Hierarchical porous carbon nanofibers via electrospinning

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Abstract
Carbon nanofibers (CNFs) with diameters in the submicron and nanometer range exhibit high specific surface area, hierarchically porous structure, flexibility, and super strength which allow them to be used in the electrode materials of energy storage devices, and as hybrid-type filler in carbon fiber reinforced plastics and bone tissue scaffold. Unlike catalytic synthesis and other methods, electrospinning of various polymeric precursors followed by stabilization and carbonization has become a straightforward and convenient way to fabricate continuous CNFs. This paper is a comprehensive and brief review on the latest advances made in the development of electrospun CNFs with major focus on the promising applications accomplished by appropriately regulating the microstructural, mechanical, and electrical properties of as-spun CNFs. Additionally, the article describes the various strategies to make a variety of carbon CNFs for energy conversion and storage, catalysis, sensor, adsorption/separation, and biomedical applications. It is envisioned that electrospun CNFs will be the key materials of green science and technology through close collaborations with carbon fibers and carbon nanotubes.

Key words: hierarchically porous carbon nanofibers, activated carbon nanofibers, electrospinning, polymer precursors

1. Introduction

Carbon fibers are high aspect ratio graphitic fibers, and possess great technical and industrial importance due to their wide-ranging properties, like very high strength to weight ratio, superior electrical and thermal conductivity, and excellent chemical resistance [1]. Besides other applications, carbon fibers have been predominantly used in high-performance fiber-reinforced composites, which are extremely desirable in automotive, aerospace, and sport industries. These fibers have gained a lot of attention since Iijima reported his results on the closely-related carbon nanotubes (CNTs), thus creating a revolution in the arena of nanotechnology in 1991 [2]. Based on valuable developments in the nano-era, wide potential applications for carbon fibers were projected and huge efforts were focused on their synthesis, in order to improve their properties, which caused an exponential increase in the number of related research articles published in the last two decades.

Generally two approaches have been adopted to fabricate carbon nanofibers (CNFs): vapor growth and electrospinning methods (Figs. 1a and a’) [3]. In the thermal growth method, the carbon fibers are fabricated through catalytic decomposition of certain hydrocarbons in the presence of catalysts (metal particles), but this method entails great difficulties for the mass-production of CNFs. Besides this, other conventional preparation methods for CNFs are also recognized to be very complicated and overpriced, including the spraying method, substrate method, and plasma-enhanced chemical vapor deposition method [4]. Therefore, a simple and low-cost electrospinning method, first patented by Cooley [5] in 1902, has gained a lot of attention during the last decade as the optimum method to produce continuous CNFs.
Due to the increase in research activities related to CNFs, herein we aim to present a comprehensive review on the development of electrospun CNFs fabricated using various polymeric precursors as well as their practical applications, and so reveal a complete picture of the research work in this field in the last few years.

3. Precursors of CNFs

Polymeric materials with a carbon back-bone can be used as the precursor of carbon fibers, but usually CNFs are produced from the polymeric precursors: polyacrylonitrile (PAN), cellulose, and pitch. Other precursors, such as poly(amic acid) (PAA), polyimide (PI), polybenzimidazole (PBI), poly(p-xylenetetrahydrothiophenium chloride) (PXTC), and poly(vinyl alcohol) (PVA), have also been reported. The microstructural characteristics of electrospun CNFs are extremely dependent on the physical and chemical natures of the precursor nanofibers. Therefore, controlling electrospinning conditions, like solution properties (temperature, concentration, solvent type, surface tension, and conductivity), processing parameters (applied voltage, spinneret diameter, solution flow rate, and tip to collector distance) as well as ambient conditions (relative humidity and temperature), can result in CNFs with relatively better or deteriorated performance [12].

3.1. PAN based CNFs

The most common polymer for the preparation of CNFs is PAN, mainly due to its high melting point and carbon yield along with the ease of attaining thermally stabilized products. In
addition, the surface of PAN-based CNFs can easily be modified using a coating or activation process [13]. Furthermore, PAN can be mixed with other polymers or be embedded with nanoscale components (nanoparticles, nanotubes, nanowires, or catalysts) to obtain multiphase precursors and subsequently to make composite CNFs. Different types of PAN-based CNFs are summarized in the following.

