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Effects of O₂ plasma and UV-O₃ assisted surface activation on high sensitivity metal oxide functionalized multiwalled carbon nanotube CH₄ sensors

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The authors present a comparative analysis of ultraviolet-O₃ (UVO) and O₂ plasma-based surface activation processes of multiwalled carbon nanotubes (MWCNTs), enabling highly effective functionalization with metal oxide nanocrystals (MONCs). Experimental results from transmission electron microscopy, scanning electron microscopy, x-ray photoelectron spectroscopy, and Raman spectroscopy show that by forming COOH (carboxyl), C-OH (hydroxyl), and C=O (carbonyl) groups on the MWCNT surface that act as active nucleation sites, O₂ plasma and UVO-based dry pretreatment techniques greatly enhance the affinity between the MWCNT surface and the functionalizing MONCs. MONCs, such as ZnO and SnO₂, deposited by the atomic layer deposition technique, were implemented as the functionalizing material following UVO and O₂ plasma activation of MWCNTs. A comparative study on the relative resistance changes of O₂ plasma and UVO activated MWCNT functionalized with MONC in the presence of 10 ppm methane (CH₄) in air is presented as well. © 2017 American Vacuum Society. [http://dx.doi.org/10.1116/1.4993579]

I. INTRODUCTION

Methane (CH₄) gas has a 100 year global warming impact factor of 24–36 compared to CO₂.¹ With the emergence of the U.S. as the world’s leading producer of natural gas, it is important to enable widespread monitoring of CH₄ emission from natural gas infrastructures. Metal oxide chemoresistive sensors are widely used to sense CH₄.²–⁴ Continuous heating is necessary for these sensors to initiate the surface chemisorption of O₂, a prerequisite to detect CH₄, often requiring 100 s of milliwatts of power.²–⁴ Carbon nanotube (CNT)-based chemoresistor sensors have demonstrated ppm levels of gas sensing at room temperature, with a power consumption of only a few milliwatts.⁵ This is a direct outcome of CNT’s high surface-to-volume ratio and outstanding modulation of electrical conductance during interaction with gas species. However, functionalizing particles (ranging from metal,⁵ metal oxides,⁶ and polymer coating⁷ to biomolecules⁸) must be deposited uniformly on the surface of pristine carbon nanotubes in order to enable effective and reversible electrical modulation in the presence of target gas species. Unfortunately, in general, the surface of CNTs shows poor affinity with the functionalizing materials.⁹–¹³ Consequently, before applying the functionalization materials, activation of the inert graphitic surface of the CNTs is necessary.⁹–¹¹

We present here a comparative analysis of novel UV-O₃ (UVO) and O₂ plasma-based surface activation processes, which enables highly effective functionalization of multiwalled carbon nanotubes (MWCNTs) with metal oxide nanocrystals (MONCs). O₂ plasma and UVO-based dry surface activation techniques have not been applied in CNT-based CH₄ chemoresistor sensors before.⁵,¹⁴,¹⁵ Weak affinity among the CNT surface and the functionalizing nanoparticles, resulting from the absence of surface activation, may greatly affect the sensor’s reversible response to low ppm

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methane concentrations. Experimental results from transmission electron microscopy (TEM), scanning electron microscopy (SEM), x-ray photoelectron spectroscopy (XPS), and Raman spectroscopy show that by forming carboxyl (COOH), hydroxyl (C-OH), and carbonyl (C=O) groups on the MWCNT surface, which act as active nucleation sites, O2 plasma and UVO-based dry activation techniques greatly enhance the affinity between the MWCNT surface and the functionalizing MONCs. We have implemented MONCs such as ZnO and SnO2 as the functionalizing material following UVO and O2 plasma activation of MWCNTs. MONCs were deposited by the atomic layer deposition (ALD) technique.16–18 These metal oxides are less expensive than previously reported functionalizing materials (e.g., Pd) used in CNT-based CH4 sensors.5,6 Electron transport is energetically favorable in the ZnO-MWCNT or SnO2-MWCNT junctions, whereas Pd forms a significant Schottky barrier with bare CNTs.5,6,19,20 A comparative study on the relative resistance changes of O2 plasma and UVO activated MONC functionalized MWCNT chemoresistive sensors, in the presence of 10 ppm CH4 in air, is presented. After CH4 exposure, the sensors were recovered to baseline resistance by N2 purging. An analysis of the effect of relative humidity (RH) on the sensor response is also presented, showing superior performance to other previously reported CNT CH4 sensors.

