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Morphology-controlled SWCNT/polymeric microsphere arrays by a wet chemical self-assembly technique and their application for sensors

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Abstract
Large-scale morphology-controlled SWCNT/polymeric microsphere arrays can be obtained by a wet chemical self-assembly technique. The loading of SWCNTs, the length of SWCNTs, and the size and nature of polymeric microspheres can easily be controlled. Similar results can also be reached using this method for MWCNTs. In both types of CNTs, they form an interesting interactive ‘net’ structure on spheres and sphere joints. The SWCNT/PS-modified Au electrode was used for detection of uric acid by cyclic voltammetry and single-potential time-based techniques. The preliminary results show that the modified electrode presents good sensitivity and stability to uric acid.

Supplementary data files are available from stacks.iop.org/Nano/17/2988
(Some figures in this article are in colour only in the electronic version)

1. Introduction
Layer-by-layer assembly is an approach based on the alternating adsorption of materials containing complementary charged or functional groups to form integrated ultra-thin films [1–5]. This method provides a powerful tool for nano- and microscale assembly of devices and novel material systems [6, 7]. Single-walled carbon nanotubes (SWCNTs) exhibit unique structural, mechanical, and electronic properties [8–11], and recent studies have demonstrated the use of SWCNTs in nanodevices and sensors [12–15]. Besides, a number of researchers have shown that they can be used to form hybrid SWCNTs/polymer materials using the electrostatic layer-by-layer assembly technique [16–19]. In particular, Correa-Duarte et al. deposited dense mono- and multilayers of multi-walled carbon nanotubes (MWCNTs) on silica, polystyrene, and melamine spherical colloids successfully using the polyelectrolyte-assisted layer-by-layer assembly technique, and measured the conductive properties of CNT-based polymeric thin films [20, 21].

Uric acid is the principal final product of purine metabolism and is commonly found in biological fluids of humans, mainly in serum, blood, and urine. Since abnormally high levels of uric acid in the human body are a symptom of several diseases like gout, Lesch–Nyhan
syndrome, immunodeficiency and gouty nephropathy [22], the
determination of the concentration of uric acid in human blood
or urine is a very important issue in diagnosing diseases.

In this paper, we present an alternative method, a wet
chemical self-assembly technique [23, 24], which
is based on the Au–S chemical bonding, condensation reaction
between the carboxylic termini of SWCNT and the amino
group [25], alternating adsorption of monolayers of individual
components attracted to each other by electrostatic and van
der Waals interactions [1–7, 20, 21], to produce large-scale
SWCNT/polymeric microsphere arrays where the nanotube
distribution density, length, and sphere size can be well
controlled. The SWCNT/PS-modified Au electrode was used for
detection of uric acid at lower concentration by cyclic
voltammetry and single-potential time-based techniques. The
preliminary results show that the SWCNT/PS-modified Au
electrode has better electrochemical performance than that of
the bare and SWCNT-modified Au electrode, which has a good
potential application in biological sensors.

2. Experimental details

2.1. Materials

The SWCNTs (1–10 nm in diameter, 5–20 µm in length)
and MWCNTs (10–15 nm in diameter, 10–20 µm in length)
used in this work were purchased from Iljin Nanotech Co., Ltd (Korea), and commercialized monodispersed polystyrene
spheres (PSs) of different particle sizes, 1.3 µm (Soken,
Chemisnow SX-130H) and 5.0 µm (Soken, Chemisnow SX-
350H) in diameter, were purchased from Soken Chemical
and Engineering Company. The other size PS spheres came
from the Alfa Aesar Company (the standard deviation of the
diameter of PSs is less than 5%). Au (2 mm in diameter) and
Ag/AgCl electrodes were purchased from CH Instruments, Inc.
(USA); the Pt wire electrode was obtained from Bioelectrochemical
System, Inc. (USA). Mercaptoethylamine and uric acid
were purchased from Sigma-Aldrich. Other reagents were
commercially available and were of analytical grade. A
phosphate-buffered saline solution (PBS, pH 7.4) was prepared
dissolving 1.6 g KCl, 64 g NaCl, 1.92 g KH2PO4, and
11.52 g K2HPO4 in 800 ml of deionized water.