3.1.1. Pure PAN based CNFs
During the last decade, detailed researches have investigated fabricating PAN based CNFs by electrospinning [14]. Similar to the production of carbon micro fibers, CNFs have been successfully prepared by electrospinning the PAN/dimethylformamide (DMF) polymer solution followed by a two-step process: stabilization and carbonization [15]. Various stabilization and carbonization conditions have been described, in which stabilization was carried out in air at temperatures between 200-300°C while carbonization was further performed in an inert atmosphere up to 2800°C [16]. Progressive and multi-stage heating methods were used to reduce mass loss and dimension shrinkage. The gradual stabilization and carbonization procedure of 5°C/min from 30-230°C, 1°C/min from 230-270°C, and then 5°C/min from 270-800°C led to very little change in fiber packing, much less planar dimensional shrinkage, with a significant increase of carbon yield as compared to the previously reported procedure [17].

Wang et al. [18,19] and Kim et al. [21,22] deeply investigated the effect of pyrolysis temperature using Raman spectroscopy, and reported that the positions and widths of the D and G peaks for electrospun PAN-based CNFs are insensitive to the pyrolysis temperature, while refractive index (RI) and crystallite width (La) was shown to have a steady decrease, and increase, with temperature, respectively. The value of La increased from 1.5 (600°C) to 2.64 nm (1200°C) and from 1.6 (700°C) to 3.17 nm (1000°C). The large negative magnetoresistance along with low La values present in the CNFs (1000°C), when positioned in a magnetic field at low-temperature, between 1.9-300 K, could establish the formation of a turbostratic structure [19]. A very high degree of carbonization is evidenced when CNFs are treated at a relatively high temperature (2800°C), as indicated by a very sharp G peak and weak D peak in the Raman spectra. Zussman et al. [20] also investigated the microstructural properties of electrospun PAN-based CNFs that had undergone heat treatment. CNFs heated from 700 to 1000°C had a low degree of carbonization with an average crystallite size (Lc) of 1.284 nm [20]. However, it was found that the RI of CNFs was 0.925 $\pm$ 0.04 and was less than 0.5 after heat treatment at 2000°C, revealing a higher degree of carbonization and a more ordered structure [21].

The fabrication of CNF webs, consisting of CNFs obtained through the activation of electrospun PAN-based CNFs having highly hierarchically porous structures, can lead to a significant expansion of applications of CNFs, such as in electrode materials, high-temperature filtration and the removal of toxic gases. Steam is widely used as an activating agent for fabricating ACNFs because of its low cost and environmentally friendly characteristics (Fig. 1b). Kim and Yang [22] reported that the specific surface area of steam-ACNFs decreased with increasing activation temperature (from 700-850°C) due to the unification of micropores at elevated temperatures.

3.1.2. Composite PAN based CNFs
In order to create some functional properties in PAN-based CNFs, a variety of nanomaterials have been integrated into precursor solutions to fabricate composite CNFs by co-electrospinning and subsequent thermal treatment. High orientation of the PAN chains were induced by creating significant interactions using a thermal treatment between PAN and CNTs. The relative orientation coefficient adjacent to multi-walled CNTs (MWCNTs) in the core region of CNFs was 0.79, higher than that without CNTs, 0.44, and the relative orientation coefficient and crystallite size increased with an increase of MWCNTs concentration. At the same time, the diameter of the as-spun nanofibers could be decreased with an increase of CNTs content due to the resulting higher electrical conductivity [23]. Later, continuous single-walled CNTs (SWCNTs) were incorporated into PAN-based CNF yarns using a drum as the collector. A non-graphitized structure including graphite layers and layered carbon was formed after graphitization at 1100°C. The modulus of the stabilized SWCNT/PAN composite nanofibers was increased from 60 to 150 GPa by increasing the SWCNT content from 0-4 wt%. Although SWCNTs might protrude out of the composite CNFs due to the shrinkage of the fibers during thermal treatment, forming a rough surface, well-distributed and oriented CNTs were achieved in almost every section investigated, which is favorable for enhancing mechanical, electrical and thermal properties [24]. Positron annihilation spectroscopy is recognized as a versatile method for checking the evolution of vacancy-type flaws and their clusters in solid materials. Ra et al. [13] first employed the method to study electrospun CNFs embedded with MWCNTs composite and found that discrete positron trapping sites in the form of vacancy-type defects formed at the interfaces among the MWCNTs and PAN matrix.

