# [O] [H] functionalization on carbon nanotube using (O<sub>2</sub>–H<sub>2</sub>) gas mixture DC glow discharge

Tamil Many K. Thandavan · Chiow San Wong · Siti Meriam Abdul Gani · Roslan Md Nor

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**Abstract** The [O] [H] functionalization of carbon nanotube (CNT) was studied using oxygen-hydrogen (O<sub>2</sub>-H<sub>2</sub>) gas mixture direct current (DC) glow discharge plasma technique for cathode/CNT-anode separation of  $0.10 \pm$ 0.01 cm. O<sub>2</sub> and H<sub>2</sub> were fixed at flowrate of 10.0 ml/min in order to obtain gas mixture ratio of 1:1. During the  $(O_2-H_2)$ gas mixture DC glow discharge, current-voltage (I-V) characteristic of gaseous species studied for various settings of gas pressures 1, 2, 3 and 4 mbar. The voltage at gap between cathode/CNT and anode, a breakdown voltage, was identified as "functionalization voltage" ( $V_{\rm func}$ ).  $V_{\rm func}$  was noticed responsible for functionalization of functional groups on sidewall of CNT. The  $V_{\text{func}}$  were recorded as 796, 707, 594, and 663 V for gas pressures of 1, 2, 3 and 4 mbar, respectively. The co-relation between  $V_{\rm func}$  and gas pressure was identified as linear relationship. But a voltage obtained due to the CNT/Cathode fall shows exponential relationship with the gas pressures. The possibility of [O] [H] functionalization was proved using Fourier transmission infra-red (FTIR) spectroscopy. Hydroxyl (-OH), carboxyl (-COOH), and carbonyl (-C=O) functional groups were identical as identified in the FTIR spectra. The field emission scanning electron microscope images show significant changes in the morphology of CNT which proves that the DC gas discharge plasma is a possible technique for [O] [H] functionalization on the sidewall of CNT.

**Keywords** Functionalization · Hydroxyl · Carbon nanotube · DC glow discharge

T. M. K. Thandavan (⋈) · C. S. Wong · S. M. A. Gani · R. M. Nor Department of Physics, University of Malaya, Lembah Pantai,

50603 Kuala Lumpur, Malaysia e-mail: tamilmanykthandavan@gmail.com

# Introduction

The discovery of carbon nanotube (CNT) in 1991 has unpaved (Iijima 1991) numerous approaches. Many investigations have been carried out to enhance diverse applications of CNT in chemical sensing (Pisco et al. 2008; Arben Merkoci et al. 2005), plastic transistors (Masashi Shiraishi et al. 2004), optoelectronic devices (Vanmaekelbergh et al. 2007), field emission (Jong Min Kim et al. 2000; Saito Yahachi Saito et al. 2002; de Jonge and van Druten 2003) and molecular electronics (Singh et al. 2006). Both single-walled carbon nanotube (SWCNT) and multiwalled carbon nanotube (MWCNT) are proven as considerable nanomaterials in these diverse applications. However the nature of CNT as agglomeration in bundle, the nonreactive and the presence of amorphous layer have inflicted their widespread use in many applications (Felten et al. 2005). Furthermore, insolubility in water, polymer resins and other solvents have showed difficulties in processing CNT into a thin film (Kakade and Pillai 2008). However, surface modification on CNT was found to be an efficient technique in introducing nanoparticle or functional groups; hydroxyl (-OH), carboxyl (-COOH) and ammonia (-NH<sub>3</sub>) on the sidewall of CNT.

The well-known surface functionalization on CNT involves process like altering the wetting and adhesion characteristics while improving dispersions of functional groups in the matrices of CNT. The attachment of functional groups on the sidewall of CNT without significantly changing the desirable properties of CNT provides a promising approach in its various fields of applications (Francesco Mercuri and Antonio Sgamellotti 2007). Conjunctionally, a number of research groups have demonstrated surface modification on CNT through both physical and chemical approaches, including polymer wrapping