Traditionally, the surface of bare CNTs is activated by exposing them to high temperature vapors10 and/or using wet chemistry.11 High temperature or air exposure may actually destroy or excessively damage the CNTs.10 Acid treatments used in wet chemistry can considerably reduce the mechanical and electric performance of the tubes by introducing large numbers of defects.21 Wet chemistry also involves additional steps, such as dissolution, sonication, mixing, and drying, which often causes undesirable agglomeration of treated CNTs.10

To increase the efficiency of CNT functionalization, two alternative dry activation processes have been proposed: (1) gas plasma9–11 and (2) UVO treatment.22

Due to the interaction of surface C atoms with active O atoms during O2 plasma or UVO activation and subsequent exposure to the atmosphere, chemical groups such as COOH, C=O, C-OH, and ether (C-O-C) are formed on the MWCNT surface.11–13,23,24 These groups act as active sites for the nucleation of MONCs.11–13,22 Both the O2 plasma and UVO exposure have no effect on the aspect ratio (i.e., length to diameter ratio) of the MWCNTs.21

II. EXPERIMENT

A. Sensor fabrication

We fabricated surface activated MONC (ZnO or SnO2) functionalized MWCNT chemoresistive CH4 sensors using the following fabrication steps: (1) lift-off based photolithography, (2) O2 plasma or UVO based surface activation, and (3) ALD based functionalization. The sensor concept is illustrated in Fig. 1(a). Figure 1(b) shows an SEM image of the sensor where functionalized MWCNTs can be seen deposited between a pair of Au electrodes.

Interdigitated Au electrodes were fabricated by photolithography. Details on the fabrication process can be found elsewhere.25

MWCNT (98% pure) with an average diameter of 12 nm, an average length of 10 μm, and a specific surface area of 220 m2/g was purchased from Sigma Aldrich. Using a microsyringe, an aliquot of 50 μl from a 1 g/50 ml solution of MWCNT-ethanol was deposited on an active area of 1 mm2 on the fabricated Au electrodes. It was followed by baking the devices at 75 °C to remove ethanol and improve adhesion.

The deposited MWCNTs were O2 plasma activated in a reactive ion etching chamber (March plasma CS-1701). The base pressure of the plasma chamber was almost 40 mTorr. O2 was introduced at a flow rate of 20 sccm, while the pressure was maintained at 160 mTorr during the process. O2 plasma was generated by applying a radio frequency of 13.56 MHz with a power of 100 watts. The duration of the plasma treatment was 5 min. A UVO cleaner (Nanonex Ultra 100) was used for the UVO treatment of the MWCNT surface where a 185 nm UV was radiated to atmosphere for generating O3 and activating the MWCNT surface. The process duration was 20 min.

Using diethylzinc [(C2H5)2Zn] as a precursor, ALD of ZnO on the surface activated MWCNTs was performed using an Arradiance Gemstar ALD tool (details can be found in Ref. 25).

ALD was also used to deposit SnO2 nanocrystals (NCs) onto the surface activated MWCNTs. The growth was carried out using an Ultratech Savannah S200 with
tetrakis(dimethylamino)tin(IV) as a precursor (details can be found in Ref. 26).

B. TEM sample preparation and imaging

Holey carbon films on Cu grids were used to prepare the TEM sample. Using a microsyringe, an aliquot of 50 µl from a 1 mg/50 µl solution of MWCNT-ethanol solution was deposited on the TEM grid. It was followed by baking the devices at 75 °C to remove ethanol and improve adhesion. MWCNTs were surface activated in a similar manner as described in Sec. II A (5 min O2 plasma or 20 min UVO).

ALD of ZnO or SnO2 was performed following a similar approach described in Subsec. II A.

A JEOL 2100F TEM operating at 200 kV was used to characterize the atomic scale morphology and crystal quality of the MONCs deposited on MWCNT surfaces.

C. XPS sample preparation

Au of 20 nm was deposited on a clean Si wafer using electron beam evaporation. Using a microsyringe, an aliquot of relatively higher density solution (1 mg/1 µl) of MWCNT-ethanol was deposited on the Au-coated Si wafers. It was followed by baking the devices at 75 °C to remove ethanol and improve adhesion. MWCNTs were surface activated in a similar manner as described in Sec. II A (5 min O2 plasma or 20 min UVO).

