows that at all ionic strengths polyelectrolytes form thinner layers on the nanotube and that the coating thickness on the nanotubes begins to approach that on flat surfaces only at the high ionic strength of 1M.

A comparison between the polymer persistence length data at different ionic strengths and the size of our nanotubes provides further evidence for the wrapping mechanism of polymer self-assembly. Measured nanotube diameters in our experiments varied from 2.1 nm to 3.7 nm, which corresponds to a range of nanotube half-circumference shown on inset of Figure 2. Remarkably, Figure 2 shows that self-assembly process starts as soon as the polymer persistence length drops below the half-circumference of the nanotube. These data lend strong support for the self-assembly mechanism based on the polymer wrapping around the nanotube: as soon as the polymer chain can bend enough to match the nanotube surface curvature, the assembly process starts. This critical polymer chain curvature determines the threshold value of the ionic strength at which layer-by-layer self assembly can proceed. We also argue that an alternative mechanism based on rod-like assembly is much less likely to produce this behavior and even if it could, the correlation between assembly threshold and the nanotube persistence length would have been purely fortuitous.

Our results indicate that polymer conformation, or more specifically, the ability of the polymer chain to bend in order to accommodate the support curvature is one of the critical parameters controlling layer-by-layer electrostatic self-assembly on one-dimensional templates. We also found polymer rigidity at low ionic strength can suppress the polyelectrolyte assembly on carbon nanotubes. Our results indicate that this process likely involves wrapping of polymer chains around nanotubes. Thus, controlling the polymer chain conformation is a powerful strategy for fine-tuning self-assembly on one-dimensional templates.

Supporting Information Available: Materials and Methods information. This material is available free of charge via the Internet at http://pubs.acs.org. This work was performed under the auspices of U.S. Department of Energy by the University of California, Lawrence Livermore National Laboratory under contract #W 7405-Eng-48.

REFERENCES
We have studied layer-by-layer polyelectrolyte self-assembly on pristine individual single-wall carbon nanotubes as a function of solution ionic strength. We report the existence of an ionic strength threshold for the deposition, below which the majority of nanotubes remain uncoated. Once the ionic strength reaches the threshold value the majority of the individual nanotubes become coated with polyelectrolytes. Our results indicate that self-assembly process likely involves wrapping of polymer chains around nanotubes and that polymer chain’s ability to bend in order to accommodate the nanotube curvature is one of the critical parameters controlling layer-by-layer electrostatic self-assembly on these one-dimensional templates.