Effect of Substrate Bias on Growth and Properties of Carbon Nanotubes Deposited under no Hydrogen Introduction by MPCVD

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In order to examine the consequence of replacing hydrogen gas with high substrate bias, processes were successfully demonstrated to fabricate various carbon catalyst-assisted nanostructures, including carbon nanotubes (CNTs), by microwave plasma chemical vapor deposition (MPCVD), under the pure CH₄ as the only source gas and without additional H₂ gas introduction. The Si wafers were first deposited with catalyst by physical vapor deposition (PVD), and then followed by H-plasma pretreatment to obtain the well-distributed catalyst particles on wafer. The pretreated specimens were then deposited in MPCVD to manipulate the various carbon nanostructures. The as-deposited nanostructures were characterized by SEM, HRTEM, Raman spectroscopy, and field emission I-V measurement. The results indicate that the various nanostructures on Si wafers can be manipulated under sufficient high negative substrate bias (< -200 V) without the additional hydrogen introduction. On important process parameters, except the substrate bias, it is interesting to note that effect of deposition time on nanostructures is essentially to extend the more CNTs into the plasma zone, which can result in different purification effect on CNTs, vary the alignment and sizes of CNTs, and expose the more nanostructures near the substrate surface.

1. Introduction
Since carbon nanotubes (CNTs) were first discovered by Iijima in 1991 [1], it has attracted a great deal of attention due to its remarkable electrical [2], mechanical [3], and field emission [4] properties. In order to fabricate CNTs, several techniques, including arc-discharge [5], laser ablation [6], and chemical vapor deposition (CVD) [7, 8] methods, were developed for various applications, such as field emission display (FED), biological sensors, fuel cell etc. However, the most of these processes required to introduce the additional hydrogen gas, especially in the CVD methods. In that, the CNTs synthesized by catalytic decomposition of hydrocarbons were covered by a thick layer of amorphous carbon, which is difficult to remove [9, 10]. Therefore, the purpose of introducing the additional gas source was proposed for etching off amorphous carbon produced during the deposition to minimize the catalyst from being poisoned. As the reasons, it is generally believed that addition of hydrogen or other gases are the required or favor condition to synthesize CNTs. On the other hand, that microwave plasma chemical vapor deposition (MPCVD) could be operated on low temperature with plasma assisting was the well-used system to fabricate CNTs in recent years [11].

In the previous literatures, it was reported that CNTs could not be grown under pure hydrocarbon gases without any additional gases.
by MPCVD. Therefore, Chen et al. used methane and carbon dioxide as the carbon source to synthesize CNTs by MPCVD [12]. In the results, it was shown that amorphous carbon could be limited with the oxygen species plasma. Kenny et al. selected a noble gas (Ar) to replace more reactive H\textsubscript{2} plasma to investigate the function of the diluted hydrocarbons plasma atmosphere by radio-frequency plasma enhanced chemical vapor deposition (r.f. PECVD) [13]. In this study, we proposed a novel process conducted pure CH\textsubscript{4} and high substrate negative bias as the effect of H\textsubscript{2} plasma without introducing any additional gas based on Fe, Co, and Ni as catalysts to fabricate well-aligned CNTs by MPCVD. In addition, various substrate negative bias and deposition time were selected to clarify the major effect to synthesize CNTs under only CH\textsubscript{4} plasma atmosphere.

2. Experimental

CNTs were synthesized under pure CH\textsubscript{4} as gas source, with Fe, Co and Ni as catalysts by MPCVD. Where the Si (100) wafer substrates were firstly deposited with catalyst by physical vapor deposition (PVD) and then followed to obtain the well-distributed catalyst nanoparticles on the substrate surface by H-plasma pretreatment. The main process parameters were substrate bias (0 to -320 V) and deposition time (1 to 20 min). Table 1 describes the specimen designations and their deposition conditions.

