Graphene Structure in Carbon Nanocones and Nanodiscs

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Carbon nanoparticles, like nanocones and nanodiscs, can be obtained by mechanical treatment of carbon nanofilaments. Microstructural studies suggest that in nanocones the conical graphene stacking with progressively increasing apex (cone) angles does not fully agree with current theoretical geometry models, such as a closed cones model and a cone−helix model. The unusual stacking form of nanocones was taken into account in a modified cone−helix model. The formation mechanism of the distinctive microstructure is attributed to the inclined anchoring effect, and the relaxation of internal stresses, which were induced by the confined pyrolysis process, resulting in easier disintegration by sonication the nanocones. This is disclosed for the first time in literature regarding the attainment of uniform carbon nanoparticles.

Introduction

Carbon nanoparticles consisting of conical graphene stacking is important in both science and engineering aspects. Potential applications such as enhanced field emission sources and catalyst supports are under rigorous development,1−3 due to their specific electronic states predicted by theoretical calculations,4,5 as well as the good balance between chemical reactivity and structural strength attributed to the graphene arrangement. Their high-efficiency electrochemical capacities was also demonstrated in comparison to other carbon materials with different microstructural alignments.6 In general, conical carbon materials were synthesized by pyrolysis of hydrocarbons with metal clusters. In this case, their conical topography is dominated by the surface morphology of catalysts.7 In addition, natural graphene cones could be found in graphite minerals.8 On the basis of the stacking form of microstructure, there are three major types of conical carbon nanomaterials: a cup-stacked (stacked closed cones) type,9,10 a lampshades (stacked open cones) type,11,12 and an acute angle (graphite cone) type.13−15 Two theoretical microstructure models, closed cone model9,16 and cone−helix model,8,17,18 have been proposed, respectively, to describe geometric organization of the conical graphenes.19 The closed cone model demonstrates the formation of conical graphene by introducing pentagonal rings into the honeycomb graphene layer. The cone−helix model illustrates a helical stacking with continuously wrapping graphene by applying a parameter of the overlap angle.

In this study, uniform carbon nanoparticles with specific shapes were obtained using sonication of the carbon nanofilaments synthesized by a template method.20 Characteristic morphologies of the nanoparticles which had never been reported in literature are disclosed herewith. Both the structure and the formation mechanism of the resultant nanodiscs and nanocones are further demonstrated and discussed. The microstructure studies showed that the nanoparticles formed an unusual graphene arrangement, which does not agree well with the existing models. This unusual stacking form would be taken into account in our modified cone−helix model.

Experimental Section

The precursor was petroleum pitch (A-240, Ashland Inc.) composed of discotic hetero-polycyclic aromatic hydrocarbons (PAHs) units, and the porous anodic aluminum oxide with nanosized channels was used as templates. The preparation of carbon nanofilaments is shown in Supporting Information Scheme S1, and the details can be found elsewhere.20,21 The nanofilaments were annealed at 2400 °C for 5 h in a graphitization furnace to attain a graphitic structure with better crystallinity. The destacking of nanocones was performed in a sonication bath at 1000 Hz for 3 h. The final products were characterized by a scanning electron microscope.
microscope (SEM, JEOL JSM-6500F at 15 kV) and a high-resolution transmission electron microscope (HRTEM, JEOL JEM-2010).

Results and Discussion

After sonication in an ethanol solvent, the as-prepared carbon nanofilaments, with two different diameters, were disintegrated entirely to form countless nanoparticles, as shown in Figure 1a,b. Disc-like nanoparticles are manifested in Figure 1a, while a mixture of conical and disc-like nanoparticles can be observed in Figure 1b. The diameters of these nanoparticles are 120 ± 20 nm (Figure 1a) and 300 ± 50 nm (Figure 1b), respectively, corresponding to those of original nanofilaments. The thickness is in the range 10–100 nm. As the large-pore-sized template was used, both conical and disc-like nanoparticles are found in the products, whereas only disc-like nanoparticles can be observed from the small-pore-sized template. Therefore, one or two kinds of nanoparticles: carbon nanodiscs and nanocones, named after their contours, can be obtained via destacking of herringbone-type nanoparticles with different diameters by mechanical treatment.

