Effect of Template on the Structure of Carbon Nanotubes Grown by Catalytic Chemical Vapor Deposition Method

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Abstract
Carbon nanotubes (CNTs) have been synthesized on an anodized aluminum oxide (AAO) template with acetylene by using catalytic chemical vapor deposition (CVD). It was found that the structure of CNTs can be mostly depends on the quality of the catalyst deposition in the pores of AAO template. Straight CNTs were observed when Ni catalyst deposited only in the bottom of holes as nanoparticles, but when Ni catalyst fulfilled the holes of AAO template as nanowires (NWs), coiled CNTs observed. The characterization of as-prepared materials was examined by SEM and Raman spectroscopy. In addition, it is realized that the nanostructure of the AAO template strongly affected the properties of as-grown CNTs. In the other word, based on the obtained results the diameter and pitch of coiled CNTs tightly depend on the size of metallic catalyst and in the results it depend on the pore diameter of AAO template.

Keywords: Nanostructures; Chemical vapor deposition; Electron microscopy; Raman spectroscopy

Introduction
Due to their unique mechanical, electrical, and thermal properties, carbon nanotubes (CNTs) have attracted particular attention among the known allotropes of carbon [1-4]. Although straight CNTs, including single walled CNTs (SWCNTs) and multi walled CNTs (MWCNTs) were commonly observed, different shapes of CNTs such as coiled, branched, and toroidal were reported [5] shortly after their discoverying by Iijima [6]. It has been demonstrated that the nanostructure of CNTs can strongly affected the characteristic properties of different shapes of CNTs, so it can limit their applications. For example, unlike straight CNTs, coiled ones can show semi metallic properties beside the typical electrical characteristics of straight CNTs, such as exhibition metallic and semi conductive properties [7,8]. Almost simultaneously with claim that there is possibility about the existence of coiled CNTs with minimized energy confirmed by theoretical calculations [9-11] coiled carbon fibers were experimentally synthesized, principally from decomposition of hydrocarbons in the presence of metallic catalyst [12-18]. It is known that coiled CNTs are MWCNTs with incomplete crystalline structures created when paired pentagon-heptagon atomic rings arrange themselves periodically within the hexagonal carbon network. We have previously reported production of a novel porous coiled CNT/Ni-NW nanostructure by ambient thermal-CVD [19] successfully. Our investigations show that this material being a sufficient structure for improvement of electrode materials for advanced energy storage systems. On the other hand, there is a great interest for growing well aligned CNTs and specially vertically aligned CNTs (VA-CNTs) for developing field emission devices and a large variety of microelectronic devices, including nanotube sensors, optoelectronic systems, batteries, and supercapacitors [20-23]. One of the common methods for growing VA-CNTs is based on using a porous filter such as AAO template. By using this kind of nano template for growing CNTs, control the shape of CNTs being easy. So, the characteristic properties of CNTs especially their aspect ratios are tightly control with the nanostructure of template, including diameter and length of nano channels. Ever since the discovery of carbon nanotubes, a variety of methods have been reported for producing different shapes of CNTs, among which may be mentioned the following main methods: arc discharge, laser ablation, and CVD [21]. In the last few years, CVD method attracted much attention as the most utilized method, because it is a cost effective method for large-scale production of CNTs on a wide range of substrates. Recently, CVD method is given special emphasis as the most used method for synthesis different shapes of CNTs, specially coiled and aligned straight ones [22-24]. In this study, acetylene decomposition was performed over Ni catalysts supported on AAO template to investigate the role of catalyst on the structure of as-grown CNTs. To evaluate the effect of nanostructure of AAO template on the appearance of as grown CNTs, different condition of anodizing template, were investigated.

Experimental
Chemicals
High-purity aluminum foil (99.9995%, Merck, Germany), Perchloric acid (60%, Merck, Germany), ethanol (96%, Jonoob, Iran), oxalic acid 2-hydrate (99%, Panreac Quimica SA, E.U.), Choromic anhydride (KANTO Chemical Co. INC, Japan), phosphoric acid (85%, Merck, Germany), copper sulfate (Merck, Germany), chloridric acid (36%, Merck, Germany), hexahydrate nickel sulfate (Merck, Germany), heptahydrate nickel chloride (Merck, Germany), boric acid (Merck, Germany) were used as received. Finally, double ionized water was obtained from an OES water purification system (Oklahoma, USA).