2.2. Preparation of SWCNT/polymeric composite

0.2 mg raw SWCNTs and MWCNTs were chemically ‘cut’
by a 10 ml mixture of concentrated sulfuric and nitric acids
(3:1, 98% and 70%, respectively [25]) that were sonicated in
a water bath for 8 h at 40°C. The shortened SWCNTs were
then collected firstly by filtering using 0.2 µm nitrocellulose
membranes (Advantec MFS, Inc. Dublin, CA) and washed
until neutral pH was achieved before redispersing in 10 ml
of ethanol. The ethanol was selected as solvent here because
the filter membrane can be dissolved in acetone. After the
ethanol was volatilized completely, the shortened SWCNTs
were dispersed in 1 l ethanol. Hence we found the as-studied
ethanol was volatilized completely, the shortened SWCNTs
were redispersed in 1 l acetone, hence we found the as-studied
SWCNT solution was as follows: taking 100 ml
0.2 mg 1–1 SWCNT/acetone solution and then evaporating
90 ml acetone at room temperature. It is worth pointing out
here that the concentration is only an approximate value, which
has no obvious effect on our experimental results (considering
the loss during the experiment).

A gold surface of polymeric sphere ordered arrays was
obtained by sputtering with 30 nm Au (99.99%) firstly. 100 µl mercaptoethylamine solution was then dropped onto
the array surface, followed by exhaustive washing using double
deionized water and acetone 6 h later and then dried at
room temperature. The mercaptoethylamine monolayer on
the polymeric microsphere surface was formed and amino groups
were introduced to the surface.

CNT solution was dropped onto the mercaptoethylamine
modified polymeric sphere surface, and then the chemically
shortened SWCNTs accompanied with carboxyl groups at the
open ends reacted with the amino groups on the sphere surface.
In the experiment, the deposition cycle is defined as the process
performed from dropping the first SWCNT solution to the
second one.

2.3. Electrochemical measurements

The as-prepared SWCNT solution was dropped onto the
mercaptoethylamine modified gold electrode surface to give
an SWCNT-modified Au electrode. The detailed information
of SWCNT/PS modified electrode preparation can be obtained
from section 2.2. Cyclic voltammetry (CV) measurements
were performed on a BAS 100W electrochemical analyser
Bioanalytical Systems, Inc., USA). All CV experiments were
carried out in PBS solution in a conventional three-electrode
cell at room temperature. An Ag/AgCl and a Pt wire were
employed as reference and counter-electrodes, respectively.

2.4. Characterization

Atomic force microscopy (AFM) was done with a Nanoscope
III (Digital Instruments, Veeco Metrology, LLC, CA) scanning
probe microscope. The FTIR spectrum was performed using
a model MB-154 analyser (Bomem Inc., CA). All of the
SEM images were obtained with a Philips XL 30 AFEG
scanning electron microscope (Eindhoven, The Netherlands)
except image (d) in figure 4 (Hitachi S-4800 scanning electron
microscope, Japan).

3. Results and discussion

3.1. Properties of shortened carbon nanotubes and composite

The mixture of concentrated sulfuric and nitric acids
were chemically ‘cut’ by a 10 ml mixture of concentrated sulfuric and nitric acids
(3:1, 98% and 70%, respectively [25]) that were sonicated in
a water bath for 8 h at 40°C. The shortened SWCNTs were
then collected firstly by filtering using 0.1 µm nitrocellulose
membranes (Advantec MFS, Inc. Dublin, CA) and washed
until neutral pH was achieved before redispersing in 10 ml
of ethanol. The ethanol was selected as solvent here because
the filter membrane can be dissolved in acetone. After the
ethanol was volatilized completely, the shortened SWCNTs
were dispersed in 1 l acetone, hence we found the as-studied
pipes can be well solubilized using a short 30 min sonication.
This procedure yielded shortened SWCNTs with terminal
carboxylate groups. The simple preparation procedure of
2 mg 1–1 SWCNT solution was as follows: taking 100 ml
0.2 mg 1–1 SWCNT/acetone solution and then evaporating

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characterized by AFM (as shown in figure 1(c)). The images indicate that the morphology of the SWCNT bundles changed from that of highly entangled strands (see figure 1(a)) to flexible-rod-like ones which were thoroughly dispersed in acetone after etching. The length of the shortened carbon nanotubes is about 0.5–1.5 \( \mu \text{m} \). The flexibility of the SWCNTs [27–29], taken together with the carboxylic group located at the end and sidewall of SWCNTs, allows them to bond fairly small particles.