D. Test setup and approach

Sensors were exposed to a 10 ppm mixture of CH4 in synthetic air (20.81% of O2 and 79.19% of N2; prepared by Praxair Inc.). The flow rate was maintained at 0.94 l/min with a residence time of 4.5 min inside the plastic test chamber (details elsewhere25). After the CH4 exposure, the sensors were flushed with N2 (the same flow rate as CH4, 0.94 l/min). The electrical signal obtained from the sensors was recorded using a custom interface circuit connected to a computer. A onset HOBO U12 series data logger temperature and RH sensor was used to continuously monitor and record the RH and temperature inside the plastic test chamber during the test.

III. RESULTS AND DISCUSSION

A. Sample characterization

High resolution XPS (hν = 650 eV) was carried out using beamline 4-ID-C at the Advanced Photon Source, Argonne National Laboratory. MWCNTs were deposited on gold-covered silicon substrates and subsequently activated by O2 plasma or UVO. Binding energies were calibrated to the Au 4f binding energy of 84.0 eV. Quantification was performed using XPS data analysis software CASAXPS. Figure 2 shows the C 1s and O 1s peaks originating from pristine [(a) and (d)] and 5 min O2 plasma activated [(b) and (e)] and 20 min UVO activated MWCNT [(c) and (f)]. The assignments of the C 1s and O 1s components were based on reported spectra containing specific oxygen functional groups.24,27–29

The measured spectra were fitted to a function having 70% Gaussian and 30% Lorentzian character, after performing a Shirley background correction. The C1 component centered at 284.3 eV represents the sp2 graphitic component. The components at 285.4, 286.5, 287.7, and 289.1 eV (C2 up to C5) were assigned to C atoms forming C-OH, C-O-C, C=O, and COOH functional groups, respectively;24,27–29 the components at 531.1 eV (O1), 532.2 eV (O2), 533.5 eV (O3), and 534.7 eV (O4) were assigned to O atoms forming C=O, C-OH, C-O-C, and H2O, respectively.24,27–29

The normalized peak areas (NPA) of various components of C 1s and O 1s spectra were calculated with respect to the area of their respective C1 component (sp2) (Table I). Comparison among the NPA of C1 component (sp2) (Table I). Comparison among the NPA of C 1s and O 1s components in several samples suggests that COOH is the primary chemical group created by the surface activation process. The NPA of C-OH components (C2 and O2) are significantly larger in the surface activated sample compared to the pristine sample, suggesting the strong presence of C-OH in the surface activated...
MWCNT as well. The NPA of C4 and O1, representing the C=O group, were found to be highest in the 5 min plasma activated MWCNT but insignificant in pristine and 20 min UVO activated MWCNT. On the other hand, the NPA of the C3 and O3, representing the C-O-C functional group, were found to be highest in the pristine MWCNT (Table I).

It is well known that active π bonds in C=C are dissociated during plasma/UVO activation and –C free radicals are produced. Subsequently, –C free radicals are oxidized by active O atoms present in the O2 plasma and UVO, resulting in C-O and C=O bonds. After prolonged interaction with plasma/UVO, C=O is further oxidized and O-C=O is formed. Due to atmospheric exposure, C-O and O-C=O stabilize by reacting with ambient H2O and generate C-OH and COOH, respectively. This is the probable cause of the strong presence of COH and COOH groups in our plasma/UVO activated MWCNTs. Surface C atoms of pristine MWCNT react with atmospheric H2O to create C-O, a probable cause of the presence of the C-O-C group in pristine MWCNT.

In summary, the XPS results corroborate that the surface activation process produces the COOH functional group along with C-OH and C=O. In later steps, these groups help in nucleating the functionalizing MONCs on the surface of the MWCNTs.

The TEM micrographs show that MONCs are not visible on the surface of the nonactivated but ALD processed MWCNTs [Fig. 3(a)]. Uniform deposition of ZnO-MONCs was found on the surface of the activated MWCNTs [Fig. 3(b)]. The clearly visible lattice fringes in the higher resolution TEM (HRTEM) image in Fig. 3(d) illustrate the wurtzite structure of the ZnO MONC and its good crystalline quality. The interplanar spacings of 2.8, 2.68, and 2.48 Å correspond to the (100), (002), and (101) planes of ZnO, respectively. The HRTEM image in Fig. 3(c) shows the atomic scale morphology of rutile SnO2 MONCs deposited on the MWCNT surface. The interplanar spacings of 2.6 and 3.3 Å correspond to (101) and (110) planes of SnO2, respectively. TEM results validate the hypothesis that surface activation of the MWCNTs is essential for effective functionalization, i.e., nucleation and stronger binding of the MONCs to the surfaces of the MWCNTs.

