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Multifunctional Chemical Vapor Sensors of Aligned Carbon Nanotube and Polymer Composites

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With the recent developments in nanoscience and nanotechnology, there is a pressing need for flexible, mechanically robust, and environmentally stable chemical vapor sensors with a high efficiency and low power consumption. This type of sensors can, for example, be used for real-time sensing of chemical warfare stimulants in a battlefield by monitoring the resistance changes in soldiers' clothing.1 The early measurements on the conductivity changes of conjugated (conducting) polymers via charge transfer with certain chemical vapors or nonconducting polymers mixed with conductive fillers through polymer swelling by gas absorption around the percolation threshold provided the basis for developing polymer-based chemical vapor sensors.¹⁻³ However, the poor environmental stability associated with most conjugated polymers limited the scope of their use for practical applications.¹ The uncertainty on the precise location of the percolation threshold in random dispersion systems remained as one of the major hindrances toward high-performance conductively filled polymer sensors.¹⁻³

The recent discovery of the gas sensing capabilities of carbon nanotubes through the charge transfer or capacitance change by gas absorption (e.g., NH₃, NO₂, O₂)⁴⁻⁶ is significantly intriguing as their small size, high surface area, good environmental stability, and excellent mechanical and electronic properties can offer many advantages for sensing applications.^{1,7} The use of the pristine nonaligned carbon nanotubes for gas sensing as reported in these studies often involves tedious processes for integrating single carbon nanotubes into sensor devices, and the number of analytes to be determined is also hampered by the limited specific interactions with the unmodified nanotubes. It will be a significant advancement if we can use *perpendicularly aligned* carbon nanotube arrays as the sensing materials. Aligned carbon nanotube arrays, in either a patterned or nonpatterned form,8 allow the development of novel sensors and/or sensor chips without the need for direct manipulation of individual nanotubes since the constituent nanotubes can be collectively addressed through a common substrate/electrode.^{8,9} The aligned nanotube structure further provides a large well-defined surface area and the capacity for modifying the carbon nanotube surface with various transduction materials^{8,9} to effectively enhance the sensitivity and to broaden the scope of analytes to be detected. Here, we report a novel concept for developing a new class of multifunctional chemical vapor sensors with a low power consumption, high sensitivity, good selectivity, and excellent environmental stability by partially coating perpendicularly aligned carbon nanotube arrays with appropriate flexible polymer films.

As schematically shown in Figure 1, the aligned multiwall carbon nanotubes produced by pyrolysis of iron(II) phthalocyanine (Figure 1a)^{8,10} were partially covered with a polymer coating top-down along their tube length (Figure 1a (I)) by depositing a droplet of polymer solution (e.g., poly(vinyl acetate), PVAc, polyisoprene,

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Figure 1. Schematic illustration of the procedures for (a) fabricating and (b) characterizing the aligned carbon nanotube—polymer composite chemical vapor sensor. SEM images of the aligned carbon nanotube array (c) before and (d) after being partially coated with a polymer (PVAc) film on top and turned upside down (cf. Supporting Information; scale bars = 5 μ m).

PI) onto the nanotube film (cf. Supporting Information). The composite film was then inverted as a free-standing film (Figure 1a (II)) for sputter-coating two strip electrodes of gold across the nanotube arrays that were protruding from the polymer matrix (Figure 1a (III)). The flexible thin film device can then be used for chemical vapor sensing through monitoring conductivity changes (Figure 1b) caused by the charge-transfer interaction with gas molecules (*without the need for direct manipulation of individual nanotubes*) and/or the inter-tube distance change induced by polymer swelling via gas absorption (*without the requirement to be operated around the percolation threshold*). SEM images c and d of Figure 1 show the aligned carbon nanotube array before and after being partially filled with PVAc, respectively.

As seen in Figure 2a, the as-synthesized aligned carbon nanotube arrays without the polymer coating do not show any obvious resistance change while exposed to various chemical vapors, nor does the pure polymer film. In contrast, Figure 2b clearly shows $\sim 130\%$ increase in the resistance change, ΔR , for a composite film of PVAc and aligned carbon nanotubes after being exposed to tetrahydrofuran (THF) vapor for several minutes. Subsequent removal of the THF vapor source caused a loss of ca. 100% in the resistance change after keeping the nanotube-PVAc composite film in air at room temperature for 2 min. It is apparent that the observed resistance changes are due to expansion of the polymer matrix upon exposure to the chemical vapor, which leads to a concomitant increase in the inter-tube distance (Figure 1b). The lost conductivity could not be completely compensated in this particular case, even after drying the nanotube-PVAc composite film in air for a prolonged period, due to the inevitable presence of some residual good solvent (i.e., THF) in the PVAc matrix. After the first vapor-air cycle with a 2 min interval, the maximum resistance and the peak height became constant (Figure 2b), indicating that the solvent absorption/ desorption had reached an equilibrium state.

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