Review

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Recent development of Supercapacitor Electrode Based on Carbon Materials

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Abstract: Supercapacitor has gained significant attention due to its fast charging/discharging speed, high power density and long-term cycling stability in contrast to traditional batteries. In this review, state-of-the-art achievements on supercapacitor electrode based on carbon materials is summarized. In all-carbon composite materials part, various carbon materials including graphene, carbon nanotube, carbon foam and carbon cloth are composites to fabricate larger specific surface area and higher electrical conductivity electrodes. However, obstacles of low power density as well as low cycling life still remain to be addressed. In metal-oxide composites part, carbon nanotube, graphene, carbon fiber fabric and hollow carbon nanofibers combine with MnO$_2$ respectively, which significantly address drawbacks of all-carbon material electrodes. Additionally, TiO$_2$ is incorporated into graphene electrode to overcome the low mechanical flexibility of graphene. In organic active compounds part, conducting polymers are employed to combine with carbon materials to fabricate high specific capacitance, long-term thermal stability and outstanding electroconductivity flexible textile supercapacitors. In each part, innovation, fabrication process and performance of the resulting composites are demonstrated. Finally, future directions that could enhance the performance of supercapacitors are discussed.

Keywords: supercapacitor, electrode, carbon materials, composite

1 Introduction

With the rapid growth of global economy, the loss of fossil fuels and the increasing environmental pollution, there is an urgent need for highly efficient energy storage devices. In this context, electrochemical capacitor, so-called supercapacitor (SC), which is one of the most promising energy storage devices, recently attracts considerable attention owing to its high power density, fast charge–discharge, and long service life [1–6].

Electrode material, which is the corresponding key for SCs, plays an important role in improving the performance of SCs [7–10]. Amongst various candidates, Carbons have been widely employed for fabricating SC electrodes [11–13]. Carbons are such electronically conductive solids that possess satisfactory corrosion resistance, low density, excellent stability and low cost [14, 15]. In addition, the porosity and morphology of carbons can be simply designed by using oxidizing agents at warm processes, which is called activation. These formed highly porous materials had a large surface area up to 2000 m$^2$·g$^{-1}$ [16–18]. The control of porosity in supercapacitors can not only obtain a high specific capacity of ion adsorption, but also measure the diffusion rate of ions in and out of the pores during the charge/discharge processes.

As one member of carbon materials, carbon nanotube (CNT) reveals high electrical conductivity, great electron transport and electrolyte accessibility. As a result, CNT is considered as a promising material to fabricate electrode material. Graphene is another attractive electrode material because of its outstanding conductivity and large specific surface area [19–22]. Nonetheless, low volumetric capacitance, hard fabrication and high costs for commercial production of these carbon materials remain stumbling blocks for making further breakthroughs in excellent performance supercapacitors. With respect to this, much work has been carried out on synthesizing reduced graphene oxide (rGO). One of the most extensive routes to obtain rGO flakes is the Hummers’ method [23]. In this method, first, graphite is oxidized in aqueous medium, then, oxidic graphite was reduced/exfoliated using, as an example, chemical [14–28], thermal [29], hydrothermal [30], electro-
chemical [31, 32] or microwave-assisted reduction means [33].

However, the introduction of rGO cannot solve all issues mentioned above. composites that combining carbon nanomaterials with micro-sized carbon particles/substrates as well as other moieties that contain redox activity can probably address these obstacles [34]. When inserting CNT between graphene nanosheets (GNs), GNs can linked into a stable, thus enhances the surface area of electrolyte ions in contrast to pure rGO. Moreover, CNT can consociate with rGO, which not only offers conductive pathways for electron transport but also cuts down the internal resistance inside the electrode [35–38].

As a matter of fact, it’s undeniable that all-carbon composite materials exist the problems of lower power density as well as lower cycling life. A satisfactory strategy to address these handicaps is to incorporate finely dispersed pseudocapacitive metal oxides such as MnO_2 and TiO_2 into a carbon matrix. MnO_2 is considered as one of the most promising metal oxides owing to its simple synthesis technology, low cost and outstanding electrochemical properties [39]. Combining MnO_2 with highly electrically conductive carbon matrices such as graphene [40], CNTs [41] and hollow carbon nanofibers [42] to increase pseudocapacitive behavior has stimulated extensive research. Here, carbon materials are utilized as scaffolds, and MnO_2 nanostructures are deposited in situ on the surfaces of carbon materials. This innovative design has been validated to reduce the electronic/ion transport way, offer larger surface area with extensive active sites for fast Faradic reactions on electrode/electrolyte interfaces, in a way, realize the enhancement of electrochemical performance [43]. TiO_2 is also widely investigated owing to its abundance, low cost, long-term thermodynamic stability, photostability, nontoxic, excellent pseudocapacitance behavior and easy fabrication [44]. Unfortunately, its low conductivity greatly influences the specific capacitance of TiO_2 electrodes [45]. Incorporating TiO_2 into graphene electrodes can help address the obstacles of low conductivity of TiO_2 and low mechanical flexibility of graphene, respectively. Herein, TiO_2 usually acts as an additive for carbon materials, especially for graphene [46].