Besides MWCNTs, other inorganic species (e.g. Ag, TiO$_2$, and Mn) have also been added into the PAN precursor solution to prepare functional composite CNFs. PAN/Ag nanofibers were obtained, and the diameter of the CNFs varied with the Ag content from 0.5 wt%; the smallest CNFs were obtained at 1 wt%, which is in disagreement with the rule that the diameter usually increases with increasing conductivity of the solution [13]. Kim and Lim [25] investigated the physicochemical and photocatalytic activities of oxidized TiO$_2$-embedded PAN-based CNFs (Ox-TiO$_2$/CNF). Brunauer-Emmett-Teller (BET) specific surface area of the Ox-TiO$_2$/CNF exhibited a large increase, from 42-223 m$^2$ g$^{-1}$ after post-oxidization, due to the loss of carbon in gaseous form [25]. Oh et al. [26] studied the pore characteristics of Mn-embedded PAN-based ACNFs. By controlling the dispersion of Mn in the resulting composite ACNFs, large specific surface area and micropore volume could be expected due to the physical migration of Mn and its catalytic activation [26]. Recently, hierarchical porous CNFs were incorporated with Fe$_3$O$_4$ nanocrystals using electrospun PAN/polybenzoxazine (PBZ) nanofibers as the composite carbon precursor. By the combination of precursor design and activation process, an extremely high surface area of 1623 m$^2$ g$^{-1}$ and a pore volume of 1.635 cm$^3$ g$^{-1}$ were obtained. Quantitative pore size distribution and fractal analysis were used to investigate the hierarchical porous structure using $N_2$ adsorption methods and synchrotron radiation small-angle X-ray scattering measurements [27,28].

The surface modification techniques applied to electrospun
CNFs can combine the advantages of the coating materials and the large specific surface area of the nanofibrous substrate to provide composite CNFs with more features. Polypyrrole (PPy) was coated on as-obtained PAN-based ACNFs, which were embedded with 5.7 wt% MWCNTs [29]. A three-dimensional (3D) network of PPy particles was well organized on the surface of the ACNFs/MWCNTs. Specific surface area, mesopore volume, total pore volume, average pore size, and electrical conductivity were all increased after the incorporation of the MWCNTs in the ACNFs. Unlike previous studies which used carbonization prior to surface coating, Shao et al. [30] fabricated zinc oxide (ZnO)-coated PAN-based CNFs using sol-gel coating followed by thermal treatment. Separated ZnO clusters with a diameter of 100 nm and perfect crystalline structure were formed after the carbonization. This special surface might be favorable for some interesting applications. However, due to the low carbonization temperature (450°C), the CNFs still exhibited an amorphous structure [30]. Recently, Shin et al. [31] synthesized two distinct 1-D CNFs encapsulating irregularly isolated SnCo nanoparticles via electrospinning of polyvinylpyrrolidone (PVP), and PAN polymers containing Sn-Co acetate precursors with subsequent calcination in reducing atmosphere. CNFs fabricated with PVP undergo structural degradation of the polymer during the carbonization processes, and have exhibited irregular segregation of heterogeneous alloy particles with a size distribution of 30-100 nm, as shown in Figs. 1c and d [31].

3.2. Pitch based CNFs

Pitch, another precursor used to fabricate CNFs, is generally obtained from petroleum asphalt, coal tar, and poly(vinyl chloride), and has a higher carbon yield and a lower cost compared to PAN [32,33]. However, impurities in the pitch must be fully removed to obtain high-performance carbon fibers, leading to a great increase in cost. Park et al. [34] reported that a DMF/tetrahydrofuran (THF) solvent mixture containing 40 wt% pitch could be electrospun and the diameter of the as-spun pitch fibers was in the micrometer range; it was difficult to make them thinner due to the low boiling point (65-67°C) of the solvent (THF). For the same pitch precursor, the performance of electrospun activated carbon microfibers was compared with that of melt-spun activated carbon microfibers and the results showed that the electrospun carbon microfibers had a higher activation reaction rate compared to melt-spun fibers, because of the finer diameter and less ordered microstructure [35]. BET specific surface area of the electrospun activated carbon microfibers decreased with activation temperature from 700-900°C, which is opposite to the increased trend obtained by Oh et al. [26] for PAN-based ACNFs treated from 800-1000°C. This was possibly due to the precursor type and the large diameter difference [26].

3.3. PVA based CNFs

PVA, another thermoplastic polymer, is a water-soluble polyhydroxy polymer and has been used as a carbon precursor in fundamental researches, even though it easily decomposes at a high temperature and gives a low carbon yield. Zhu et al. [36] fabricated FeOx-filled PVA-based composite CNFs through carbonization at a relatively low carbonization temperature of 600°C considering transition metals as a catalyst. The resultant carbonized CNFs have shown good conductivity, magnetic, and hydrophobic properties. Theoretically, the increased electrical conductivity of the spinning solution with increasing FeOx content could be predicted to reduce the diameter of the electrospun hybrid nanofibers. However, the diameter of the electrospun FeOx containing nanofibers was increased with an increase of FeOx content [36].