The side view of a nanodisc in Figure 2a shows the thickness of ~100 nm. Thinner nanodiscs were difficult to see, since they tend to lie flat on the supporting film of TEM grids and become quite "transparent" to the electron beam. The lattice image (Figure 2b) enlarged from Figure 2a shows that the orientation of stacked graphenes of the nanodisc is parallel to the circular plane. The inset diffraction pattern in Figure 2c reveals that the nanodisc is amorphous. The blurred rings can be approximately assigned to (100) and (110) planes, indicating that the microstructure of the nanodisc is stacked in a turbostratic arrangement. The cross-angle between the sets of spots, known to correlate directly with the average apex angle of the nanocone, is about 34°. Yoon et al. found that the catalytically grown carbon nanofibers consist of assemblages of small substructures, which are called carbon nanorods and nanoplates. The substructures are considered the basic building units for diverse fibrous nanocarbons. Despite our nanoparticles being made from templatesynthesized nanofibers, they can be considered the fundamental building blocks for the original nanofibers as well. However, the circular periphery of our nanoparticles is distinct from the above-mentioned substructures, which usually show a polygonal contour.

The graphene arrangement of the nanoparticles is studied in detail by considering the current conical graphene models. According to the closed cones model, the planar topography of the nanodisc should reveal stacking with graphenes composed of hexagonal carbon rings only. As the pentagon rings were introduced in part into hexagonal networks, the pentagonal "defects" will curve the planar structure to form the conical shape. In terms of Euler’s theorem and the symmetry of graphenes, there are five possible apex angles that are geometrically permitted by inserting pentagons from one ring to five rings into the hexagonal framework, resulting in apex angles \( \theta = 19.2^\circ, 38.9^\circ, 60.0^\circ, 83.6^\circ, \) and 112.8° respectively. The values of apex angle, \( \theta \), are given by

\[
\theta = 2 \arcsin(1 - n/6) \tag{1}
\]

where \( n \) is the number of incorporated pentagons. The simulated molecular structure of the graphenes with pentagon defects is built by Hyperchem and illustrated in Figure 3. Nanocones with distinguishable apex angles corresponding to the molecular models can be found in our samples, as shown in Figures 2e and 3a,b,d. The apex angles were measured to be 38°, 57°, 78°, and 112°, respectively. The measurement error of 0.7–7% is due to the random inclination of the nanocones with respect to the observation direction.

Although the closed cones model defines five kinds of seamless folding of graphene cones, various apex angles that do not coincide with this model are still observed. The cone–helix model


Figure 1. SEM images of the resultant products: (a) disc-like nanoparticles and (b) mixture of the disc-like and conical (arrowed) nanoparticles found after sonication treatment.
is then considered because it is able to account for conical graphenes of most apex angles, including the five from the closed cones model. The cone–helix model is constructed by introducing a parameter, the overlap angle, to estimate the overlapping degree of wrapped graphene sheets, resulting in the various conical graphenes. Another parameter, the degree of graphitic alignment, was adopted to justify if the overlapped graphenes are energetically stable enough. It evaluates how many carbon atoms among the layers can achieve the alignment of lattice points in graphite crystal. When the two parameters are combined, a series of conical graphenes with all possible apex angles can be generated (see Table S1 in Supporting Information). Our products always have a pair of tapered and funnel-shaped ends with different apex (cone) angles. The angles of any ends agree with one of proposed angles from the cone–helix model. For instance, the angles of the nanocone in Figure 2e are 30° (funnel-shaped end) and 38° (tapered end), which can be assigned to the overlap angles of 240.0° and 267.8°, respectively.