Room temperature Raman spectroscopy was performed using a Renishaw Invia micro-Raman system with a 514nm laser. Three types of ZnO ALD functionalized MWCNT samples were used in the Raman characterization: (1) O2 plasma activated, (2) UVO activated, and (3) nonactivated, i.e., untreated. Raman spectra, illustrated in Fig. 4, reveal that after the surface activation and ALD functionalization, the D, G, and G' band peaks of the MWCNTs are preserved, while additional Raman peaks originating from the ZnO NCs appeared. The characteristic Raman peaks of ZnO NCs represented by m, n, o, and p were observed only on the O2 plasma activated samples (1), n, o, and p were observed on the UVO activated samples (2), while none of these peaks were visible on the untreated MWCNT samples (3), consistent with the hypothesis that the ZnO NC functionalization is enhanced in surface-activated MWCNTs. The peaks described in Fig. 4(b)—200.6 cm−1 (m), 324.25 cm−1 (n), 430.84 cm−1 (o), 569.87 cm−1 (p)—correspond to 2E2(LO), E2(high) − E2(low), E2(high), and A1(LO) modes of ZnO, respectively, suggesting that surface activated samples have ZnO NCs with high crystalline quality. In addition, the characteristic ZnO Raman peaks are sharper in O2 plasma activated MWCNTs than in UVO activated MWCNTs. The full width at half maximum of the A1(LO) peak was found to be 50.5 and 62.29 cm−1 for O2 plasma activated and UVO activated samples, respectively, also suggesting the superior crystal quality of ZnO on O2 plasma activated MWCNTs. Consequently, Raman characterization results also validate the hypothesis that surface activation of the MWCNTs is essential for effective functionalization, i.e., stronger nucleation and binding of the MONCs onto the MWCNT surfaces.

The G peak represents the movement in the opposite direction of two neighboring carbon atoms in a graphitic sheet, hence indicating the presence of crystalline graphitic carbon in MWCNTs, while the D peak represents the defects in the curved graphite sheet, sp3 carbon, or other impurities. The I(D)/I(G) ratio, where I corresponds to the peak area of the Lorentzian functions, is an estimate of the relative structural defects. Our preliminary characterization suggests that due to O2 plasma activation, the relative intensity of the D-peak with respect to the G-peak (I(D)/I(G)) ratio of the MWCNT increases 13.5%. The results are presented in the supplementary material. The probable reason for the increase in the intensity of the D-peak with respect to the G peak is the presence of reactive sites on the surface of the MWCNTs created by O2. These sites are supposed to

| Peak component | Location (eV) | Functional group | NPA |
|----------------|--------------|-----------------|-----|
| C2             | 285.4        | C-OH            | 0.29|
| C3             | 286.5        | C-O-C           | 0.26|
| C4             | 287.7        | C=O             | 0.03|
| C5             | 289.1        | COOH            | Not present |
| O1             | 531.1        | C=O             | 0.02|
| O2             | 532.2        | C-OH            | 0.20|
| O3             | 533.5        | C-O-C           | 0.77|
FIG. 3. (Color online) (a) TEM image of untreated but ZnO deposited MWCNTs. ZnO nanocrystals were not visible on the surface of these untreated MWCNTs. (b) Uniform distribution of atomic layer deposited ZnO nanocrystals on the UVO activated MWCNT surface. (c) High resolution TEM image of a SnO$_2$-MWCNT (ALD at 175 °C) sample showing interplanar spacings of 2.6 and 3.3 Å corresponding to (101) and (110) planes of SnO$_2$. (d) High resolution TEM image of a ZnO-MWCNT (ALD at 175 °C) sample showing interplanar spacings of 2.8, 2.68, and 2.48 Å corresponding to (100), (002), and (101) planes of ZnO.