3.4. PI based CNFs

Unlike these thermoplastic precursors, electrospun thermosetting nanofibers like PI can directly undergo carbonization for the fabrication of CNFs without the need for the expensive stabilization process, and have thus gained increasing attention in recent years [37-39]. Kim et al. [40] reported the formation of yellow electrospun webs of PAA at voltages between 13-15 kV, which were then imidized with 81% yield. The imidized webs were then carbonized at 700-1000°C under a nitrogen atmosphere with yields greater than or equal to 53%. The carbonized webs were activated under steam in the temperature range of 650-850°C resulting in a specific surface area of 940-2100 m²/g [40]. Xuyen et al. [37] described that pressure during the carbonization process could play an important role in the electrical conductivity of PI-based CNF mat [37]. Chung et al. [41] embedded iron(III) acetylacetonate (AAI) into PI-based CNFs and investigated the effect of AAI on the degree of carbonization. It was found that AAI acted as an effective promoting agent for the carbonization as indicated by a sharper diffraction peak, lower RI and larger crystallite size [41].

3.5. PBI based CNFs

Aside from PI as a precursor, a series of investigations of PBI-based CNFs and ACNFs were conducted by Kim and co-workers [42]. The electrical conductivity of PBI-based CNFs showed a steady increase from 5.74 × 10⁻¹ to 35 S cm⁻¹ as the carbonization temperature was increased from 700-3000°C. The major reason might be the rigid-chain structure possessed by PBI. Even though the PBI-based CNFs treated at 3000°C are still non-graphitizable carbons, as proved by X-ray diffraction, Raman spectra indicated a very high degree of graphitization. BET surface area kept on increasing with increasing temperature to 800°C and then decreased if electrospun PBI was further heated to 850°C. This trend is different from PAN-based CNFs, which showed a steady decrease in the same temperature range.

3.6. PXTC based CNFs

Aligned nanofiber yarn several centimeters long was obtained by electrospinning another thermosetting polymer precursor (PXTC) [43]. The formation of the yarn might result from the iononic condution of PXTC. The presence of D and G peaks in the Raman spectra signaled a successful change from electrospun PXTC yarn to CNFs in the temperature range from 600-1000°C; yarn carbonized at 500°C did not show these two peaks in the Raman spectra. The mole fraction of graphite for the carbonized nanofibers was determined to be 0.21-0.24.

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and 241 m$^2$ g$^{-1}$, and therefore showed a capacitance of 245 F g$^{-1}$ with a current density of 1 mA cm$^{-2}$[47]. MWCNTs dispersed in CNFs have given improved electric double-layer capacitor performance in aqueous electrolytes. The addition of 3 wt% MWCNTs in PAN precursor increased the surface area to 1170 m$^2$ g$^{-1}$ and electrical conductivity to 0.98 S cm$^{-1}$, and consequently increased electrical double-layer capacitor capacitance to 180 F g$^{-1}$ in 6 M KOH [29]. A coating of polypyrole (PPy) on these MWCNT-embedded CNFs led to a further rise in capacitance, to 333 F g$^{-1}$. For MWCNT-embedded CNFs, electrical conductivity and capacitance increased to 5.32 S cm$^{-1}$ and 310 F g$^{-1}$, respectively, from 0.86 S cm$^{-1}$ and 170 F g$^{-1}$ for the nanofibers without MWCNTs [48]. Composite CNFs containing V$_2$O$_5$ prepared via electrospinning revealed that the content of V$_2$O$_5$ was the major factor responsible for the morphology and pore structures [49]. An electrode made of these C/ V$_2$O$_5$ composite nanofibers led to the highest specific capacitance of 150 F g$^{-1}$ and energy density of 18.8 W h kg$^{-1}$ over a power density range of 400-20 000 W kg$^{-1}$. In another report, boric acid (H$_3$BO$_3$) and urea were used to introduce boron and nitrogen functional groups in CNFs and to increase total surface area [50]. The electrode with these characteristics demonstrated better supercapacitor performances, with specific capacitance of 180 F g$^{-1}$ and energy density of 17.2-23.5 W h kg$^{-1}$ in the power density range of 400-10 000 W kg$^{-1}$.

Continuous graphene-embedded CNFs were fabricated by electrospinning PAN-DMF solution with oxidized graphene nanosheets followed by carbonization at 800°C. The electrochemical measurements revealed the maximum specific capacitance of 263.7 F g$^{-1}$ in 6 M KOH aqueous electrolyte

4. Applications of Electrospun CNFs

Advanced energy conversion and storage systems, like supercapacitors, rechargeable lithium-ion batteries (LIBs), and fuel cells, are in urgent demand to fulfill newly emerging applications such as portable electronics, electric vehicles, and industrial power management. The performance of these devices strongly depends on the properties of the electrode materials used. CNFs have advantages similar to a 1D carbon nanostructure with the additional benefits of being inexpensive, continuous, and relatively easy to use in applications.

