Synthesis of carbon nanotubes by CVD over Fe catalyst prepared in spin coating on SiO$_2$/p-Si(100)

Ahmed N. Abd

Department of physics, college of Science, University of Al-Mustansiriyah. Baghdad-Iraq

Email address: ahmed_naji_abd@yahoo.com

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Abstract. Carbon nanotubes (CNTs) have been grown by the decomposition of C$_2$H$_2$ over thin catalyst film in order to investigate the growth mechanism of CNTs by chemical vapor deposition (CVD). The catalyst was prepared from an iron chloride FeCl$_3$ precursor solution that was spin – coated on SiO$_2$ substrate. The growth mechanism and the length and the diameter of the CNTs were greatly influenced by the precursor concentration, the time deposition and the temperature and the ratio of C$_2$H$_2$:N$_2$ are fixed. Transmittance electron microscopy and Scanning electron microscopy and Raman spectroscopy and X-ray diffraction have been carried out in order to investigate the behavior of the growth process. The iron chloride film contact with the C$_2$H$_2$:N$_2$ reaction atmosphere.

1. INTRODUCTION

Synthesis of carbon nanotubes (CNTs) has been extensively investigated by a number of research groups, since the first observation in 1991 [1]. Carbon nanotubes (CNTs) have been studied intensively due to their importance as building blocks in nanotechnology since they were discovered. Their special geometry and unique properties offer great potential applications, including nanoanical transistors [2], energy storage [3], field emission devices [4], gas sensors [5], CNTs show great promise for future electronic applications, [6], CNTs are expected to find applications in all industrial areas, also provide rich research subjects. CNTs can be produced by different techniques; among them most widely used ones are arc- discharge [7], laser ablation [8], and chemical vapor deposition (CVD) [9]. We chose chemical vapor deposition (CVD) technique which has some distinct advantages over other techniques. The CNTs can grow at relatively low temperature, and the size of CNT can be controlled by varying the size of the metal catalyst. Furthermore, it is relatively easy to obtain vertically aligned CNTs. However, the process parameter control for the growth of CNT still remains in an empirical manner, because the reaction kinetics and the growth mechanism are not fully understood. Behavior of CNT in CVD process can be compared with that of the carbon filaments [10].

2. EXPERIMENTAL

2.1. Description of the CVD apparatus

The CVD apparatus used in our experiments is basically composed of a quartz glass tube with diameter 3.5cm, and tube furnace with diameter 4.5cm and bulbar show in fig.1
2.2. Synthesis of nanotubes

The deposition of nanotubes was carried out in four steps:

a. Silicon (1×1×0.05)cm substrate(100)/p-type with resistivity of 15(Ω.cm) was used. The substrate was cleaned in acetone 99.99 purity 100 ml inside and put in the digital ultrasonic at 40 °C for 10 min with deionised water, ultrasonic with alcohol 99.99 purity for 5min ,washing with deionised water ,last step substrate were then dried with dry nitrogen. After cleaning substrate put it into a reactor, we have allowed the temperature increase to about 900 °C, for removing again the remaining dirtiness. In this stage we inject pure oxygen in reactor for about 20 min, Later we turn off furnace and cooling the substrate. The thickness of the SiO$_2$ layer was estimated at approximately 300nm.

b. Preparation the catalyst nanoparticles we used spin coating method to deposition films on the silicon substrate. Solutions of concentration (15, 30, 45, 60, 75 and 90) mM were sonicated for 30 min before placed onto substrate. It was found that FeCl$_3$ has better solubility in ethanol. The solution was dropped on p-Si(100)/SiO$_2$ substrate in number of circular 1000 r/min for each once with spin coater at room temperature. The thickness of the deposited films can be changed with varying concentrations of FeCl$_3$.

c. The coated substrate was carried out into quartz tube and introduces N$_2$ high purity 99.99 with flow rate 85 sccm in tube after this heating the tube with heat rate 20 °C/min until to reach the final temperature at 750 °C. Stay in this temperature 30 min to forming Iron oxide clusters.

d. The synthesis of nanotubes was performed by adding a constant flow rate of acetylene (15 sccm) to the nitrogen carrier gas during 15 min. After this step the acetylene switch off and the N$_2$ continuously in flow until to cooling in room temperature.

3. RESULTS AND DISCUSSION

The images of AFM in fig. (2) Show a good coverage of catalyst on P/100/SiO$_2$ and an obvious dependence of size and density of nanoparticles on the surface. The average roughness and the average grain size is increase with increase concentration of the solution. The average roughness increase that beginning from 0.039 nm and end in 4.11nm and average grain size beginning from 4.37nm and end in 8.28nm for (15-90) mM. This must mean that there are some other parameters that are important in the dispersion. The distribution of clusters is strongly dependent on the type of salt, support and the distance from the center of the support to the edge [11]. The most obvious is the quantity of liquid dropped on the surface which will affect the quantity of iron molecules per unit area.
Figure 2: Show the catalyst at concentration A. 15mM and B. 30mM.

