Synthesis of Carbon Nano Fiber from Organic Waste and Activation of its Surface Area

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Abstract

Carbon Nano fibers (CNFs) have recently attracted a lot of attention due to their widespread range of technological applications attributed to their unique physical and chemical properties, such as, small size, high strength, high adsorption linked with their large specific surface area, high temperature tolerance and corrosion resistance. CNFs have been used in energy conversion and storage, reinforcement of composites and self-sensing devices. The complete removal of entrapped metallic impurities and amorphous carbon incorporated with CNFs has been a long-standing issue. We have developed a new approach for preparing graphitic CNFs and its activation of surface area by purification. This approach entails Thermal Decomposition (TD) based synthesis of CNFs from organic solid waste, such as, stems of rice plants. CNFs are synthesized from organic waste precursor (Rice Stems) at 900 °C under inert atmosphere. The active surface area was measured using a Surface Area Analyzer. Morphology of CNFs was studied with using SEM and XRD. The SEM image shows that the synthesized CNFs have diameter ranging within 45-60 nm.

Introduction

Properties of porous carbon and carbon based nano-materials strongly depend upon the methods of synthesis employed. Carbon materials have attracted much attention due to their wide range of physical and chemical properties, which lead to broad range of utilities in advanced engineering materials, such as, in aerospace, transportation, actuators, sensors, fuel-cells, radar-absorbing materials, wind-turbine blades, electromagnetic interference shielding, and expensive sporting goods. They also find applications in catalysis, electronic devices, gas and liquid separation, and memory storage devices [1-3].

Carbon fibers and nano fibers are the carbonic materials with high porosity and high specific surface area [1]. Carbon Nano Fibers (CNFs) have unique tubular structures with their diameter in nanometer range, and effectively have large specific surface area [4]. The basic shape of CNFs is cylindrical, with varying lengths and diameters. CNFs have superior physical and chemical properties, and very high mechanical strength. These unique properties make them suitable for a number of applications, e.g., as sensors and probes, thermal resistance, energy storage, and in many other optical, electronic and medical applications [4].

Vapor-grown CNFs have attracted much interest because they have the potential to provide solutions to many problems in composite applications. Unlike glass fibers, they are electrically conducting and therefore suitable for applications that require the ability to discharge electrostatic potentials, provide sufficient conductivity for electrostatic painting, or even shielding from radio frequency interference or lightning strikes. Moreover, their thermal conductivity is excellent [5,6].

In recent years, nanoparticles have been drawing attention [7] in the composite industry sector as they have the ability of enhancing the mechanical and physical properties of fiber-reinforced composites synthesized through older methods. Their nanometer size, connecting to high specific surface area, combined with extraordinary mechanical, electrical and thermal properties, make CNFs unique nano-fillers for structural and multifunctional composites. While there is so much investigation reporting wide range of utilities of carbon nanotubes and CNFs in composites, the potential of these nano-materials to enhance the damping properties of composites still remains relatively less explored [8-10].

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More Information

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Numerous research works for the synthesis of Carbon Nano Materials (CNMs) have been carried out using precursors obtained from fossil fuel and petroleum products. The availability of such precursor and its cost are factors that determine the cost-effectiveness of finally produced carbon nanomaterials. Hence, there is a need to search for new sources of precursors, which are not fossil-fuel based. Plant derived precursors can yield CNMs of similar, and sometime even better, quality than one would normally get starting with fossil-fuel materials and petroleum [11].

In the present work, we synthesized the CNFs using stems of Rice plants by the method of Thermal Decomposition (TD). Stems of Rice plants are usually dumped and incinerated by the farmers, which leads to a number of environmental issues. Transformation of such biomass to emerging renewable materials is advantageous compared to the traditional dumping and incineration [9].

Among the various resources available for synthesis of nanostructured materials, plant waste is superior in terms of cost-effectiveness and it is eco-friendly also. Hence, we tried to synthesize the CNFs using plant based biomass through the TD route. The active surface area analysis has been done using Surface Area Analyzer instrument before and after the treatment with 50% Conc. HCl and HNO3. Morphology of CNFs was investigated through Scanning Electron Microscopy (SEM), Atomic Force Microscopy (AFM) and X-Ray Diffraction (XRD) analyses.

**Materials and Methods**

CNFs were synthesized from organic waste precursor, i.e., stems of the rice plant collected from Kalyan, India. The stems of the rice plant were washed with distilled water to remove surface contamination (e.g., dust, pesticide, etc.). It was then soaked in 10% KOH solution for 12 h. The soaked samples were washed with distilled water several times to reach the neutral pH and dried in oven at 120 °C. Finally, the dried stems of rice plant were thermally decomposed at 900 °C under inert atmosphere of hydrogen gas for 2 h in a horizontal furnace. The CNFs were treated with 50% Conc. HCl and HNO3 to remove the alkal, amorphous carbon and metals present. Finally, the CNFs were filtered and washed with distilled water several times to reach the neutral pH, and finally dried in an oven at 100 °C.

**Characterization techniques**

**FT-IR Spectroscopy:** The CNFs was mixed with potassium bromide (KBr) (FTIR grade) in the ratio of 1:2 and pressed with the help of hydro-press to make pellets. The sample pellet was positioned into the sample holder and FTIR spectra were recorded on FTIR-4100 (Jasco).

**Scanning Electron Microscopy (SEM) and Energy-Dispersive X-Ray Spectroscopy (SEM-EDX):** SEM was done at IIT Bombay, India on a Jeol JSM -7600F FEG-SEM Model No. JSM-7600F with SEI Resolution 1.0 nm at 15 kV, 1.5 nm at 1 kV, in GB mode having magnification as low as 25X to 10,000X to as high as 100X to 1,000,000X with accelerating voltage ranging from 0.1 to 30 kV and probe current ranging from 1 pA to ≥ 200 nA [3].