Equipments
DC power supply (MP6003, Megatek, Germany) was used in order to apply electric field in preparation of AAO template. Electro analyzer system SAMA 500 (Iran) was used for electrochemical deposition of Ni catalystic particles in the pores of AAO template. The samples investigation were done by using scanning electron microscopy (TESCAN, VEGA, Czech Republic), field- emission scanning electron microscope (TESCAN, Mira II and 3 LMU, Czech Republic) and Raman spectroscopy (BRUKER, SENTERA, Germany).

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Preparation of template and growing CNTs

As illustrated in Figure 1, a porous AAO template was obtained by two-step anodization of a 3 × 3 × 0.3 mm³ high purity (99.9995%, Merck, Germany) aluminum foil which degreased with acetone by using ultrasonication and then annealed at 500°C for 5 hr. Before proceeding Aluminum by anodizing the sample must pretreated using electropolishing for smoothing and brightening the surface of metal. In first step of anodization, the sample was anodized in an aqueous solution of oxalic acid (0.3 M) under a DC constant voltage (45 V) at low temperature (below 5°C) for 20, 22, and 25 hr. After the first step of anodization, the anodized surface layer was removed by electrochemical etching step which occurred in a mixture of chromic acid (1.8 wt %) and phosphoric acid (6 wt %) at 75°C for 3 hr. In the second step, anodization was done for 4 hr under the same conditions as mentioned in the first step. In order to facilitate the uniform electrodeposition of Ni which acts as catalyst in growing CNTs, immediately at the end of the second anodization, the voltage was dropped from 45 to 14 V at the rate of 0.5 V min⁻¹. The remaining Al substrate was mostly removed in a saturated copper sulphate aqueous solution in chloridric acid at room temperature [23]. Finally, the AAO template was etched in a 1 M aqueous solution of phosphoric acid at a room temperature for 40 min for the further thin the barrier layer and widen the pore size. Catalytic Ni particles deposited in the nanochannels of as prepared AAO template from Watt bath which contained from hexahydrate nickel sulphate (330 g l⁻¹), heptahydrate nickel chloride (45 g l⁻¹) and boric acid (35 g l⁻¹) at 23°C and pH=2. The electrochemical deposition of Ni-nanoparticles in the pores of AAO template was performed by using a Ni deposition template which sputtered with a layer of Au, as working electrode, platinum as counter electrode, and Ag/AgCl as reference electrode, and all of the potentials refer to the reference electrode. Ni nanoparticles were electrochemically deposited in nanochannels of AAO template applying at scan rate of 50 mV s⁻¹ in the watt bat contained NiSO₄·6H₂O, H₃BO₃, and NiCl₂·7H₂O. The deposition time was selected by considering the deposition rate of Ni nanoparticles which deposited in the pores of AAO template, by adjusting the deposition time. The deposition of Ni nanoparticles were carried out in the potential range between -1.1 to 0 V applying 5 cycles for deposition Ni nanoparticles and 60 cycles for growing Ni nanowires (Ni-NWs) which fulfilled the holes of AAO template. Along with fulfilling pores with Ni-NWs, the color of surface changes to silver, so the Ni deposition must be stopped after the first change of color. Figure 2 shows the cyclic voltammograms recorded for deposition Ni nano catalysts in the pores of AAO template at 23°C and pH=2. CNTs were grown by using a catalytic pyrolysis of acetylene as hydrocarbon source and a mixture of hydrogen and argon as carrier gas. In the first step, catalytic Ni particles were reduced in a carrier gas (contains 10% H₂ and 90% Ar) at 500°C for 1 hr. In the second step, which started immediately after heating up to 650°C, CNTs were grown by catalytic decomposition of 10% C₂H₂ and 20% H₂ in an Ar carrier gas at a total flow rate of 150 sccm during 40 min. After growing CNTs, the furnace stay at 650°C for 10 min without any hydrocarbon source, then the furnace was cooled down slowly in presence of Ar. To have free standing VA-CNTs, surrounding AAO template was removed by using KOH solution in room temperature Figures 3a and 3b illustrate the preparation steps of growing VA-CNTs and coiled CNTs by using the CVD method, respectively. Our previous studies show that coiled carbon nanotubes grow directly on an anodized aluminum oxide (AAO) template which one side of it is covered with conducting Au layer and the other side is filled by using catalytic Ni nanowires.