Figure 3: Show SEM images of CNTs grown on the catalyst particles formed on the p(100)/SiO$_2$. We show that the nanotubes are oriented perpendicularly and helical shapes on the substrate surface for high densities of nucleation centers. A high nanotube density can be achieved with a high density of iron oxide clusters that are nucleation centers for the growth of nanotubes. The nanotubes synthesized by this method have diameters about between 20 and 80 nm. The highest nanotube densities and film thicknesses of about 50µm were observed with the lowest iron chloride concentration at high temperatures 750°C [12].

Figure 3: Show the growth of CNTs.

The TEM images shown in Figure 4 confirmed that the carbon structures were multiwalled carbon nanotubes, which hollow cavity, diameter about 80nm, and revealed the presence of iron particles mainly in the tip of MWCNTs, suggesting that a tip-growth mechanism might be predominant. The CNTs were not complete graphitization of the walls visible in the TEM imaged the presence of small amounts of amorphous carbon was confirmed by Raman spectroscopy and XRD analysis. The layers of CNTs, which are not clear separated probably caused by more defective growth [13]. Temperature annealing, up to 2200°C, has demonstrated benefits on mechanical and physical properties. These treatments are increasing in the crystallite size, corresponding to an increase in the graphitic domains. Some prior literature exists in high temperature heat treatment of nanotube-like materials described the graphilization of entangled, sinuous nanotubes at 2800°C [14].
An XRD measurement (2θ scan) was carried out using Cu Kα radiation to examine the structure of the CNTs. Fig.5.A. show three clear characteristic peaks of the graphite structure in MWNTs at 26.2°, 43.6° and 50.7° [15]. Two graphite peaks corresponding to the (002) and (100) planes at 2θ = 25.8°, 43.5° can be clearly seen and the broad diffraction peak of (002) graphite planes at 2θ = 25° gradually increases with increasing annealing temperature [16]. Fig 5.B. show the wider peak of 002 centered at 24.97° is induced by the disordered carbon and indicates to presence defect level in the atomic carbon structure. The significant diffraction pattern of the MWCNTs non pure function is appearing at and near 2θ of 25.3° [17]. The peaks at 43.2° for 100 plane indication of the low quality of carbon nano materials [18]. The peaks at 52.53°, 54.16°, and 55.36° are not originated from the CNTs. It should be noted that the peaks between 2θ = 50° and 60° are from the Si substrate [18]. The crystalline size of CNTs estimated from an X-ray diffraction peak using Scherrer’s equation (d = 0.9λ/ B cos Θ) where d is the size of crystals, B is the full width at half maximum of a peak in radians at the diffraction angle Θ and λ is the X-ray wavelength, which is ≈ 0.15418 nm in this study (CuKα) [18].

The Raman spectra of MWCNTs were non-functionalised are shown in Figure 6. The spectrum of carbon structure contains two main bands: the G-band (1596 cm⁻¹ MWCNT), which is assigned to the E₂g C–C (sp² - bonded) stretching mode of a well-ordered graphitic structure and the D-band.
(1306 cm\(^{-1}\) for MWCNT), attributed to the \(A_{1g}\) (sp\(^3\) -bonded) stretching mode, resulting from the presence of a disordered structure or lattice defects in the graphite structure (substitutional hetero-atoms, vacancies or chemically bonded hetero-atoms). The bands at 2604 cm\(^{-1}\) (G' band) for MWCNTs constitute an overtone of the D band. Due to such a highly defective MWCNT structure. At the second order mode observed between 2604 cm\(^{-1}\) assigned to the first overtone of the D mode (2D) and often called G’ mode, which is highly dispersive. The G’-band feature (operative for the graphite) is based on double resonance mechanism. The D-band and G’-band show, for graphite or nanotubes, a strong dispersion as a function of laser excitation energy (their frequency are sensitive to the excitation wavelength), while the G-band is not dispersive. The peak 454cm\(^{-1}\) and the broad peaks around (300-700) cm\(^{-1}\) belong for \(\text{Fe}_2\text{O}_3\) and other types of iron oxide nanoparticles.

![Figure 6: Raman spectrum of MWCNT/ Fe\(_2\)O\(_3\) nanostructure.](image)

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4. CONCLUSION

The density of catalyst nanoparticles deposited depended mainly on salt concentration and speed of spin coating. The density of particles decreased away from the center to the edge of the wafer. The diameter of the carbon nanotubes was correlated to the size of clusters deposited. A higher catalyst loading also produces MWNTs at 750 °C. A Raman peaks and XRD indicate the presence of some form of carbon because non availability pure function of MWCNTs. This result emphasizes the need to maintain on the purification CNTs in the study.
References