3. Results and discussion

3.1 SEM morphologies and TEM images

Figure 1 shows SEM images of the as-grown Ni-assisted carbon nanostructures under different substrate biases (Specimens A1 to A4). It indicates that the substrate bias plays an important role to form CNTs. Under the lower negative bias (less than -200 V) and without introducing additional hydrogen gas source, it is difficult to synthesize CNTs (Figs. 1(a) to 1(c)). This is in agreement with the previous results in the literatures. However, when the substrate bias was raised to -320 V, the well-aligned CNTs can be obtained, as shown in Fig. 1(d). In other words, this is the first work to prove that CNTs can be synthesized by MPCVD under pure CH\textsubscript{4} as source gas without introducing additional gases, if the substrate bias is high enough. In order to examine the deposition time required to form detectable CNTs under a -320 V substrate bias, SEM images in Figs. 1(d), and 2(a) to 2(c) for Ni as catalyst show that deposition time of 4 min is long enough to form CNTs. The results also reveal that the morphologies of CNTs vary from the well-aligned manner with gradually enlarging in size to become spaghetti-like in structures and smaller in sizes (Fig. 2(c)) at longer deposition time (above 20 min). The change in carbon nanostructures after longer deposition time may be due to more severe bombardment to become too smaller in sizes to stand vertically by themselves. The typical SEM morphologies of the nanostructures for Fe as catalyst are shown in Fig. 3(a) and 3(b) after 10 min and 20 min deposition times, respectively. It examines no CNTs presenting except amorphous carbon on the substrate surface (Specimen C2). Especially, the CNTs will be extended more into the plasma atmosphere after longer deposition time to suffer greater etching effect. In order to further clarify the deposited nanostructures, Figs. 4(a), 4(b) and 4(c) show the HRTEM images of the nanostructures with Ni, Co and Fe as
catalysts (Specimen A6, C1, and B3), respectively. These micrographs further support that the nanostructures are all tube-like tip-growth CNTs with catalysts at the tips of the tubes.

3.2 Raman spectra and field emission properties

In order to examine the difference in bonding structures of the as-deposited CNTs for different deposition times, Fig. 5 shows three curves of $I_D/I_G$ ratio of the Raman peaks as a function of deposition time for CNTs with Fe, Co and Ni as catalysts, respectively. It is interesting to note that there are existences of minimum ratio values (1.29, 1.76 and 1.78, respectively) at the moderate deposition time, which are corresponding to the best quality of CNTs in the present deposition conditions. By comparing the SEM morphologies with the corresponding $I_D/I_G$ ratios, it seems to imply that the $I_D/I_G$ ratio is a function of sizes, degree of alignment and amorphous carbon contents of the CNTs themselves, and also the surrounding nanostructures after becoming spaghetti-like CNTs. From Fig. 2(b), 1(d) and 2(c), which are corresponding to the Ni-assisted CNTs with 5, 10 and 20 min deposition times, respectively, the $I_D/I_G$ ratios are about the same (1.8 ~ 1.9). This can be explained by the following reasons: The amorphous carbon contents within CNTs themselves in Fig. 2(b) are relative higher than in Fig. 1(d) and 2(c), but the exposed amorphous carbons embedded on the nanostructures near the substrate surface are the least. The opposite situation is for Fig. 2(c). The CNTs themselves in Fig. 2(c) with less amorphous carbon contents and sizes are due to the preferential amorphous carbon removal by extending more CNTs into the plasma atmosphere. In contrast, for Fe-assisted CNTs, the SEM morphology in Fig. 3(a) indicates dense bundles of CNTs, which are much smaller in sizes with good quality (Specimen B1). The most of the exposed nanostructures in Fig. 3(b) are basically the etched structures near the substrate surface; therefore reveal a much higher $I_D/I_G$ ratio (Specimen B2).