Despite the fact that microstructure at each end of the nanocone was expounded well by a cone–helix model, it still cannot explain why the two ends have different apex (cone) angles. In fact, the apex (cone) angles of the conical graphene stacking in nanocones

![Figure 2. TEM microstructure of the carbon nanoparticles: (a) the cross-sectional view of a nanodisc, and (b) lattice image enlarged from (a) revealing the orientation of stacked graphenes; (c) top view of the nanodisc; (d) lattice image of a nanocone, and (e) the low-magnification image of the nanocones; two sets of (002) diffraction spots are observed in the inset diffraction pattern.](image-url)
increase continuously from funnel-shaped end to tapered end. This is different from any reported conical carbon materials with regular apex angles, which can be perfectly interpreted by the cone–helix model. Furthermore, it is noticed that the average interlayer spacing has a ~4.6% difference between the peripheral and central parts of the nanocones (see Figure S1 in Supporting Information). This difference is small and not easily observed. In contrast, the interlayer spacing among stacked graphenes developed from the cone–helix model or closed cones model should be identical everywhere. To summarize the discussion with the stacking form of the nanocones, a possible extension of the cone–helix model is proposed. The nanocone is constructed by the stacking of conical graphenes with progressively increasing apex (cone) angles (decreasing overlap angles) from funnel-shaped end to tapered end. The comparative schematics of current and modified cone–helix structure are shown in Scheme 1.

The formation mechanism of the nanoparticles is attributed to both confined pyrolysis process and inclined anchoring effect. In the softening stage, the channels of the template were impregnated with liquid-phase precursor. The arrangement of discotic hetero-PAHs units in softened precursor was dominated by the competition between the self-assembly arisen from the van der Waals force and the bonding to the template surface by the anchoring effect. 24 The previous studies have demonstrated that in our experiment the PAH units in the liquid-phase precursor would show the edge-on state but not anchor on the template surface with an orthogonal angle. 20 The “inclined” edge-on anchoring in our products is distinctive from that mentioned in other literature. 24 This competition caused the hetero-PAHs units, which are not stacked, to follow the lowest energetic way as the cone–helix model defines, and the pyrolysis process that followed forced the graphene framework to be established simultaneously in the confined channels, generating internal stresses that result in structural instability. Usually, the conical carbon nanomaterials were synthesized by a catalytic pyrolysis method in which the graphenes were stacked layer by layer on the metal catalysts, so the internal stresses were released during the growth process. The relaxation of internal stresses plays a crucial role in disintegrating the nanofilaments forming the nanoparticles under an applied mechanical force. It is suggested that whether or not sliding occurs at interface of the conical graphenes is dependent on whether the cone angles are energetically stable or not, in a relative sense.

Conclusions

Carbon nanoparticles, like nanodiscs and nanocones, with specifically controllable morphologies can be prepared by sonication treatment of herringbone-type nanofilaments synthesized by the template method. The formation mechanism is attributed to the relaxation of internal stresses generated by both the confined pyrolysis process and the inclined anchoring effect, resulting in structural instability. The arrangement of stacked

Figure 3. Microstructural models of the nanocones using the closed cones model: introducing (a) 3 pentagons, (b) 2 pentagons, and (c) 1 pentagon into the hexagonal graphene sheets, generating the conical graphenes with apex angles of $\theta = 57^\circ$, 78°, and 112°.
graphenes of the nanodiscs and nanocones was studied and compared with the current theoretical models. The unusual stacking form in nanocones does not fully agree with the familiar cone–helix models; thus, it should be taken account in a modified model with progressively increasing apex (cone) angles.

To the best of our knowledge, no similar synthesis of carbon nanocones and nanodiscs has been reported yet. Thus, the sonication of nanofibers can be taken into account as a new approach to produce uniform and controllable carbon nanoparticles. With morphological and microstructural controllability, the nanoparticles may have the potential to develop novel anode materials and other potential applications.

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**Supporting Information Available:** Experimental processes. Table of the overlap apex angles. The studies on interlayer spacing. This material is available free of charge via the Internet at http://pubs.acs.org.

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