**In Situ** Synthesis of Carbon Nanotubes on Heated Scanning Probes Using Dip Pen Techniques

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**Summary:** Carbon nanotubes (CNT) were synthesized on heated scanning probes and under ambient conditions without requiring Chemical vapor deposition (CVD) apparatus or process gases. In this study, dip pen nanolithography (DPN) techniques were utilized for deposition of catalyst precursors on the scanning probe tips in the form of aqueous solution of metal salts—prior to the synthesis of the CNT. A layer of fullerene ($\text{C}_60$) of $\sim 200$ nm thickness was vapor deposited on the scanning probe tip prior to the deposition of the metal catalyst. During the *in situ* synthesis of the CNT on the scanning probes, the temperature of the heated scanning probes reached 350–400 $^\circ$C. Hence the scanning probes were heated in an inert atmosphere to prevent potential oxidation of the deposited fullerene layer. The synthesized CNTs were subsequently characterized using SEM and Raman spectroscopy. The Raman spectroscopy showed peaks in the Radial breathing mode (RBM), as well as the defect (D) and graphitic (G) bands. The RBM peaks indicate that the single walled carbon nanotube (SWCNT) ranged in diameter from 0.9–1.5 nm. The peaks in the Raman spectra are indicative of SWCNT mixtures (metallic and semiconducting) and possibly multiwalled carbon nanotube (MWCNT). Hence this process can be optimized to synthesize SWCNT of a specific chirality (metallic or semiconducting). This study differs from an earlier study reported in the literature involving synthesis of CNT on scanning probes where the process temperatures typically exceeded 700 $^\circ$C, and resulted in synthesis of highly graphitic MWCNT (Sunden, *et al*., 2006). SCANNING 30: 151–158, 2008. © 2008 Wiley Periodicals, Inc.

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**Introduction**

Carbon nanotubes (CNT) are known to form naturally during the combustion processes (Singer and Grumer, 1959; Su and Chen, 2007)—though their initial discovery was attributed to the arc discharge process on graphite electrodes (Iijima, 1991). This technique was refined (Hafner *et al*., 1999) and subsequently several other processes were developed for manufacturing CNT in “bulk” quantities—such as “CVD” (Ebbesen and Ajayan, 1992), and the HiPCo
process (Ouellette, 2002). However, these processes yield a mixture of small proportions of MWCNT and SWCNT while the major proportions of the reaction products are carbon fibers and soot. Also, in these processes the synthesized SWCNT is obtained as a mixture of metallic and semiconducting type. Hence, conventional techniques require expensive separation processes, which are an impediment to their implementation in novel commercial devices that have been envisioned for decades (Robertson, 2007).

Applications of CNTs have been envisioned in superconductors, field-effect transistors (FETs), memory elements, MEMS (microelectromechanical systems), micro/nanosensors and micro/nano electronic devices (Robertson, 2007). Microelectronic applications require precise location and chirality of synthesized CNTs with high spatial density. Hence successful implementation of SWCNT in nanoelectronic circuits has not been possible using current synthesis techniques primarily due to the following three factors (Robertson, 2007): (i) the inability to pattern SWCNT at a precise location, (ii) the inability to precisely control the chirality of the synthesized SWCNT, i.e. metallic or semiconducting type (separation of SWCNT mixtures is expensive and complicated), and (iii) the inability to synthesize SWCNT with high spatial number density.

In earlier attempts (Sunden et al., 2006) nanotube bundles were synthesized on heated microcantilevers using CVD techniques. An iron catalyst layer of 10-nm thickness was vapor deposited on the microcantilever followed by self heating of the cantilever to 800°C in a quartz tube that is typically used for CVD, and in the presence of organic precursors of CNT (e.g. methane and acetylene). Graphitic nanotube bundles were synthesized on the microcantilever, which was reported to be verified by Raman spectroscopy.