FIG. 4. (Color online) (a) Raman spectra obtained from the O$_2$ plasma activated ZnO functionalized MWCNT (top panel), UVO activated ZnO functionalized MWCNT (middle panel), and untreated but ZnO functionalized MWCNT (bottom panel). (b) Enlarged ZnO peaks in the range of 100–850 cm$^{-1}$ frequency shift.
B. Methane sensing

Our hypothesis is that MONCs facilitate chemisorption of methane gas molecules on the surface of the activated MWCNTs. We found that the resistance of the MONC-MWCNT sensors increases in the presence of a mixture of methane in dry air. The relative resistance change has been defined as

\[ \Delta R / R = (R_{\text{methane}} - R_{\text{air}}) / R_{\text{air}} \]

A series of experiments were conducted to evaluate the effect of UVO and O₂ plasma treatments on the performance of the MONC-MWCNT sensor. To decouple the sensor response to CH₄ from the interference of variable RH, the RH inside the test chamber was kept constant and monitored in real time during each test period. The relative resistance of the sensor monotonically increased at room temperature when 10 ppm CH₄ in air was introduced to the test chamber. While maintaining a constant flow rate, when the incoming gas was switched from CH₄ to N₂, the relative resistance of the sensor decreased and returned to the baseline [Figs. 5(a)–5(c)]. The sensing mechanism could be elucidated from this phenomenon: the monotonic increase in the sensor’s relative resistance is a result of absorption of CH₄ molecules on the MONC functionalized MWCNT surface.

Figure 5 also corroborates the assumption that surface activation is essential for effective functionalization of the MWCNT by MONCs and for the sensor to act reversibly in the presence and absence of 10 ppm CH₄ in air. Figures 5(a) and 5(b) show reproducible changes in the relative resistance of the surface activated ZnO-MWCNT sensor during alternating exposure to CH₄ and N₂. Figure 5(c) illustrates a surface activated SnO₂-MWCNT chemoresistor sensor alternatively exposed to CH₄ and N₂ showing similarity to the ZnO-MWCNT results. No discernible signals were observed in the untreated (but ZnO NC deposited) MWCNT sensor [Fig. 5(d)].

At room temperature, the average relative resistance change \[ \Delta R / R = (R_{\text{methane}} - R_{\text{air}}) / R_{\text{air}} \] was found to be 1.91 ± 0.98% for UVO activated and 10.5 ± 1.01% for O₂ plasma activated ZnO-MWCNT sensors. The results show that the O₂ plasma activation significantly enhances the affinity of the MONCs (in this case ZnO NCs) to the MWCNT surface in comparison to UVO activation. This enhanced affinity causes stronger electron transport through the ZnO-MWCNT junction, i.e., a larger resistance change in the presence of CH₄ at room temperature [Figs. 5(a) and 5(b)]. This is likely due to the better crystal quality of the ZnO NCs on O₂ plasma activated MWCNTs compared to UVO activated MWCNTs (as also observed from the Raman results in Fig. 4).

A novel UV-based recovery technique was recently presented by our group. The sensor was first exposed to 10 ppm CH₄ in air for 30 min, and without interrupting the flow of CH₄, the sensor was irradiated with UV light until the sensor returned to its baseline resistance. No N₂ flow was used during the recovery. A recovery time of about 3 min was observed. The improvement in the recovery time, we believe, was due to the UV induced reduction of the desorption energy barrier of the CH₄ molecules at the sensor surface.

To verify the sensor response to methane, the ZnO-MWCNT sensor was tested at varying CH₄ concentrations (2, 5, and 10 ppm in dry air) at room temperature. Methane of 2 and 5 ppm was obtained from the dilution of 10 ppm methane in synthetic air. The sensors were exposed to CH₄ for 10 min (gas phase) and then to nitrogen for 10 min.

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**Fig. 5.** Comparison of the dynamic sensor response \[ \Delta R / R = (R_{\text{methane}} - R_{\text{air}}) / R_{\text{air}} \] of MONC-MWCNT sensors to 10 ppm CH₄ in dry air, followed by flushing with N₂ for sensor recovery. Each section of the figure shows \[ \Delta R / R \] for: (a) O₂ plasma treated ZnO functionalized, (b) a UVO treated ZnO functionalized, (c) a UVO treated SnO₂ functionalized, and (d) an untreated but ZnO deposited MWCNT sensor.
A possible sensing mechanism has been reported by MWCNT nanohybrid decreases in the presence of oxidizing resistance of the SnO2-MWCNT nanohybrid reported by Lu et al. The change in resistance is in accordance with the change in sensors changes in the presence of a mixture of methane in air.