**X-Ray Diffraction (XRD) analysis:** XRD measurements were accomplished at IR-Tech Lab, Mumbai, India on MiniFlex machine unit with Cu-Kα radiation (λ = 1.54178 Å) at 40 kV and 15 mA, at a scan-speed of 04.00 °/min.

**Raman spectroscopy:** Raman spectroscopy is based on the Raman Effect, which results from interactions of the vibrational modes of molecules with electromagnetic radiation. Raman spectroscopy is well known as a powerful tool for the characterization of carbon structures. The ratio of intensities of the D and G bands, R, gives interesting information on the structure of carbon materials [3]. It was carried out at University of Mumbai, India.

**Surface area analysis:** Active surface area was analyzed through BET Surface Area Analyzer from Smart Instruments Co. PVT. LTD.

**Results and Discussions**

In this work we developed easy, reliable and eco-friendly methods for synthesis CNFs. The waste Stems of the rice plant were pyrolysed at 900°C in an inert atmosphere to get CNFs.

**FTIR Spectroscopy**

FT-IR analysis (Figure 1) show that synthesized treated CNFs have C–C and C≡C stretching which was indicated by peaks at 1595 cm-1 and 1397 cm-1, 705 cm-1, 917 cm-1, 993 cm-1 also indicate the C–C binding stretching in finger print region and peak at 3431 cm-1 indicate that presence of amine (NH2) group on the smooth surface of CNFs [8].

**Surface morphology:** SEM analysis (Figure 2a,b) shows that the average diameter of the CNFs is 40-60 nm and they have smooth surface. Figure 2a shows matrix residue on CNFs Surface, which was removed by treatment with acid as shown in figure 2bc. The EDAX spectra in figure 2d shows the absence of other impurities like metal salts, such as, halides and oxides.

Figure 2a further shows the presence of CNFs of a few micrometer in length. Figure 2b depicts a closer view of the
same CNFs showing length in the range of a few hundreds of nm or longer. Figure 2c demonstrates a clear picture of one of the CNFs, showing aperture of around 35.8 nm size for a fiber, and the wall measuring around 73.2 nm in another one.

**X-Ray diffraction analysis:** The X-Ray Diffraction (XRD) pattern for as synthesized CNFs (Figure 3a) show broadening of spectra as compared to the one shown in figure 3b for purified CNFs sample. This indicates the presence of impurities in as synthesized CNFs. Figure 3b further shows a sharp peak at 26.46° corresponding to carbon crystallographic structure of the carbon fibers. The sharp peaks at 26.46°, 42.35°, 44.41° and 54.49° for Hexagonal shape (JCPDS No. 00-008-0415) marked by their indices [0 2 2], [1 0 0], [1 0 1], [1 0 4] were also observed. The diameter of the CNFs calculated from Scherrer formula was found to be around 33 nm.

**Raman spectroscopy:** The ratio of the intensities of disorder-induced phonon mode (D-band) and graphite band (G-band) in Raman spectra is a good indicator of the quality of samples. Figure 4 shows the intensities of these Raman bands that indicate a high quantity of structural defects present in the samples [9].

The G-band is visible at 1550–1600 cm⁻¹, which is produced from the high degree of symmetry and order in carbon materials. The D-band at 1250–1450 cm⁻¹ is related to the disorder-induced phonon mode at the boundaries of Brillouin zone. The second order D-band (also called the D”-band), which is related to the boundary point K in Brillouin zone of graphite and depends upon the packing in three-dimensional space, is evident around 2500–2900 cm⁻¹. Moreover, the peaks at 2785 cm⁻¹ and 3111 cm⁻¹ are due to symmetric and asymmetrical C–H stretching vibrations of the CH₃ group, and asymmetrical C–H stretch vibrations of the CH₂ group, respectively. The band at 1600 cm –¹ is assigned to the conjugate C=C bond, and the band around 1388 cm–¹ is due to the CH₃ twisting.

**Atomic Force Microscopy (AFM) analysis:** The size and shape of the CNFs were obtained directly from tip-corrected AFM measurements, and the shape of the CNFs is estimated on the basis of AFM images and line scans. Figure 5 show the tube like structure of CNFs with average diameter (inner and outer wall) of 149.4 nm.
Synthesis of Carbon Nano Fiber from Organic Waste and Activation of its Surface Area

Surface area analysis: Surface area of as synthesized CNFs was found to be 106 m²/g which increase on treatment with 50% Conc. HCl and HNO₃ to be 250 m²/g.

Effect of acid treatment

Acid treatment is one of important ways to improve the graphitization of CNFs and remove the amorphous carbon. The effects of acid treatment on synthesized CNFs were shown by the increased surface area and the quality of the synthesized CNFs. The surface area of synthesized CNFs from rice stem was about 106 m²/g and acid treated CNFs have 250 m²/g. Clearly the specific surface area was found to increase after acid treatment, which may be attributed to more efficient graphitization of the fibers.

Conclusion

Qualitative CNFs can be synthesized from organic waste precursors, i.e., stems of rice plant. The average diameters of the CNFs synthesized in this investigation were found to be in the range of 40-60 nm. The treatment of CNFs with 50% Conc. HCl and HNO₃ shows that matrix residue on the surface of CNFs was completely removed, and a smooth fiber surface was obtained. Acid treatments not only removes the amorphous carbon but also it improved the active surface area. Such CNFs may find applications in the areas of sports, automotive, construction industries, aerospace, marine, and electrical energy sectors etc.

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