4.1. Electrode materials for electrochemical capacitors

Porous carbon materials are essential for electrodes materials for electrochemical capacitors; in commercially available capacitors, activated carbons are commonly used. Iijima and Ichihashi [44] first used webs of PAN-based CNFs as electrodes of electric double-layer capacitors, using 30 wt% potassium hydroxide (KOH) aqueous solution. The 700°C activated webs gave a very high capacitance of 173 F g$^{-1}$ at a low discharge current density, as low as 10 mA g$^{-1}$, but at the higher current density of 1000 mA g$^{-1}$ the 800°C activated webs gave a high capacitance of 120 F g$^{-1}$. They reported that the former had a surface area of 1230 m$^2$ g$^{-1}$ containing micropores, but the latter had a surface area of 850 m$^2$ g$^{-1}$ consisting of mesopores. Similar results were obtained in later works [45,46]. PAN-based CNFs prepared by mixing PAN with 15 wt% cellulose acetate gave surface areas of 919 m$^2$ g$^{-1}$ and 241 m$^2$ g$^{-1}$, and therefore showed a capacitance of 245 F g$^{-1}$ with a current density of 1 mA cm$^{-2}$[47].

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Fig. 2. (a) Scheme for the processing of activated carbons (ACs), hierarchically porous carbon nanofibers (CNFs), and nitrogen-doped hierarchically porous CNFs. Scanning electron microscope micrographs of porous CNFs formed by carbonizing nanofibers at different Nafion:polyacrylonitrile (PAN) blend compositions: (b) 80:20 electrospun at 25% total solid concentration, (c) 80:20 electrospun at 20% total solid concentration. Cyclic voltammetry of (d) carbonized 60:40 Nafion:PAN and (e) carbonized 80:20 Nafion:PAN. (a) Reprinted with permission from [54]. Copyright © 2013, Elsevier. (b-e) Reprinted with permission from [57]. Copyright © 2013, Elsevier.
specific gravimetric and volumetric capacitances of up to 210 F g\(^{-1}\) and 60 F cm\(^{-3}\) in 1 M H\(_2\)SO\(_4\) at a high cyclic voltammetry scan rate of 100 mV s\(^{-1}\) due to the large fraction of mesopores, as shown in Figs. 2b-e [57].

4.2. Electrode material for dye-sensitized solar cells

One-dimensional CNFs were prepared and used as a low-cost alternative to platinum counter electrodes in dye-sensitized solar cells (DSSCs) [58,59]. Electrochemical measurements revealed that the counter electrode made with CNFs exhibited low charge-transfer resistance, large surface area, and fast reaction rates in DSSCs applications. A counter electrode made of Pt/CNFs was developed by solution-depositing Pt nanoparticles onto CNFs [60]. This type of electrode reduced the overall series resistance, decreased dark saturation current density, and increased shunt resistance of the DSSCs. Correspondingly the Pt/CNFs-based DSSCs achieved an energy conversion efficiency of ~8%, which was improved over those of pure Pt- or pure CNFs-based DSSCs. In another research, MWCNT embedded mesoporous CNFs were prepared followed by consecutive stabilization at 280°C, and carbonization at 800°C. A cell using the prepared MWCNT-embedded mesoporous CNFs as a counter electrode material exhibited even higher overall energy conversion efficiency than that of the cell using a Pt counter electrode material [61]. Recently, Yun et al. [54] fabricated hierarchically porous CNFs with the large surface area of 2862.1 m\(^2\) g\(^{-1}\) and 20.6 at% electroactive oxygen atoms. These hierarchically porous CNFs exhibited excellent electrochemical performance, and their thermal treatment with melamine results in nitrogen-doped hierarchically porous CNFs with 9.1 at% nitrogen and improved electrical properties. According to the results, the electrochemical performance of nitrogen-doped hierarchically porous CNFs is better than that of hierarchically porous CNFs, displaying a specific energy of 113 Wh kg\(^{-1}\), specific power of 105 kW kg\(^{-1}\), and stable cycle life of over 5000 cycles (Fig. 2a) [54]. Different morphological structures of CNFs are also presented in Figs. 1e and f [55,56]. Later, Tran and Kalra [57] reported the fabrication of uniformly porous CNFs for supercapacitors by using blends of PAN and sacrificial nafion. The resultant porous CNFs exhibited a specific surface area of up to 1600 m\(^2\) g\(^{-1}\) with a large fraction of mesopores (2-4 nm). Electrochemical measurements showed large