The sensitivity of a semiconducting oxide gas sensor is defined as follows:5,6

\[ \Delta R / R = \left( R_{\text{gas}} - R_{\text{air}} \right) / R_{\text{air}} \]  

(1) for reducing gas

\[ \Delta R / R = \left( R_{\text{air}} - R_{\text{gas}} \right) / R_{\text{gas}} \]  

(2) for oxidizing gas

where \( R_{\text{air}} \) is the resistance of the sensor in air and \( R_{\text{gas}} \) is the resistance of the sensor in the presence of gas and air.

We found that the resistance of the MONC-MWCNT sensors changes in the presence of a mixture of methane in air. The change in resistance is in accordance with the change in resistance of the SnO2-MWCNT nanohybrid reported by Lu et al.,6 where they show that the resistance of the SnO2-MWCNT nanohybrid decreases in the presence of oxidizing NO2. A possible sensing mechanism has been reported by Lu et al.6 Target molecules, in this case NO2, get directly adsorbed onto the SnO2-MWCNT surface, facilitate electron transfer, and change the electrical conductivity of the hybrid nanostructure.

One can find references that describe the electronic properties of MWCNTs as metallic35,36 or semiconductive.6,37–39 However, ZnO and SnO2 are widely known as n-type materials,6,32 and the presence of a reducing gas, such as CH4, alters their charge concentration, resulting in a change in the resistance of the MONC-MWCNT conglomerate, which was observed experimentally.

A deeper investigation on the methane-functionalized CNT surface interaction is in order; however, this is beyond the scope of this paper. Furthermore, we are currently conducting experiments to study the electronic properties of surface pretreated metal oxide nanocrystal functionalized MWCNTs, which will help us to understand their methane gas sensing mechanism more thoroughly.

Understanding the effect of RH is important in estimating the outdoor performance of microfabricated gas sensors. To examine the effect of RH on sensor performance, the change in the baseline relative resistance of the surface activated ZnO-MWCNT sensors was measured at room temperature due to a change in RH. Humidity was provided by a controlled flow of moist air (flow rate: 0.94 l/min) into a plastic test chamber (residence time: 4.5 min). The baseline relative resistance of the sensor increased by about 4% as the RH was increased from 10% to 91% and returned back to the original baseline once the RH was reduced back to 10%. This suggests a strong electron transfer between the MONC functionalized MWCNTs and water molecules (Fig. 7). Our ongoing research involves fabricating a network of MWCNTs selectively functionalized with various metal oxide nanoparticles with different sets of sensitivities to CH4 and H2O. By deconvoluting the constructive/destructive interference at various RH levels, the RH contribution can be effectively determined and separated from the device’s response to CH4.

We explored the relative resistance change for 10 ppm methane at two different RH levels.36 While exposing the SnO2-MWCNT sensor to 10 ppm of CH4 in dry air at the higher RH (approximately 70%), a monotonic resistance increment was observed which was similar to the low RH tests where RH was held constant at 5%, as seen in Fig. 8. The sensor also equilibrated to its original response in a similar fashion when the chamber was purged with N2. The response to CH4 and signal-to-noise ratio reduced in comparison to those at lower RH, which we believe was a result of adsorbed H2O molecules on the SnO2-MWCNT sensor. Although the sensor showed a reduced response, it was still capable of detecting 10 ppm CH4 in air at 70% RH. We observed a monotonic increase in the sensor resistance, while it was exposed to CH4 at low and high RH, as well as a monotonic decrease (return to baseline) when CH4 was purged with N2 (Fig. 8). The sensor behavior is similar at both low and high humidities although the sensitivity is reduced at high RH. This is likely the result of adsorbed H2O molecules on the sensor surface. Water molecules are found to behave as electron donors on the surface of carbon.

