Synthesis and Characterization of Carbon Nanospheres

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Abstract

An experiment was conducted to synthesize carbon nanospheres of different sizes by controlling the temperature in polycondensation reactions of glucose under hydrothermal processes. Morphological, structural, thermal, and optical properties of as-synthesized particles were characterized using a transmission electron microscope (TEM), scanning electron microscope (SEM), X-ray diffractometer (XRD), thermal gravimetric analysis (TGA), differential thermal analysis (DTA), and Raman spectroscopy. Thermal study indicates that particles change their phase from amorphous to crystalline before achieving a thermal stability at 520 ˚C. The presence of D-band and G-band in the Raman spectrum also verifies amorphous and crystalline natures of the sample. It has also been observed that the particle’s morphology, dispersity, and size were modified after calcination of samples at 500 ˚C in comparison to those not calcinated. Overall results indicate that the carbon nanospheres are hard solids and highly dispersed with size ranges from 50 nm to 260 nm.

Subject Areas
Nanometer Materials

Keywords
Carbon Nanosphere, X-Ray Diffraction, TGA

1. Introduction

Carbon, a life forming element on earth, has ability to bond to itself. This property can be used to produce a myriad of structures that can be exploited in medical and energy applications. Carbon has many natural structures such as soot, amorphous structure, diamond, a crystalline structure, and graphite, a honeycomb lattice structure. It is possible to synthesize carbon of such structures
with different shape and size by mere control of carbon containing reactants and experimental conditions. The past two decades have witnessed a number of breakthroughs in carbon research that include fullerenes, graphene, and nanotubes. With the continuing technological developments and understanding the properties of carbon, scientists are able to synthesize verities of carbon containing materials from the micrometer to nanometer range [1]-[8]. The shape and size of carbon nanomaterials can easily be tuned during the growth process to produce carbon nanoparticles in the form of nanotubes, nanoflakes, nanoplates, nanorods, nanoballs, nanoonions, nanospheres, and many more. Among all these shapes and sizes, carbon nanospheres (CNs) show potential applications in optoelectronics, medicines, gas storage and sensors, heterogeneous catalysts, and encapsulating electrodes [1] [4] [9] [10].

Carbon nanospheres (CNs) are dispersible in water, and have shown high efficiency to breach the nuclear membrane while transporting drug molecules into the cell. The CNs are also intrinsically fluorescent and can be tracked easily inside cells without using any fluorescent tags [11]. One of the major problems in the synthesis of carbon nanospheres is their agglomeration in nanorange. Hence, we have used a developed synthesis technique to fine tune the intrinsic properties of CNs by changing their size and crystallinity.

2. Experiment

Various synthesis processes have been developed for carbon nanomaterials such as chemical vapor deposition (CVD), polymer pyrolysis, and hydrothermal treatment. Here we have used a hydrothermal (HTs) treatment to control the shape and size of CNs. We aim to tune the size and dispersion of CNs by controlling the growth parameters in HTs processes. The controlled size, monodispersity, chemical composition, and microscopic structure of a synthesized product can improve the performance of many devices like electrodes, photothermal materials, drug delivery carriers, and so on. Size of CNs in HTs processes can be controlled by adjusting the concentration of carbohydrate (glucose) solution, reaction time, and/or temperature. The synthesis process may be regarded as “green-chemistry,” as it uses no toxic solvents or surfactants [3] [12] [13] [14] [15].

We have synthesized carbon nanospheres by polycondensation reactions of glucose under hydrothermal conditions. Fifty mL of 0.5 M aqueous solution of glucose was kept in a 100 mL Teflon lined autoclave and heated for 6 hours at 140˚C to 190˚C. The autoclave was cooled to room temperature before centrifuging the brown products with water. Finally, the product was rinsed in ethanol and dried at 60˚C for 5 hours. The morphological, structural, thermal, and optical properties of as-synthesized particles were characterized using a transmission electron microscope (TEM), scanning electron microscope (SEM), X-ray diffractometer (XRD), thermal gravimetric analysis (TGA), diffraction thermal analysis (DTA), and Raman spectroscopy.
3. Results and Discussion

The TEM images indicate that the carbon spheres are hard solids and are highly dispersed with size ranges from 50 nm to 260 nm on average. The samples which were calcinated at 500 °C after a synthesis process have shown an improvement in surface morphology, reduction in size, and enhancement in dispersion compared to those which were not heat treated. The TEM images (a, b, c) in Figure 1 represent CNs synthesized at 170 °C, 180 °C, and 190 °C, respectively for 6 hours in hydrothermal pot. The TEM images (a, b, c) in Figure 2 were calcinated at 500 °C in nitrogen atmosphere after the synthesis process. In Figure 3, TEM image (a) represents CNs synthesized at 160 °C, SEM image (b) represents CNs synthesized at 140 °C, and a histogram (c) represents size distribution of as synthesized carbon nanospheres before and after calcination. A node-like structure on the surface of a particle in Figure 3(a) may be responsible for interconnection of carbon spheres. The SEM images shown in Figure 3(b) possess highly porous hollow particles of sponge-like surface indicating immature particles with the size range in micrometer. This sample was grown at 140 °C for 6 hours.