Fig. 3. (a) The fabricating process and i\(^{\text{1}}\) reduction of activated carbon nanofibers (ACNFs) with hollow core/mesoporous shell structure. (b) I-V curves of hollow ACNF (HACNF), Meso-HACNF1, Meso-HACNF2, and Pt counter electrode. (c) High-resolution transmission electron microscope images of donut-shaped lithium face centered cubic (FCC) metal crystallites and (d) stress-strain curves of hollow CNF (HCNF). (e) Cycling performances of freestanding HCNF anodes. (a and b) Reprinted with permission from [59]. Copyright © 2013, Elsevier. (c-e) Reprinted with permission from [70]. Copyright © 2013, American Chemical Society.
Hierarchical porous carbon nanofibers fabricated a 3D free-standing net and -1 versus reaction times for the reduction of 4-nanoparticle, CNFs (squares), Au nanoparticles (circles) and CNFs@Au (stars); (e) catalytic activity of the CNFs@Au network for the reduction of 4-nanoparticle after three rounds of cycling. (a-e) Reprinted with permission from [73]. Copyright © 2013, Royal Society of Chemistry.

### 4.3. Anode materials for lithium-ion rechargeable batteries

LIBs are attractive power sources for a wide range of electric devices from cell phones and laptop computers to hybrid electric vehicles. Kim et al. [21] prepared CNFs by combining electrospinning and thermal treatments. Due to the particular nanotexture, these PAN derived CNFs exhibited the highest reversible capacity (ca. 450 mA h g⁻¹), and a good rate capability (350 mA h g⁻¹ at a charge current of 100 mA h g⁻¹) [21]. CNFs prepared from (PAN/PPy) also showed a relatively large reversible capacity (ca. 360 mA h g⁻¹) with high rate capability, good cycle performance, and structural integrity after 50 charge/discharge cycles, attributable to their large surface area and extended interface with electrolyte, leading to a short pathway for charge and electron transport [62]. CNFs have also been doped with silicon and tin nanoparticles. [63] The fine dispersion of nano-sized particles within CNFs allow them to act as structural buffers, particle stabilizers, and electroactive materials, and also eliminate the need for binding or conducting additives in the electrode materials. Wang et al. [64] firstly used electrospun C/Si composite nanofibers as an anode material for lithium-ion batteries and the composite nanofibers exhibited reversible capacity (1240 mA h g⁻¹) with stable capacity retention even after 40 cycles [64].

Yu et al. [65] prepared tin nanoparticle-dispersed carbon (Sn/C) nanofibers by stabilization of electrospun SnCl₂/PAN fibers and subsequent carbonization at different temperatures. Sn/C nanofibers at 700 and 850°C presented the highest charge (785.8 and 811 mA h g⁻¹) and discharge (1211.7 and 993 mA h g⁻¹) capacities due to the unique feature of reticular nanofibers geometries. Electrospun CNFs anodes loaded with transition-metal oxide nanoparticles have been widely investigated as anode materials in lithium ion batteries, because of their high theoretical capacities, safety, non-toxicity and low cost [66]. One example is the fabrication of C/SnO₂ composite nanofibers with adjustable sheath thicknesses, involving the electrodeposition of SnO₂ on CNFs [67]. These composite nanofibrous anodes have delivered nearly 800 mA h g⁻¹ discharge capacity at the first cycle at 50 mA g⁻¹ current rate, with 69% capacity retention even after 100 charge/discharge cycles, reflecting a major improvement over pure SnO₂ powder-based anodes [68]. Recently Kong et al. [69] prepared C/SnO₂/C core/shell/shell hybrid nanofibrous mats via single spinneret electrospinning followed by carbonization and hydrothermal treatment. The embedded and de-aggregated SnO₂ nanoparticles in the carbon phase, which were less than 10 nm in size, provided a huge number of reaction sites for lithium ions and ensured complete alloying with them [69]. Recently, Lee et al. [70] investigated the mechanical sustainability of a carbon based anode subjected to repeated electrochemical reactions with Li ions via nanotensile tests of individual hollow CNFs, as shown in Figs. 3c–e. Surprisingly, the mechanical properties of such electrodes were improved by repeated electrochemical reactions with Li ions, which is contrary to the conventional wisdom that the mechanical sustainability of carbon-based electrodes should be degraded by repeated electrochemical reactions [70].