In this paper, we introduce a method for the synthesis of CNT, wherein no CVD is required, and the precise positioning of the synthesized CNT is managed through the strategic placement of metal catalysts on a MEMS platform. The MEMS platform consists of a microcantilever scanning probe tip integrated with a microheater at the base of the microcantilever. Thus this technique obviates some significant hurdles presented by other methods (Robertson, 2007) that prevent the exploitation of CNT in micro/ nano electronics. In this study, we report the synthesis of CNT on commercially available heated scanning probe tips (or “probes”). The catalyst precursors were deposited in a “wet” form on the probes from the aqueous solution of metal salts using dip pen techniques. The metal salts served as precursors of the catalysts required for synthesis of CNT. To obviate the CVD process, fullerene precursors were vapor deposited on the probes. The probes were self-heated using the microheaters located at the base of the scanning probes to enable the synthesis of CNT from fullerenes. Thus, this process did not require any process gases for the synthesis of CNT.

Materials and Methods

Experimental Apparatus and Procedure

A scanning probe platform consisting of a microcantilever array with integrated microheaters at the base was used for in situ synthesis of CNT. The heated microcantilevers used in this study consists of an array of eight microcantilevers (ActivePen system, Manufactured by Nanoink Inc.). The silicon nitride microcantilevers are 100 microns long, with 20 microns width and on a pitch of 40 microns. Each cantilever has an electronic serpentine gold heating element at the base, a gold heat spreader (10 microns by 60 microns), which facilitates the heat distribution, and a pyramid shaped oxide sharpened probe at the tip. Figure 1 shows the magnified image (100X) of a microcantilever from the array. The scanning probe tip is located at the distal end and the microheater is fabricated at the base of the microcantilever.

The Mbraun Labmaster double glove box system with gas purification (located at the Center for Nanomanufacturing, University of Texas, Austin); was
used for the vapor deposition of fullerene (C-60) onto the active pens. This instrument consists of a two-chamber glove box system. A spin coater (specialty coating systems model P6700) is available inside the smaller glove box chamber. An Edwards thermal deposition chamber with four sources is available inside the larger glove box. A 150 °C vacuum oven is mounted at one end of the glove box. The ActivePen is placed in the thermal deposition chamber and the C60 fullerene powder (Manufactured by Nano-C Inc., Westwood, MA) was placed in the target container. The target container was electrically heated at a pressure of 1 mTorr to a temperature of 550 °C to enable sublimation of the fullerene powder. The deposition rate by sublimation was estimated to be ~20 nm/s for a total deposition time of 10 s. The fullerene thickness was estimated to be ~200 nm at the end of the vapor deposition process using a quartz microbalance.

A passive microfluidic platform (Inkwells) was used to deposit metal catalysts onto the scanning probe microcantilever array using an atomic force microscope (AFM) platform. Various experimental parameters such as the position of the active probes, actuation current, deposition speeds, deposition patterns, etc., can be varied and controlled through the interfacing software. An Inkwell is a microfluidic platform for delivering solutions of specific chemical precursors to an array of scanning probe tips for their subsequent deposition using dip pen techniques (Banerjee et al., 2005; Rosner et al., 2006). In the Inkwell device, an array of reservoirs connect to microwells through microchannels. A hydrophobic layer is deposited on the top of the substrate to prevent any cross-contamination.

After mounting the scanning probe array on the AFM, the individual microcantilevers can be addressed and actuated by passing current through the respective microheaters at the base. Using this scheme each tip can be retracted or extended and all the tips can be actuated individually or simultaneously. At the beginning of the synthesis experiments all the microcantilevers in the array are actuated sequentially to verify if any of the probes are damaged. The deflection of each cantilever is observed through the real time video window of the interfacing software.

The apparatus is placed in an environmental control chamber, which is connected to the nitrogen supply line from a pressurized nitrogen cylinder. The instrument is equipped with temperature and humidity sensors and an exhaust fan for controlling the humidity of the environment surrounding the probes mounted on the instrument. The values of temperature and humidity can be varied, controlled and monitored through the control software.