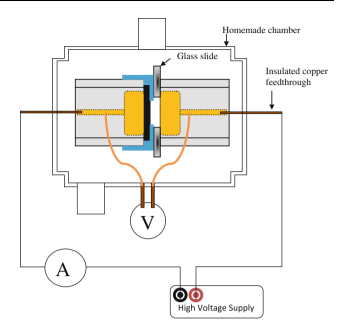
(O'Connell et al. 2001; Inna Gurevitch and Simcha Srebnik 2007), fluorination (Hamwi et al. 1997; Touharaa et al. 2002), mechano-chemical treatment (Konya et al. 2002; Chen et al. 2004) and plasma treatment (Bogaerts et al. 2002). Especially, the plasma treatment has exhibited advantage of non-polluting, which is not trifling for industrial fabrication. This technique also allows increase in quality and production quantity which are required for commercial use. In addition, the processing period of plasma treatment is shorter than any other techniques mentioned earlier. It also enables wide range dispersion of functional groups on the side wall of CNTs.

In this paper, we detailed the preparation of cathode/ CNT-anode electrode system for DC gas discharge plasma treatment of  $\rm H_2$  and  $\rm O_2$  gas mixture. The processing parameter, gas pressures, was related with the  $V_{\rm func}$  to show the –OH functionalization on the sidewall of CNT.

## **Experimental arrangement**

SWCNTs dissolved in an acetone solution were dispersed on silicon, Si(001) small chips cut into 1 by  $1 \pm 0.1$  cm dimension. The Si(001) chips were cleaned in an ultrasonic bath of acetone solution to remove the presence of any oxide layers and impurities on the surface. The substrates further rinsed in deionised (DI) water to remove existing acetone solution. Three drops of CNT solution dispersed onto the cleaned silicon chip and further dried for 24 h in desiccators filled with drying agent, silica gel. This drying process could remove any moisture content from the CNT sample. A piece of non conductive tape was cut into a desired size and a small square hole  $5 \times 5$  mm was placed at the centre of the tape. Then the tape was stuck onto the CNT sample and clamped to the brass electrode which acts as a cathode. The cathode/CNT was separated from an anode brass electrode using 1 mm thick glass slides. The prepared electrode system as shown in Fig. 1 was placed into a homemade stainless steel chamber that consists two sets of copper electrical feedthroughs. The chamber was then vacuumed to pressure about  $10^{-5}$  mbar. The cathode/ CNT and anode electrodes were connected to the high voltage supply via the copper electrical feedthroughs. O<sub>2</sub> and H<sub>2</sub> gases are flowed at gas mixture ratio of 1:1 into the vacuumed chamber. Potential difference from high voltage supply was slowly increased to maximum to obtain a current-voltage (I-V) characteristic of [O] [H] plasma between the cathode/CNT and anode for gas pressures of 1, 2, 3 and 4 mbar. The I-V characteristics were used to study and evaluate the [O] [H] functionalization on the sidewall of CNTs. The plasma-treated CNTs were compared with the non-plasma-treated CNT using FESEM (Quanta 200F) and FTIR spectroscopy.





**Fig. 1** Illustration of electrode system inside the home made stainless steel chamber. The electrode system consists of cathode brass electrode, anode brass electrode, two glass slides and Si (001) substrate contents of CNT shows the cathode/CNT-anode separation of  $0.10 \pm 0.01$  cm. The electrical connection setup consists of 5 kV power supply, ammeter, and voltmeter

#### Results and discussions

## **FESEM**

Figure 2 exhibits a comparison of FESEM images of nonplasma-treated CNT (Fig. 2a) and plasma-treated CNT (Fig. 2b, c, d) which reveals structural and morphological changes on the surface of CNTs. The morphology of nonplasma-treated CNT shows that almost all CNTs are not aligned in parallel to one another and found to be intersecting one another to form agglomeration in bundle. The CNTs are visible and randomly oriented and spread on the Si(001) exhibiting small grooves and small bumps of CNT bundles. But this morphology has been slightly altered after treated with DC gas discharge plasma of O2 and H2 at the gap of cathode/CNT and anode. Figure 2b shows morphology of CNT after the DC gas discharge plasma for gas pressure 1 mbar. The morphology of CNTs bulging outward forming larger grooves and larger protuberant are vertically aligned from the Si substrate. Figure 2b1 shows magnified version of Fig. 2b. A closer look at the image reveals formation of flashes that cover large surface of CNT and few CNTs in a range of 10-15 nm joint together at point T (refer Fig. 2b1) and formed a bundle of CNTs. Even two CNTs are aligned in parallel. Figure 2c, c1 (magnified image) shows morphology obtained after DC gas discharge plasma for gas pressure of 2 mbar. Larger protuberant is visible and more flashes cover the surface of Appl Nanosci (2012) 2:47–53