Results and Discussion

Influence of the nanostructure of catalytic Ni particles on the structure of CNTs

The Figure 4 show the SEM images of two AAO based templates after deposition of Ni nano catalysts in their pores as nanoparticles and NWs, respectively. As it can be seen in Figure 4a, the catalyst particles deposited uniformly at the bottom of pores and in Figure 4b, the catalytic Ni-NWs were deposited in the pores of AAO template and filled the holes. Figures 4c and 4d show SEM images of as-grown straight VA-CNTs on the supported Ni nanoparticles by acetylene decomposition at 650°C before and after removing AAO nano template by using KOH aqueous solution in room temperature, respectively. Cross sectional and plane views of AAO/Ni-NW based template after growing coiled CNTs on the Ni-NWs were illustrated in Figures 4e and 4f, respectively. Raman spectroscopy is a method used to obtain suitable information about the structure of as-prepared two kinds of carbon nanotubes. Figure 5 represents the Raman spectra of grown CNTs on the AAO template using Ni nanoparticles and Ni-NWs as catalysts. The multi-walled structure of grown CNTs is identified by the sharp G bands which appeared at 1500-1600 cm⁻¹ region of the wave number and attributed to the graphite E'g optical mode and also by the one observed at the wave number of 1200-1300 cm⁻¹ is known as D bands which corresponds to the disordered sp² carbon of graphite [26]. The ID/IG ratio of the as-grown coiled and straight CNTs is ~1.45 and ~0.865, respectively. It reveals that the crystallinity of the CNTs grown using Ni nanoparticles as catalyst is higher than that of coiled CNTs grown using Ni-NWs, and it should be noted that increasing the deposition time of Ni nanoparticles to fill the pores of AAO template can increase the possibility of creating pentagon-heptagon atomic rings within the hexagonal carbon network, which are known as defects [27,28]. Thermogravimetric analysis (TGA) was performed to precisely determine the content of coiled CNTs in the coiled CNT/Ni-NW electrode. The results of this analysis has previously been presented by using our group.

Influence of the pore diameter of AAO template on the nanostructure of coil CNTs

In the CVD process, the size of nano pores which were fulfilled by Ni-NWs play an important role for growing coiled CNTs. SEM observations indicated that the diameter and pitch of as-grown coiled CNTs (Figure 6) were strongly affected by the size of metallic nano catalysts, which was controlled significantly by the pore size.
Figure 2: The Ni nano catalysts deposited in the pores of AAO template in Watt bath in scan rate of 50 mV s\(^{-1}\).

Figure 3: A schematic illustration of growing (a) VA-CNTs and (b) coiled CNTs, by using a catalytic CVD method.

Figure 4: SEM images showing (a) AAO template after deposition Ni nanoparticles in the bottom of pores, (b) and (c) CNTs grown at 650°C for 40 min in nanochannels of AAO template before and after removing AAO template, respectively, (d) Ni-NWs deposited in the nanoholes of AAO template, (e) side view and surface morphology of AAO/Ni nanowire based template after growing coiled CNTs.
of AAO template [29]. As it can be seen in Figure 6 diameter, and pitch of coil CNTs were increased by increasing the pore size of AAO template. The insets of Figures 6a and 6c show the surface morphology of AAO templates. By increasing the diameter of pores and also the diameter of Ni-NWs from ~50 to ~80 nm, the diameter of coiled CNTs were increased from ~100 to ~400 nm, respectively. Based on the above results we can claim that the diameter and pitch of coiled CNTs are tightly depend on the pore diameter of AAO template [30].

Conclusion

In summary, two different types of CNTs were successfully fabricated on an AAO template by using CVD method. This study reveals that the structure of metallic catalyst and the size of nanoparticles of metallic catalyst are important parameters in growing CNTs which can strongly affect the microstructure of MWCNTs and also their alignment on the substrate. On the other hand, the diameter and pitch of coil CNTs can significantly controlled by the diameter of nano pores of AAO template. Therefore, results from this investigation can provide additional insight into control of the shape of CNTs and their application in various industrials.

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References