After vapor deposition of the fullerene layer, X-ray photoelectron spectroscopy (XPS) was used to characterize the array of scanning probe tips on the ActivePen. Figure 2 shows the representative XPS data for a scanning probe tip. A distinct peak at 286 eV is observed on all the tips confirming that the tips are coated with a layer of fullerene (Cox et al., 1991; Chiang et al., 1992; Mochida et al., 1995).

**Dipping Procedure**

Several metals, which form carbide eutectic are known to catalyze CNT growth, such as, Pd, Pt, 

![Fig 2. An XPS analysis of the scanning probes tips showing a peak at 286 eV, which confirms the presence of the sublimated coating of fullerene.](image-url)
In this study, the individual probes were coated with aqueous solutions of three different metal salts as catalyst precursors (for Pd, Co and Ni). These catalyst precursors were chosen due to their low reduction temperatures to form bare metal from the salt. The catalyst precursors were obtained from 50 mM aqueous solutions of NiCl₂, CoCl₂ and PdCl₂. These salt solutions were used to dip coat the fullerene-coated scanning probes. Inkwells are necessary for the purpose of dipping the probe tips in the desired solution and for confining the dip coating over a localized area on the scanning probe tips. The Inkwell used in this study consists of six reservoirs connected to six microwells through separate microchannels. The catalyst precursor in the form of an aqueous solution of the metal salt is used for coating the scanning probe tips. The salt solutions were obtained by mixing a metal salt with water in a 10-mL Eppendorf tube, followed by decanting the salt solution using a micropipette. A droplet of the salt solution (50–100 nL) is deposited in the reservoir using a syringe. The salt solution flows from the reservoir into the microwell by microcapillary wicking in the microchannels. The Inkwell is then mounted in the AFM. The scanning probes were then mounted on the AFM, and vertically aligned on top of a filled microwell in the Inkwell.

Individual probes are sequentially dipped into respective microwells for coating each tip with a specific catalyst precursor. The aligned cantilever was then lowered into the microwell in steps of 3–5 microns. The dipping of the scanning probe tip was detected when the cantilever snaps into the microwell due to capillary forces. The scanning probe is pulled into the microwell instantly when it makes contact with the liquid meniscus. This can also be observed in the video window of the interfacing software by a change in the reflectivity of the cantilever compared to the other microcantilevers in the array. Figure 3 shows a schematic of the scanning probe tip dipped into the microwell. Figure 4 shows the cantilever array before and during the dipping process.

The dipping procedure was repeated sequentially for all the eight microcantilevers. Of the eight microcantilevers in the ActivePen array, three of the scanning probe tips were dipped into NiCl₂ solution, three were dipped in CoCl₂ solution and two were dipped in PdCl₂ solution. Performing the dipping operation using the microwells helps to constrain the area over which the catalyst is deposited in each scanning probe tip.

After accomplishing the dipping of the scanning probe tips into the catalyst precursor solutions, the Inkwell was dismounted. The humidity was recorded to be 42% indicating ambient moisture and air was in contact with the probe tips. Hence, Nitrogen was passed into the chamber, and the relative humidity was monitored. When the humidity dropped to ∼1% RH after approximately 15 min, indicating that the environmental chamber was filled with nitrogen, the synthesis experiments were performed sequentially for each cantilever. Initially, the current supply to an individual microheater (at the base of a microcantilever) was set at 25 mA. The current was increased in the steps of 2–3 mA, until the limiting value of 50 mA was attained for the supply current (the software setting limited the supply current to this value). This resulted in a heat spreader of the microcantilever attaining a temperature of ∼400 °C, while the scanning probe tip reached a temperature of ∼350 °C (temperature data was obtained using infrared microscopy measurements). The supply current to the microheater at the base of each microcantilever was maintained at 50 mA for a period of 5 min and the power was then switched off. This procedure was repeated sequentially for all the scanning probe tips, before dismounting the probe array.

The low synthesis temperature provides several advantages. It has been reported that lower synthesis temperatures yield a narrower distribution of the diameters (and correspondingly the chirality) of the
synthesized CNT (Robertson, 2007). Also, lower synthesis temperatures can enable easier integration with conventional microelectronics manufacturing and packaging processes.