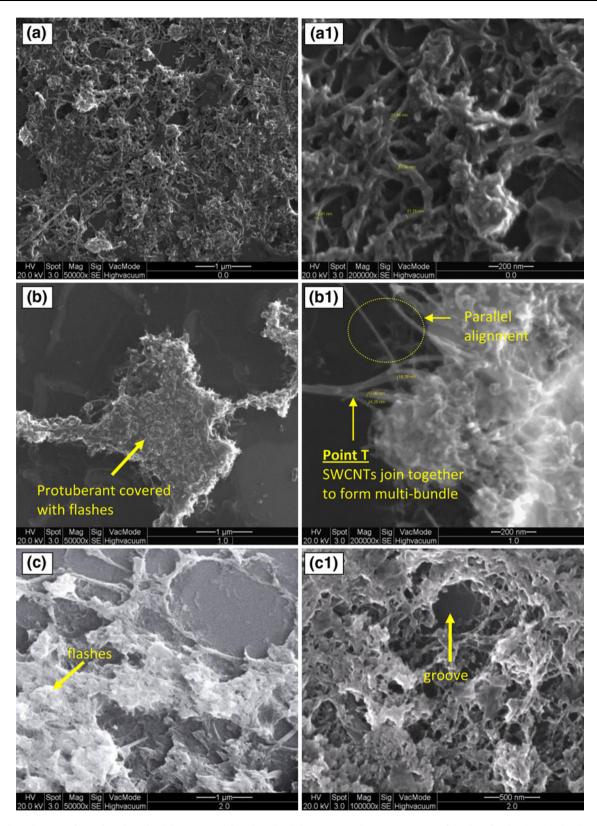


Fig. 2 SEM images of non-functionalized CNT (a) and functionalized CNT prepared at gas pressures of 1 mbar (b), 2 mbar (c), 3 mbar (d) and 4 mbar (e). a1, b1 are  $4 \times$  magnification of a and b, respectively. c1, d1 and e1 are  $2 \times$  magnification of c-e, respectively



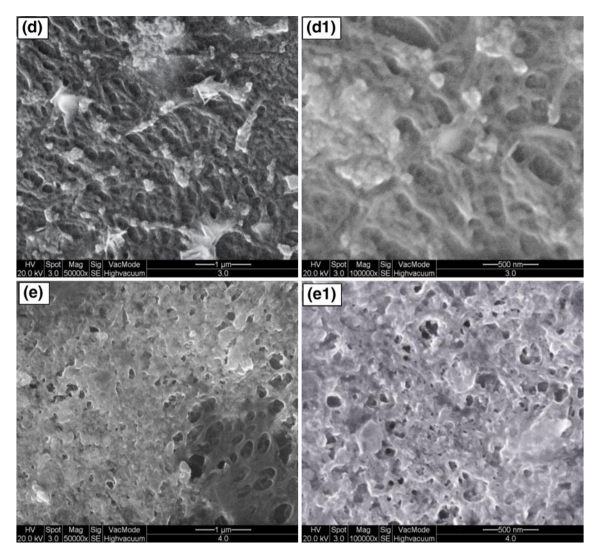


Fig. 2 continued

CNTs compared to Fig. 2a1 (non-plasma treated), b1 (plasma treated at 1 mbar). Figure 2d, e shows DC gas discharge plasma treated at pressures of 3 and 4 mbar, respectively. The respective magnified image of Fig. 2d, e is shown as Fig. 2d1, e1 exhibits formation of flashes that cover larger surface area of CNT. Comparatively, Fig. 2d1, e1 shows flashes that cover larger surface area than the Fig. 2b1, c1.