Patterning the resulting ordered arrays with a continuous and homogeneous distribution of carbon nanotubes and conductive properties can be achieved, as outlined in figure 2. Firstly, the colloid monolayer of polymeric spheres was fabricated on the substrate and then coated with a 30 nm thick gold layer; subsequently, it was modified by mercaptoethylamine to introduce the amino group. Lastly, carboxylated SWCNT solution was deposited onto the surface in a controllable way.

3.2. Assembly at different SWCNT concentrations

The monolayer arrays of polystyrene (PS) microspheres with large area around cm\(^2\) in size were fabricated by the spin coating method on the silicon wafer and then heated at 130°C for 40 min in order to guarantee the hard adherence of these arrays to the substrate as we previously reported [30–33]. This colloid monolayer was used as a substrate for the assembly of mono- and multi-layer SWCNTs, which provides a large specific surface area. The PS sphere surface was coated with a 30 nm gold layer firstly in a clean room, and then functionalized with a primary monolayer of mercaptoethylamine. Finally, the SWCNTs were assembled on the PS sphere surface by the covalent bond of –NH–CO–.

Figure 3 shows the typical SEM images of microspheres with diameter of 5 \( \mu \text{m} \) after one deposition cycle under 0.2 and 2 mg l\(^{-1}\) SWCNT concentrations. It is obvious from the figure that SWCNTs can be assembled on the polymeric microsphere surface continuously and homogeneously by controlling the concentration at the deposition step. Firstly, SWCNTs are assembled on the 2D microsphere surface easily due to the condensation reaction between the –COOH and –NH\(_2\); this step is different from physical adsorption, as evidenced by the stability of the layers when ultrasonicated. Owing to the chemical bonds, a few SWCNTs are chemisorbed onto the mercaptoethylamine-modified PS microsphere surface.
adapting themselves to the morphology of the interdigitated thin layer which is different from the multi-layer in the thickness (see figure 3(a)). The elasticity of CNTs has been reported recently, and the existence of the bending mechanism was demonstrated by predominantly tethering the CNTs to the functionalized surface through their carboxyl end groups [21].

With increasing SWCNT concentration, many SWCNTs were sequentially adsorbed onto the sphere surface to form an interdigitated multi-layer because of electrostatic and van der Waals interactions and formed 3D structures (see figures 3(b) and (c)). As can be seen from the SEM images, carbon nanotubes distribute on the surface uniformly and serve as connecting lines between neighbouring spheres within the resulting composite of thin and multi-layers. In addition, the SWCNTs form an interesting interactive 'net' structure on spheres and sphere joints (see figure 3(d)), which can make a structure with good conductivity for future applications of electrochemical sensors and electronic devices.

3.3. Assembly at different deposition cycles

In our experiments, we can control not only the amount of SWCNTs (different SWCNT concentration for each deposition cycle) but also different deposition cycles at a given concentration to obtain SWCNT/PS microsphere arrays. Figure 4 shows SEM images for the SWCNT layers on the monolayer PS microsphere arrays under different deposition cycles (one cycle for image (b), five cycles for image (c), and 20 cycles for images (a) and (d)) at the given concentration of 0.2 mg l$^{-1}$. Obviously, carboxylated carbon nanotubes can be assembled on the surface of micro-polymeric sphere arrays uniformly by controlling the deposition cycles at the given concentration. It is better to point out two important structural characteristics here. First of all SWCNTs uniformly cover the entire surface of the PS spheres without phase separation, and secondly the surface packing layers of the shortened SWCNTs increased rapidly with increasing deposition cycles. This also indicates that a steady SWCNT layer can be obtained by controlling the deposition cycles. Figure 4 shows an interdigitated layer of SWCNT which is deposited in each deposition cycle due to the alternating adsorption of anionic and cationic bipolar components onto PS surfaces via the electrostatic attraction and van der Waals interactions. The final morphology of the multi-layers can be described as a mixture of shortened carbon nanotubes intricately interwoven together on the PS microsphere arrays (see figure 4(d)). We suggest that the increase in the surface packing layers with the deposition cycles was achieved by changing the acting force from the Au–S covalent bond to electrostatic and van der Waals interactions between the carbon nanotubes. Therefore, the wet chemical self-assembling technique should in principle and in experiments, enable us to prepare uniformly dispersed SWCNT/PS microsphere arrays with a controllable packing density by simply regulating the number of deposition cycles or relative amounts of the cut SWCNT components within each of the constituent layers. Apparently, the assembly of SWCNTs with different lengths onto microsphere surfaces of various sizes and natures as templates can also be realized using this method (supplementary information is available at stacks.iop.org/Nano/17/2988).