### 4.4. Catalysis

Because of their ultra-high specific surface area, excellent electrical properties, very high chemical resistance, and acceptable mechanical properties, electrospun CNFs are materials of choice for catalytic support. For example, CNFs were developed as an electrode support for redox enzymes immobilization which is applied in bioelectrocatalytic O₂ reduction. The valuable effects of these CNFs on the electrical performance of the electrode were attributed to the high loading of active enzymes and fast kinetics at the electrode surface [71]. Without any other catalyst, the porous CNFs showed high activities for nitrogen oxide (NO) removal at room temperature. In this case, carbon acted both as catalyst and adsorbent which enabled the catalytic oxidation of NO into NO₂ or the reduction of NO into N₂ [72]. Recently Zhang et al. [73] fabricated a 3D free-standing network composed of cross-linked carbon@Au core-shell nanofibers and the as-prepared CNFs@Au network exhibited excellent catalytic activity for the reduction of 4-nitrophenol (4-nanoparticle) based on the electron-rich catalytic platform, arising from
the synergistic effect between carbon and Au. Notably, the free-standing 3D nanofibrous cross-linked network structure could improve the catalyst’s separation and reuse (Figs. 4a-c) [73]. Composite Pd and Pt/CNFs with nanocactus- and nanoflower-like morphology were fabricated to act as a highly active catalyst toward the redox reactions of hydrogen peroxide (H₂O₂) and β-nicotinamide adenine dinucleotide [74]. Carbon/Ag nanoparticles composite electrodes have also shown high catalytic activity in the reduction of 4-nitrophenol with NaBH₄ which might be attributed to the high surface areas of Ag nanoparticles loaded on the CNFs and the synergistic effect on delivery of electrons between Ag nanoparticles and CNFs [75]. One dimensional ZnO/CNFs with high photocatalytic activity were successfully prepared to degrade organic pollutants under UV light irradiation [76]. It was demonstrated that CNFs with surface-coated ZnO exhibited higher photocatalytic property than pure ZnO for the degradation of Rhodamine B. In later research, 1D hetero-structures of Bi₂MoO₆/CNFs and In₂O₃/CNFs were fabricated to use in various catalytic activities [77]. Enhanced photocatalytic activity was perceived for those hetero-structures under visible light compared to pure Bi₂MoO₆ or In₂O₃.

4.5. Sensors

Sensors/detectors with suitable operation, simple construction, steady response, high sensitivity, with good selectivity are always required for the determination of various chemicals and biomolecules. CNFs modified carbon paste electrode (CNF-CPE) was developed by casting a water suspension of CNFs onto the surface of a CPE and such electrodes were used to directly detect the three amino acids l-tryptophan, l-tyrosine, and l-cysteine using cyclic voltammetry and constant potential amperometric methods [78]. The modified electrodes have shown excellent electrocatalytic activity and good analytical performance toward the oxidation of amino acids with a detection limit of 0.1 μM. CNF-CPE was also used to build an amperometric sensor device without any enzyme or medium to detect xanthine (Xa) [79]. The dynamic linear range of Xa detection was 0.03 to 21.19 μM with a detection limit as low as 20 nM. The system was effectively applied to estimate the freshness of fish and determine Xa in human urine. Recently, Hood et al. described the development of a bimetal (Fe and Cu)-grown hierarchical web of CNFs electrode for glucose sensing applications, with a sensitivity of 8.29 μA/mM cm⁻² and a linearity up to 8.5 mM (R² = 0.991) (Figs. 5a and b) [80].

For bio sensing applications, carboxylic acid group functionalized CNFs were combined with other nanostructures to fabricate composite electrodes, and then used to incorporate hydroxyapatite (HA) or prussian blue to make a composite. These composites were coated on a polished Au electrode followed by immobilization of cytochrome c. The resulting biosensor established a good electrocatalytic activity and fast response to H₂O₂ sensing, with a detection limit of 0.3 μM [81]. Beside pure CNFs, metal nanoparticle-loaded CNFs could also be used to modify electrodes; for example, Pd nanoparticle-loaded-loaded CNFs was used to coat CPE (Figs. 5d and d’) [55]. The as-modified

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4.6. Adsorption/separation

Carbon materials, particularly activated carbons, have been used as a powerful adsorbent for a long time [87-89]. Lee et al. [90] demonstrated that the ACNFs possess superior capability for formaldehyde gas adsorption even at a low concentration and in a humid environment. ACNFs have also been actively used as an effective tool to remove NO in polluted air [71]. The reported percentage of NO removed by 0.1 g of porous CNFs material was higher than 60% when inlet NO concentration was 20 ppm. CNFs adsorption media have also been used as a promising alternative for bioseparation. Surface-functionalized CNFs with weak acid cation exchange ligand were capable of adsorbing approximately ten-times more protein than their microfiber counterparts [91]. Recently, magnetic Fe₃O₄ loaded CNFs based on PBZ precursors have been synthesized by a combination of electrospinning and in situ polymerization. The benzoxazine electrode exhibited excellent electrochemical catalytic activities toward dopamine (DA), uric acid (UA), and ascorbic acid (AA), which frequently coexist in biological samples, by decreasing their oxidation overpotentials and improving their peak currents. The lowest detection limits of DA, UA, and AA were observed at 0.2, 0.7, 15 μM, respectively. Similarly rhodium nanoparticle-loaded CNFs were dispersed in DMF under ultrasonic agitation and then cast on the surface of a pretreated pyrolytic graphite electrode. The as-modified electrode established excellent electrocatalytic activity toward hydrazine oxidation [82]. In a more recent research, a cobalt nanoparticles-decorated CNFs modified electrode showed a pH-controlled electrocatalytic activity toward the oxidation of cysteine and N-acetyl cysteine, which was used to fabricate an enzymeless sensor for amino acid [83].