![Figure 1](image1.png)

**Figure 1.** TEM images for as synthesized CNs for 6 hours, images (a), (b), and (c) represent the CNs synthesized at temperature 170 °C, 180 °C, and 190 °C, respectively.

![Figure 2](image2.png)

**Figure 2.** TEM images for CNs calcinated at 500 °C, images (a), (b), and (c) represent the CNs synthesized for 6 hours at temperature of 170 °C, 180 °C, and 190 °C, respectively.
Figure 3. Image (a) is the TEM of CNs synthesized at 160˚C for 6 hours, image (b) and (c) are the SEM of CNs synthesized at 140˚C for 6 hours, and that at 170˚C for 15 hours, respectively.

The surface morphology and porosity of spheres seem to form “a pool of thread” while growing at elevated pressure in the polycondensation reaction. In polycondensation reactions, carbon containing materials may have formed a long polymer chain which would curl up around a “seed” to form carbon spheres until the conditions are favorable. As this temperature was not favorable to grow CNs for 6 hour durations, we have continued our experiment from 160˚C to 190˚C. However, the particle grown at 170˚C for 15 hours has a good surface morphology but extra-large size which is not of our interest at this time as shown in Figure 3(c). The histogram in Figure 4(b) depicts that 170˚C was the optimal condition to grow carbon nanospheres in our experimental setup.

The X-ray diffraction patterns of the sample contains a strong peak at 11.6˚ and weak peak at 22.2˚ associated to (002) as shown in Figure 4(a) and (110) reflection planes in a tetragonal arrangement of carbon atoms. The XRD was taken with Cu-Kα (λ = 1.5406 Angstrom) radiation and the Raman spectrum as shown in inset of Figure 4(a) was recorded using 514 nm laser excitation at the room temperature. Raman spectrum of carbon spheres grown at 140˚C comprises sharp peaks at 1580 cm⁻¹ (G-Band) representing a graphitic structure followed by a broad shoulder at 1330 cm⁻¹ (D-Band), which corresponds to the presence of both sp² and sp³ hybrid carbons and minor peaks correspond to amorphous carbon structure. The TGA/DTA curves as shown in Figure 5 were taken in N₂ atmosphere at a heating rate of 15˚C/min. The TGA was performed to determine the thermal behavior of carbon nanospheres [16]. The TGA curves indicate a slow rate of weight loss up to 160˚C indicating the evaporation of volatile substance and water contents in the samples. Between 160˚C and 260˚C, the samples may have released nitrogen or oxygen contents. Weight loss down to 520˚C indicates a phase change of carbon materials from amorphous to crystalline structure. The sample achieved thermal stability above 520˚C. Fast weight loss at 265˚C results from a TGA of CNs grown at 170˚C for 15 hours, but only 20% of weight loss has been observed. Such fast weight loss can support the idea of hard solid spheres. Two broad exothermic peaks of the DTA curve at 320˚C
Figure 4. The XRD patterns of CNs synthesized at 170˚C for 15 hours, figure (a), and inset shown in figure (a), is the Raman spectrum of CNs synthesized at 140˚C for 6 hours, and a histogram of size distribution of CNs before (bottom bar) and after (top bar) heat treatment, figure (b).

Figure 5. TGA/DTA Curve taken in N2 atmosphere at heating rate of 15˚C/min. and 500˚C indicate the phase transition of samples from amorphous to crystalline, and a thermally stable structure of nanospheres, respectively.

4. Conclusion

Carbon nanospheres with clean surface morphology were successfully synthesized using a green-chemistry method. This method involves the hydrothermal decomposition of glucose in water at considerably low temperatures from 140˚C to 190˚C. The product obtained was characterized by TEM, SEM, TGA, X-ray diffraction, and Raman spectroscopy. The TEM images indicate that the carbon spheres are hard solids and highly dispersed with size ranges from 50 nm to 260 nm on average. The carbon nanospheres synthesized at 170˚C for 6 hours have shown high quality. Carbon active centers on the surface of spheres may be the
cause of fusion of two spheres as seen on the sample grown at 160°C. The presence of a D-band and a G-band in the Raman spectrum supports the TGA/DTA curves which indicate the thermal stability of carbon spheres and phase transition from amorphous to crystalline structure. Our study shows that the post heat treatment of hydrothermally synthesized carbon nanospheres will enhance the surface morphology, reduce the particle size, and become well dispersed.

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References