The wet chemical self-assembling technique can also be used here to assemble multi-walled carbon nanotubes (MWCNTs) on the monolayer PS microsphere arrays. Similarly, prior to assembly, raw MWCNTs were oxidized by an acid mixture to form carboxylated carbon nanotubes (see section 2 for the detailed procedure). The MWCNT layers can easily be controlled by the deposition cycles or MWCNT concentration. Certainly, MWCNTs can also be assembled on any other micro- or nano-spheres with different size. Figure 5 shows the SEM results of assembling MWCNTs on the PS microsphere arrays under 20 deposition cycles at the concentration of 0.2 mg l$^{-1}$. Obviously, PS spheres are connected by 3D MWCNT ‘net’ structures, indicated by yellow arrows.

3.4. Preliminary electrochemical behaviour

The redox reactions of potassium ferrocyanide usually serve as a benchmark in investigating electrochemistry at different carbon electrodes [34]. Here, Fe(CN)$_6^{3-}$/Fe(CN)$_6^{4-}$ as redox pairs were used to investigate the electrochemical performance and electron transfer properties of the SWCNT/PS electrode. Figure 6(a) indicates the potential window of the bare Au electrode, SWCNT-modified Au electrode, and SWCNT/PS electrode for K$_3$Fe(CN)$_6$ in 0.1 mol l$^{-1}$ PBS. Firstly, it is found that the current separation between the forward and backward scans becomes greater that of the bare gold electrode and the SWCNT modified gold electrode. This agrees with the large capacitance feature reported before [35, 36]. The capacitance is calculated from the CV curves, with $C = i/v$, where $i$ is the current and $v$ the scan rate (V s$^{-1}$). At the potential of $-0.05$ V, for example, the effective capacitance is about two orders of magnitude higher than that of the SWCNTs, and over three orders of magnitude higher than the conventional bare gold electrode. This high capacitance is consistent with the large surface area of the SWCNT/PS composite [36, 37]. Secondly, the voltammetric peak at the SWCNT/PS electrode is broad, suggesting slow electron transfer kinetics, presumably due to the macroscopic plane-like structure. Finally, we observed that the SWCNT/PS electrode has a good stability; the result is shown in figure 6(b). This is because SWCNT assembly on the substrate is via the Au–S chemical bond, electrostatic and van der Waals interactions.

Figure 5. SEM images of MWCNTs assembled on PS sphere surface. Image (a) is for MWCNT/PS ordered arrays, whereas image (b) corresponds to multilayer MWCNTs on an individual sphere. Concentration, 0.2 mg l$^{-1}$; deposition cycles, 20.
the electrochemical reaction is very slow. Secondly, at the bare gold electrode, two anodic peaks occur at about 320 and 460 mV. Finally, at the SWCNT/PS-modified Au electrode, there is only a weak and broad oxidation peak around 360 mV. Therefore, we suggest that the electrochemical behaviour was improved by our present effort on SWCNT/PS introduction. Such advantages enable us to study its application for biosensors such as uric acid, aspartate aminotransferase (GOT) and alanine aminotransferase (GPT) detection, etc.

Figure 6. (a) Potential windows for bare gold, SWCNT, and SWCNT/PS electrodes in $5 \times 10^{-4}$ mol l$^{-1}$ K$_3$Fe(CN)$_6$/PBS; (b) the stability of as-studied SWCNT/PS electrode, $5 \times 10^{-4}$ mol l$^{-1}$ K$_3$Fe(CN)$_6$/PBS; scan rate of 100 mV s$^{-1}$.