**Results**

The materials characterization was performed to verify the synthesis of CNT achieved from heating fullerene in the presence of the metal catalysts. It is expected that the catalyst precursors (e.g. PdCl₂) will reduce to form pure metal (e.g. Pd) when heated in presence of fullerenes. The metal catalyst will then nucleate CNT from the fullerenes. Scanning electron microscopy (SEM) was performed to verify if successful synthesis of materials was achieved on the scanning probe tips. Figure 5 shows the SEM image of the scanning probe tip after coating with fullerenes. Figures 6 and 7 shows the SEM image of the scanning probe tip after the synthesis experiments were completed with PdCl₂. At lower resolution the SEM image (Figure 6) shows that precipitated particles were observed on the scanning probe. At a higher resolution (Figure 7), thread-like structures are observed at the base of the scanning probe tip—potentially indicating the formation of CNT bundles. The diameter of the bundles is observed to be between 50–100 nm. The SEM images for the other scanning probe tips were similar to Figure 6. The maximum operating (accelerating) energy of the SEM was 40 kV and the images were obtained from secondary electron emission. However, at higher resolution other structures were observed in Figure 7, which were not detected on the other scanning probe tips. These additional nanostructures (as shown in Figure 7) consist of “hair-like” or “wire” type of structures. These nanostructures were subsequently verified to be SWCNT bundles by micro-Raman spectroscopy.

Figure 7 shows that the spatial density of the synthesized CNT is sparse. Such sparse density is useful for studying the field-emission characteristics from individual CNT as well as for measuring the thermophysical characteristics of individual CNT. This synthesis technique provides the feasibility study for such novel characterization studies. Energy dispersive X-ray spectroscopy (“EDS”) was performed along with SEM imaging of the tips following the synthesis experiments. Table I lists the results from the EDS study. The results show that the tip surface is coated with an abundance of carbon (potentially as fullerene or carbon nanotubes) and small quantities of palladium (which is a remnant of the PdCl₂ solution deposited by dip coating).

Subsequently, micro-Raman Spectroscopy was performed to verify the quality of the synthesized CNT
Fig 6. An SEM image of a microcantilever after the synthesis experiments. The SEM images are obtained at different magnifications. The figure shows catalyst particles formed on the microcantilever after the dipping operation.

Fig 7. An SEM image of a microcantilever after the synthesis experiments. The figure shows carbon nanotube bundles, which appear as hair or wire shaped material synthesized at the base of the scanning probe.

bundles (Model: LabRam IR Confocal Raman Spectroscopy—Microscopy Instrument, Manufacturer: JY Horiba, Japan). The instrument is equipped with a 635-nm wavelength laser with a probe area of 1 micron². This laser was used for the optical excitation of the material synthesized on the probes, and each cantilever was analyzed sequentially. In the Raman spectra, the quality of the synthesized CNT was ascertained by tracing the peaks in the radial breathing mode (RBM), the defect (D) and the graphitic (G) bands. The scanning probe array was mounted on a glass slide for performing Raman spectroscopy.

The RBM mode in Raman spectra is unique to cylindrical symmetry, and is widely used to determine the diameter of SWCNT. Distinct peaks for RBM, D and G were obtained for the two tips coated with PdCl₂ while distinct peaks were not observed for the other scanning probe tips. This indicates that synthesis of CNT using PdCl₂ is feasible at lower synthesis temperatures (i.e. 350–400 °C). The Raman spectra of the scanning probe tips coated with PdCl₂ are shown in Figures 8 and 9.

Figure 8 shows that in the micro-Raman spectra RBM peaks are observed in the range 150–300 cm⁻¹ for the scanning probe coated with PdCl₂ catalyst precursor and fullerene. Since the microcantilevers are composed of silicon nitride, a peak is also observed at 300 cm⁻¹ corresponding to silicon. In the RBM mode distinct peaks were seen at 167, 180, 232, 253, and 275 cm⁻¹.