In addition to the electrochemical detection of chemicals, CNFs can detect gases through resistance change. Ultrafine CNFs decorated with ZnO/SnO₂ nanoparticles were deposited on an interdigitated electrode array and constructed to detect dimethyl methylphosphonate with a minimum detectable level of 0.1 ppb [84]. Recently, CNFs attached with Pd nanoparticles have shown excellent hydrogen sensing capability at room temperature [85]. Pd nanoparticles could be alternatively deposited on CNFs via supercritical CO₂ method for the sensing of hydrogen (Figs. 5c and c’) [86].

Fig. 6. (a) Illustration showing the synthesis of A-Fe@carbon nanofiber (CNF) by a combination of electrospinning and in situ polymerization. The C/C₀ versus time plots for adsorption of (c) methylene blue (MB) and (d) rhodamine B (RhB) dye solution. The insets show the magnetic responsive performance (60 s) of A-Fe@CNF-3 after adsorption of MB (10 min) and RhB (15 min). (a-c) Reprinted with permission from [92]. Copyright © 2012, Elsevier.
monomers could effortlessly form thermosetting nanofibers by in situ ring-opening polymerization, and subsequently be converted into CNFs by carbonization (Fig. 6). The resultant fibers with an average diameter of 130 nm are comprised of CNFs with embedded Fe$_3$O$_4$ nanocrystals, and could have a high surface area of 1885 m$^2$ g$^{-1}$ and a porosity of 2.3 cm$^3$ g$^{-1}$. The Fe$_3$O$_4$ loaded CNFs exhibited efficient adsorption for organic dyes in water and excellent magnetic separation performance, suggesting their use as a promising adsorbent for water treatment, and also provided new insight into the design and development of a carbon nanomaterial based on a PBZ precursor [92].

4.7. Biomedical application

The application of carbon fibers in the biomedical field has a history of more than 30 years to date [93,94]. CNFs containing β-tricalcium phosphate nanoparticles have shown good biocompatibility for cell growth [95]. The results indicated that bioactive HA strongly interacted with the CNFs through coordination bonds and would provide strong interfacial bonding to host tissues. Fracture strength of the CNFs/HA composite reached 67.3 MPa with 41.3% CNFs. Bioactive glass (BG) is a bioceramic that has been investigated for applications in bone regeneration. CNFs with BG were developed as a substrate for bone regeneration uses. Biomimeralization in simulated body fluid and in vitro co-culture with MC3T3-E1 osteoblasts revealed the improved ability of CNFs/BG composites to promote the in vitro formation of apatite and MC3T3-E1 proliferation (Figs. 7a-c) [96]. Natural bone is a fiber-reinforced hybrid material composed of type-I collagen fibers and HA minerals. CNFs were also used to prepare a CNFs/HA composite to mimic the collagen fiber/HA composite structure of natural bone (Figs. 7d-f) [56]. The fracture strength of the CNF/HA composite with a CNF content of 41.3% reached 67.3 MPa. CNF/HA composites with such strong interfacial bonding and high mechanical strength can be potentially useful in the field of bone tissue engineering.

5. Concluding Remarks and Future Perspectives

CNFs having diameters intermediate between CNTs and carbon fibers with hierarchical pore structures can be conveniently fabricated by electrospinning polymeric precursors followed by stabilization and carbonization. This new class of carbon nanomaterial with 1D nanostructure and associated high specific surface area has rapidly found application in energy conversion and storage, catalysis, sensor, adsorption, and biomedical fields. However, few studies have focused on the stretching of electrospun precursor nanofibers before stabilization, and tension is rarely applied to the nanofibers assembly during stabilization to prevent shrinkage of the fibers and to ensure a large degree of molecular orientations along the fiber axis. Therefore, there is still considerable room for further improvement of the microstructural, electrical and mechanical properties of the final CNFs. Current attempts reveal there is a strong expectation of achieving even higher strength carbon fibers. Future research will be focused on the fundamental correlations between processing conditions and the nanofibers’ structures and it is envisioned that electrospun continuous CNFs are going to contribute greatly to the emerging family of carbon materials.

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