![Fig. 6. (Color online) Relative resistance change of an O2 plasma activated ZnO functionalized MWCNT sensor under exposure to 2, 5, and 10 ppm of CH4 in the dry air mixture at room temperature. RH was kept constant at 2% during the experiment. A different metal electrode/CNT configuration was used, resulting in a smaller relative resistance change compared to Fig. 5(a).](image1)

![Fig. 7. (Color online) Relative resistance change \( \Delta R / R = (R_{\text{RH}} - R_{\text{air}}) / R_{\text{air}} \) of the ZnO-MWCNT sensor, while the relative humidity inside the chamber was varied by a controlled flow of moist air. The right hand y-axis represents the RH inside the chamber during the test.](image2)
It was reported that a hydrogen-bonded water monolayer forms around the nanotube at a fully water covered condition. Na et al. presented the change in electrical resistance as a function of relative humidity, which agrees with the result presented in Fig. 8, i.e., a decrease in the relative change in resistance ($\Delta R/R$) at a high RH condition. This can be attributed to electron donation by the H$_2$O molecules on the sensor surface.  

Our ongoing work focuses on studying the response of the MONC-MWCNT sensor at a fixed ppm methane for multiple RH%.

The CH$_4$ sensor described here uses MWCNTs functionalized by MONCs to sense methane. We used two well known MONCs, ZnO and SnO$_2$, that are widely used methane sensing materials and are inexpensive. ZnO and SnO$_2$ promote energetically favorable electron transport at the MO–MWCNT junction. The work function of ZnO was reported to be almost 4.64 (Ref. 43) or 5.2 eV, while SnO$_2$ has a work function of 4.7 eV. The work functions of these MONCs are almost equal to the work function of MWCNTs (4.7–4.9 eV). Therefore, the Schottky barrier height at the MONC-MWCNT junction is low, facilitating electron transfer between MWCNTs and MONCs. The low Schottky barrier improves the overall sensitivity of the sensor (i.e., high $\Delta R/R$ at low ppm), making the hybrid MONC-MWCNT system a potentially superior sensing element to either of its constituent components.

Table II compares the performance of our sensor with other published CNT CH$_4$ sensors. Note that although the sensor presented in Ref. 46 shows an equivalent performance to our sensors, it used wet chemically treated single walled carbon nanotubes (SWCNTs). Wet chemical treatment may be undesirable in CNT sensor fabrication as it is well known that acid treatments used in wet chemistry can considerably reduce the mechanical and electric performance of the tubes by introducing large numbers of defects, which might limit its reproducibility and reliability, as well as increase the cost of the sensor. Wet chemistry also involves additional steps, such as dissolution, sonication, mixing, and drying, which often causes undesirable agglomeration of treated CNTs.

IV. CONCLUSION

In summary, O$_2$ plasma activation has a stronger impact than UVO activation on enhancing the MONC functionalization of MWCNTs and thus on the response of the chemoresistive sensors to 10 ppm CH$_4$ in air. The strong relative resistance change in the presence of 10 ppm of CH$_4$ at room temperature is a consequence of: (1) strong electron transfer to the MONCs from CH$_4$ molecules, (2) energetically favorable electron transport at the MONC-MWCNT junction, and (3) enhanced affinity of the dry surface activated MWCNT to MONCs as a result of formation of active chemical groups. The O$_2$ plasma and UVO-based activation processes give rise to COOH, C=O, and C-OH functional groups on the MWCNT surface and hence enhance the nucleation and bonding of MONCs with the MWCNT. These treatments produce a strong reversible relative resistance change in the chemoresistors under iterative exposure to 10 ppm CH$_4$ in air and a relatively reduced response to lower concentrations. The response varies with RH, with a lower response at high RH%.

Table II. Comparison of CNT chemoresistor sensors.

| Material | Lowest detection limit | Operating temperature | Interference gases |
|----------|------------------------|-----------------------|-------------------|
| Pd-SWCNT (Ref. 5) | 15 ppm | Room temperature (RT) | Not mentioned (NM) |
| Pd-MWCNT (Ref. 14) | 3 vol. % | RT | H$_2$, NH$_3$ |
| Pd-MWCNT (Ref. 11) | 2 vol. % | RT | NM |
| Carbon nanofiber (Ref. 47) | 500 ppm | RT | NM |
| Pt-CNT, Ru-CNT, Ag-CNT (Ref. 48) | 0.7 vol. % | 150°C | CO$_2$, CO, NO$_2$, NH$_3$ |
| SWCNT (wet chemical treated) (Ref. 46) | 2 ppm | RT | CO, SO$_2$, NH$_3$ |
| ZnO-MWCNT and SnO$_2$-MWCNT (This work) | 2 ppm | RT | H$_2$O |
higher RH as well as a lower detection limit. At low RH, the detection limit is between 2 and 5 ppm.

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