The diameter \( d \) of the SWCNT can be derived from the location of the Raman shift peaks \( \omega \) in the RBM mode by the following equation (Jorio et al., 2001):

\[
d = \frac{248}{\omega}
\]

where the units of \( d \) is in nm and \( \omega \) is in cm⁻¹. For the range of peaks observed in our experiments—this corresponds to SWCNT of diameters ranging from 0.9 to 1.5 nm.

Typically, Raman spectra reported in the literature that the D band is observed at a Raman shift of \( \sim 1350 \text{ cm}^{-1} \) and the G band at \( \sim 1600 \text{ cm}^{-1} \). If the D band is more prominent (higher or comparable intensity) than the G band, it is indicative of presence of MWCNT. Conversely, if the G band is significantly more prominent with respect to the D band, along with the presence of RBM bands it is indicative of SWCNT (Jorio et al., 2001; Jorio et al., 2004; Thomsen et al., 2004).

Figure 9 shows that for the PdCl₂ there is a peak corresponding to the D band at 1350 cm⁻¹ and a peak corresponding to the G band at 1582 cm⁻¹. This confirms that PdCl₂-coated scanning probes catalyze CNT growth at lower temperatures. Lower processing temperatures were also reported in the literature to provide a narrower range of chirality distribution for SWCNT.

### Table I. Energy dispersive X-ray spectroscopy (EDS) analyses of the microcantilever tip after carbon nanotube synthesis

<table>
<thead>
<tr>
<th>Element</th>
<th>Weight (%)</th>
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<tbody>
<tr>
<td>Si</td>
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<tr>
<td>C</td>
<td>96.85</td>
</tr>
<tr>
<td>Pd</td>
<td>3.15</td>
</tr>
<tr>
<td>Ni</td>
<td>0.00</td>
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<tr>
<td>Co</td>
<td>0.00</td>
</tr>
<tr>
<td>Total</td>
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In situ synthesis of carbon nanotubes using dip pen techniques

Fig 8. Micro-Raman spectra in the RBM mode of the fullerene-coated scanning probe tip (dipped in PdCl$_2$), before (red) and after (blue) the synthesis experiment. The graphs show peaks at the following locations in the RBM mode (cm$^{-1}$): 167, 180, 232, 253, and 275.

Fig 9. Micro-Raman spectra of the fullerene-coated scanning probe tip (dipped in PdCl$_2$), before (red) and after (blue) the synthesis experiment.

(Robertson, 2007). The peaks for G and D band are of comparable intensity, which is indicative of MWCNT being synthesized in this process. However, it can be argued that since the G band has a higher intensity than D band—purely SWCNT are synthesized. The Raman data is also indicative of semiconducting (or mixture of conducting and semiconducting) SWCNT being synthesized in this process.

Conclusion

This investigation provides the proof of concept for the synthesis of SWCNT and possibly MWCNT on scanning probe tips using dip pen techniques. This technique obviates CVD apparatus while enabling synthesis of CNT under ambient conditions. By coating the tips with fullerene and metal catalyst precursors, the process enables the synthesis of CNT without requiring any process gas flow (except for maintaining an inert atmosphere) while the synthesis temperatures can be as low as 350°C. The SEM images show that the synthesized CNT are constrained to a localized area near the scanning probe tip. The process presented in this study proves the feasibility for developing a portable CNT synthesis platform (or instrument) where CNT synthesis can be performed “on demand”.

The Raman spectroscopy showed peaks in the RBM mode in the range of 100–300 cm$^{-1}$, and the D and G peaks were observed at 1350 and 1582 cm$^{-1}$ respectively. The RBM data shows the SWCNT diameter ranges from 0.9 to 1.5 nm. The intensity of D band was slightly higher than the G bands, suggesting the existence of MWCNT, while the presence of the RBM mode is indicative of SWCNT. Hence this process can be optimized to synthesize either SWCNT or MWCNT.
on the scanning probe tips. The process can be optimized for growing CNT of a single chirality (i.e. metallic or nonmetallic SWCNT).

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References