The $I-V$ curves and two corresponding Fowler-Nordheim (F-N) plots of the as-grown Ni-assisted CNTs are shown in Fig. 6 for deposition times ranging from 4 min to 20 min (Specimens A4 to A7). It indicates that the CNTs of 10 min deposition time give the best field emission properties (turn-on field = 5 V/μm, field enhancement factor $\beta$ = 912), though the $I_D/I_G$ ratios for these four specimens are about the same. The corresponding nanostructures for the best field emission are shown in Fig. 1(d) (Specimen A4), where the CNTs remain in well-aligned manner and are etched to become slightly smaller in sizes, to result in the proper tube number density and higher aspect ratio, and to minimize the screening effect. The nanostructures with the worst field emission are shown in Fig. 2(c) for deposition time of 20 min (turn-on field is beyond the instrument upper limit, Specimen A7), where the CNTs are the smallest, but scarce and lie down on the nanostructures embedded by amorphous carbons.

4. Conclusions

A novel process were successfully developed to fabricate the catalyst-assisted CNTs by MPCVD under high negative substrate bias (above -200 V) and pure CH₄ as the only gas
source, but without introducing any additional gas, e.g. H₂. The results indicate that the sufficient high negative substrate bias can replace effect of the additional hydrogen source to form various carbon nanostructures. The deposition time can be an important parameter to manipulate the bombardment effect to purify the CNTs structures and to vary the alignment and sizes, so the desired final nanostructures can be obtained. The deposited nanostructures can vary from the well-aligned CNTs of various sizes, the bundles of CNTs, to become the spaghetti-like nanostructures embedded on the amorphous carbon matrix. The well-aligned CNTs with proper tube number densities, purities and sizes can be the best nanostructures to behave excellent field emission properties. Under the present deposition conditions, the best turn-on field strength can go down to 5 V/μm and the maximum current density at field strength of 10 V/μm is beyond our instrument limit (35 mA/cm²), though there are rooms for much better improvement. On the other hand, the formation of CNTs without additional hydrogen introduction provides the data base to clarify the roles of hydrogen and substrate bias in CNTs growth mechanism.

References

Table 1

<table>
<thead>
<tr>
<th>Specimen designation</th>
<th>Catalyst type</th>
<th>Substrate bias (V)</th>
<th>Deposition time (min)</th>
<th>Substrate temperature (°C)</th>
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H-plasma pretreatment conditions: substrate = 10 nm thick catalyst-coated Si, microwave power = 400 W, substrate bias ≥ 0 V, flow rate of H₂ = 100 sccm, process time = 10 min
Other deposition conditions: base pressure = 5 mTorr, working pressure = 9 Torr, microwave power = 800 W, CH₄ flow rate = 1 sccm
Fig. 1. Effects of substrate bias and substrate temperature on SEM images of the as-deposited Ni-assisted nanostructures, (a) 0 V, 525°C (Specimen A1), (b) -100 V, 550°C (Specimen A2), (c) -200 V, 550°C (Specimen A3), and (d) -320 V, 596°C (Specimen A4).

Fig. 2. Effect of deposition time and the substrate temperature on SEM images of the as-grown Ni-assisted CNTs, (a) 4 min, 481°C (Specimen A1), (b) 5 min, 578°C (Specimen A2), and (c) 20 min, 603°C (Specimen A4).

Fig. 3. Effect of deposition time and the substrate temperature on SEM images of the as-grown Fe-assisted nanostructures, (a) 10 min, 596°C (Specimen B1), and (b) 20 min, 603°C (Specimen B2).

Fig. 4. TEM micrographs of the as-deposited CNTs for using different catalyst materials, (a) Ni (Specimen A1), (b) Co (Specimen C1), and (c) Fe (Specimen B3).

Fig. 5. $I_D/I_G$ ratio of Raman peaks of the as-deposited CNTs as a function of deposition time.

Fig. 6. $I-V$ curves and the corresponding $r-q$ plots for the as-grown Ni-assisted CNTs under different deposition times (Specimen C4 to C7).