Figure 7. (a) Cyclic voltammograms of bare gold, SWCNT-modified, and SWCNT/PS-modified Au electrodes in $1 \times 10^{-8}$ mol l$^{-1}$ uric acid/PBS solution at a scan rate of 100 mV s$^{-1}$; (b) real-time detection of the current for the SWCNT/PS-modified Au electrode in uric acid solution for concentrations from $2.5 \times 10^{-7}$ to $1.75 \times 10^{-6}$ mol l$^{-1}$ (each addition $2.5 \times 10^{-7}$ mol l$^{-1}$); scan rate, 100 mV s$^{-1}$; initial potential, 320 mV. The inset shows a plot of the current versus concentration; the black square points are experimental data, and the line is a linear fit through these data.

Therefore, we suggest that the electrochemical behaviour was improved by our present effort on SWCNT/PS introduction. Such advantages enable us to study its application for biosensors such as uric acid, aspartate aminotransferase (GOT) and alanine aminotransferase (GPT) detection, etc.

Figure 7(a) indicates the comparison of bare gold, SWCNT-modified, and SWCNT/PS-modified Au electrodes in $1 \times 10^{-8}$ mol l$^{-1}$ uric acid/PBS solution at a scan rate of 100 mV s$^{-1}$. Firstly, no current peak for uric acid can be observed at the bare gold electrode, suggesting that it is hard for uric acid to accumulate at the gold electrode surface and/or the electrochemical reaction is very slow. Secondly, at the SWCNT-modified Au electrode, there is only a weak and broad oxidation peak around 360 mV. Finally, at the SWCNT/PS-modified Au electrode, two anodic peaks occur at about 320 and 530 mV, and two cathodic peaks occur at around 240 and 460 mV. It can be inferred that uric acid can effectively accumulate at the SWCNT/PS electrode surface and that its electrochemical oxidation was promoted. This is likely due to the fact that the as-studied SWCNT/PS composite has an interesting interactive ‘net’ structure on spheres and sphere joints; on the other hand, the acid treatment creates more oxide defects on the tube sidewalls [38] and ends [38–40], which can be involved in the redox reaction and make the SWCNT more hydrophilic so that the sample can contact the surface better.

Figure 7(b) shows the measurement of current as a function of time and solution concentration at the SWCNT/PS modified Au electrode. It is obvious that the current increases stepwise with discrete changes in concentration of uric acid from $2.5 \times 10^{-7}$ to $1.75 \times 10^{-6}$ mol l$^{-1}$ and that the current is constant for a given concentration; the response is very fast in reaching a dynamic equilibrium upon each addition of the sample solution, generating a steady-state current signal within several seconds; the changes in current are also reversible for increasing and/or decreasing uric acid concentration. A typical plot of the peak current versus concentration (the inset in figure 7(b)) shows that this concentration dependence is linear over concentration from $2.5 \times 10^{-7}$ to $1.75 \times 10^{-6}$ mol l$^{-1}$.

4. Conclusions

In summary, the results reported here demonstrate that large-scale morphology-controlled SWCNT/polymeric microsphere arrays can be obtained by a wet chemical self-assembly technique. The loading of SWCNTs can be easily controlled by the deposition cycles or relative amounts of the cut SWCNT components within each of the constituent layers and the length of the studied SWCNTs can be controlled by the oxidization conditions such as temperature, the amount of reactant (tubes and acid), sonication time, etc. On the other hand, the morphology of the composite can also be controlled.
by using arrays of microspheres with different sizes. Similar results can be reached by using the wet chemical method stated above as for MWCNTs. Besides, both types of CNTs form an interesting interactive ‘net’ structure on spheres and sphere joints. Preliminary electrochemical results show that the SWCNT/polymeric microsphere array electrode has better electrochemical activity to uric acid than that of bare gold and SWCNT-modified electrodes even at lower concentration $1 \times 10^{-6}$ mol l$^{-1}$. In addition, a good linear relationship between current and concentration was obtained from 2.5 $\times 10^{-7}$ to 1.75 $\times 10^{-6}$ mol l$^{-1}$. Future studies will investigate the detailed properties of biosensors such as selectivity, effect of pH, effect scan rate, electron transfer mechanism, etc.

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