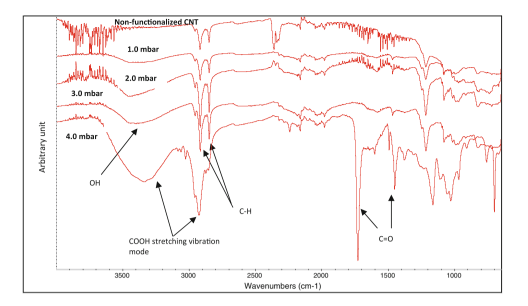
# **FTIR**

Figure 3 shows significant changes in the FTIR spectra of non-plasma-treated CNT and plasma-treated CNT at gas pressures of 1, 2, 3 and 4 mbar. The spectra of plasma-treated CNTs are compared to non-plasma-treated CNT spectrum to identify the possibility of chemical bonding in the matrices of CNT. Non-plasma-treated CNT only exhibits C-H vibration mode around 2,857

and 2,917 cm<sup>-1</sup>. The CNT samples treated by DC gas discharge plasma of O<sub>2</sub> and H<sub>2</sub> at gas pressures of 1, 2, 3 and 4 mbar show broaden peak around 3,400 cm<sup>-1</sup> that ascribes the -OH stretching vibration (Men et al. 2008; Hsin et al. 2006). Presence of peak at wavenumber 2,850 and 2,917 cm<sup>-1</sup> shows the presence of C-H stretching frequencies (Niu et al. 2007) which correspond to alkyl groups in both plasma-treated CNT and nonplasma-treated CNT. The intensity of the C–H stretching peak noticed to be increasing as the gas pressures increased from 1 to 3 mbar. But for gas pressure of 4 mbar the two peaks of C-H vibration further stretched and broadened to -OH broaden peak to form -COOH stretching vibration mode. Besides this, strong vibration mode at 1,728 cm<sup>-1</sup> shows the evidence of -C=O chemical bonding in the matrices of CNT treated with DC gas discharge plasma for gas pressures of 4 mbar only.



Fig. 3 FTIR spectra of nonplasma-treated CNT in contrast with DC gas discharge plasmatreated CNT for gas pressures varies from 1, 2, 3 and 4 mbar. The spectra were arranged in ascending order as the pressures stack down



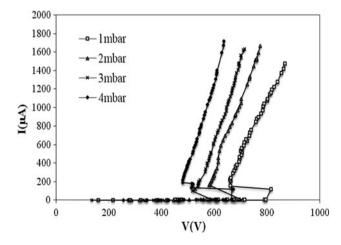
## *I–V* characteristics

Figure 4 shows the *I–V* characteristics of the CNT samples for various settings of gas pressures 1, 2, 3 and 4 mbar. The I-V characteristics are not smooth, indicating instabilities of emission current, presumably from cathode bombardment during discharge of O2 and H2 gases at the gap of cathode/CNT-anode. The breakdown voltages are identified as 796, 707 and 594 V for gas pressures of 1, 2, and 3 mbar show a descending trend as the pressures increased. But the breakdown voltage for gas pressure at 4.0 mbar shows higher voltage (663 V) than at the 3.0 mbar (594 V). I-V characteristics of 1, 2, 3 and 4 mbar show similar trend. Immediately after the breakdown voltage of the gas mixture, a sharp increase in voltage and current are seen at (815 V, 115  $\mu$ A) and (667 V, 114  $\mu$ A) for the gas pressures of 1.0 and 4.0 mbar. As the potential difference is increased, the gap voltage of cathode/CNT-anode shows sudden decrease to 660 to 580, 518 and 512 V for the respective gas pressures of 1, 2, 3 and 4 mbar. Further increase in potential difference of high voltage supply has exhibited proportional relation between the current and gap-voltage.

## Discussion

The *I–V* characteristic of (O<sub>2</sub>–H<sub>2</sub>) gas mixture DC glow discharge at the cathode/CNT-anode separation is discussed based on the typical *I–V* characteristics reported for DC gas discharge plasma of Argon (Bogaerts et al. 2002). A very slight increase in current (almost constant current) is noticed in the gap between cathode/CNT-anode brass electrodes for the increased voltage of main supply. The

saturation regime is almost unnoticeable due to a very low current  $<10^{-10}$  A. This explains that emissions of primary electrons from the CNTs (attached to cathode brass electrode) instead of cathode brass are due to the omnipresent cosmic radiation. It is suggested that active sides  $C^+$ ,  $C^{2+}$ , and  $C^{3+}$  on the exterior sidewall of CNTs ( $C \rightarrow C^{n+} + e^{-}$ ; where n = 1, 2, and 3) are created. The combination of saturation and Townsend regime is a non-selfsustaining regime due to low electric field across at the gap of cathode/CNT-anode. At this regime, ionization of O<sub>2</sub> and H<sub>2</sub> could not be established. However, increasing the main voltage supply to the gap above 180 V has caused a transition from non-selfsustaining discharge to selfsustaining discharge. This could be due to the increased electric field at the gap of cathode/CNT-anode. At higher electric field, O<sub>2</sub> and H<sub>2</sub> are ionized by the impact of emitted primary



**Fig. 4** *I–V* characteristics of O<sub>2</sub> and H<sub>2</sub> gas discharge at the cathode/CNT-anode separation of 1 mm for pressures of 1, 2, 3 and 4 mbar



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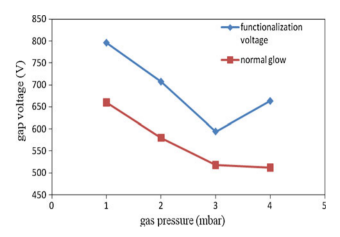


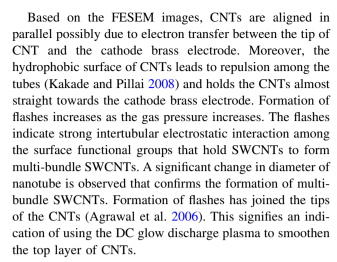
Fig. 5 Gap voltages of CNT-anode; functionalization voltage and normal glow voltage at gas pressures of 1.0, 2.0, 3.0 and 4.0 mbar

Table 1 Gap-voltage of CNT-anode during [O] [H] functionalization

Gas pressure (mbar)	Gap-voltage of CNT-anode	
	Voltage breakdown (V)	Normal glow voltage (V)
1.0	796	660
2.0	707	580
3.0	594	518
4.0	663	512

electrons from CNTs (Bogaerts et al. 2002). The primary electron collides inelastically with  $O_2$  gas to form oxide ions which ascribed as  $O_2$  breakdown ( $O_2 + 4e^- \rightarrow 2O^{2-}$ ) whilst  $H_2$  breaks down into positive ions ( $H^+$ ) and electrons. These ion species ( $O^{2-}$  and  $H^+$ ) chemically could attach to the active sites of sidewall of CNT to bond with functional groups through the phenomena called as "functionalization". During the functionalization, the gap voltage of cathode/CNT-anode marked as gas breakdown voltage which is not selfsustainable and not visible showed a sudden drop to normal glow voltage. Therefore, the breakdown voltage can be referred as functionalization voltage ( $V_{\rm func}$ ).

The  $V_{\rm func}$  and normal glow voltage are further analyzed (refer Fig. 5) as per detailed in Table 1. Figure 5 shows the corresponding relationship between the functionalization voltage and pressures. The  $V_{\rm func}$  decreases linearly to the gas pressure ranged from 1.0 to 4.0 mbar. This explains that by setting the gas pressure to higher value, the presence of number of gas particles density per cm<sup>3</sup> at the gap has increased and distance between the particles is decreased. This allows the particles to collide inelastically and more frequently with each other at the low gap voltage.



## **Conclusions**

[O] [H] functionalization of CNT using (O<sub>2</sub>-H<sub>2</sub>) gas mixture DC glow discharge plasma technique is obtained at cathode/CNT-anode separation of  $0.10 \pm 0.01$  cm. The I-V characteristics of (O<sub>2</sub>-H<sub>2</sub>) gas mixture DC glow discharge showed breakdown voltages 796, 707, 594, and 663 V for gas pressures of 1, 2, 3 and 4 mbar. The breakdown voltage is identified as "functionalization voltage" that responsible for the functionalization of functional groups on the sidewalls of CNTs. FTIR has confirmed structural modification on the sidewalls of CNT which corresponds to increasing in the intensity of -OH stretching mode that broadens from 3,200 to 3,600 cm<sup>-1</sup>. Increased (O<sub>2</sub>-H<sub>2</sub>) gas mixture pressure up to 4 mbar has functionalized the sidewalls of CNTs with -COOH and -C=O functional groups. FESEM images proved the formation of bulging and grooves after the applied voltage between cathode/CNT and anode.

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