Conducting polymer, especially polyaniline (PANI), has captured intensive attention due to its low price, convenient synthesis, high pseudocapacitance and excellent chemical doping/undoping [47–50]. To overcome some of the drawbacks of pure PANI [51], PANI is usually employed to combine with carbon materials to fabricate flexible textile supercapacitors with excellent performance [52], including enhancing specific capacitance, increasing thermal stability, improving electroconductivity and so on. As an example, Lu et al. fabricated a PC/CNTs/PANI nanocomposite, this composite held the specific capacitance of 1090 F·g⁻¹ and revealed a specific energy density of 97 Wh·kg⁻¹ [53].

Many reviews of supercapacitor electrode materials have been reported. However, they usually focus on one single material, such as graphene, CNT, MnO_2 and so on. With the booming development of supercapacitors, one material can not satisfy the demand of supercapacitors, and studies of combinations with different materials need to be urgently intensified. In this work, compositions that use at least one carbon basic and one active basic are demonstrated.

2 All-carbon composite capacitors

Kim et al. [54] fabricated 2D porous graphene/CNT networks with vertically aligned honeycomb structures, the resulting networks could be used to fabricate stretchable supercapacitors electrodes. The schematic illustration of the fabrication process was shown in Figure 1. Here, reentrant structures were synthesized via a directional crystallization process followed by a radial compression. In contrast to conventional frameworks, reentrant structures, which were inwardly protruded frameworks in porous networks, offered fascinating structure-assisted stretchability, like accordion and origami structures. In addition, the resulting structures revealed great conductivities under bi-axial stretching conditions. The 2D auxetic cellular and vertically aligned structures overcome the tradeoff among conductivity, ion-accessible surface area and tensile properties of electrodes and realize the fabrication of high-performance stretchable supercapacitors, which is essential to realize implantable and epidermal electronics.

Diez et al. [55] demonstrated an easy method for the preparation of a partially reduced graphene oxide/carbon nanotube (prGO–CNT) self-standing binder-free film. This film was prepared from a mixture of GO and CNTs by hydrothermal treatment at 210°C to generate a hydrated self-assembled composite, after compressing, this composite finally required high density with tuned areal carbon loading. In contrast to CNT-free electrode, due to the incorporation of a handle of CNTs (only 2 wt%), prGO–CNT film had a greatly enhancement in capacitance retention at high current densities and the stability during cycling, this improvement was beneficial to supercapacitors with high mass loading electrodes. Moreover, this film obtained high volumetric capacitance and excellent capacitance retention (250 and 200 F cm⁻³ at 1 and 10 A·g⁻¹, respectively),
which make it a promising electrode material for supercapacitors.

Li et al. [56] demonstrated a method to fabricate a novel fibrous CNT-aerogel with large specific surface area, high mechanical strength, and admirable electrical conductivity by electrochemical activation and freeze-drying and employed it as fiber supercapacitors (FSCs). In the normal works, the enhancement of energy density was mainly realized by increasing the specific capacitance of the electrode, such as compounding with pseudocapacitive materials. Here, the introduction of ionogel electrolyte made the CNT-aerogel//CNT-aerogel FSC store energy at the interface between the electrode surface and the electrolyte, instead of by electrochemical reactions. This innovation made its power density 1–2 orders of magnitude higher than previous reported FSCs (Figure 2). The higher energy density had been close to the level of lithium-ion batteries and the power density had reached the highest value of
Table 1: Comparison of energy density and power density of different carbon materials. Values were obtained from the respective references

| Electrode material                  | Energy density (Wh Kg⁻¹) | Power density (Wh Kg⁻¹) | Reference |
|-------------------------------------|--------------------------|-------------------------|-----------|
| Carbon foam/CNT                     | 28                       | 3700                    | This work |
| Active Carbon/CNT                   | 0.016                    | 14.4                    | [63]      |
| Graphene/CNT/Nickel foams           | 19.24                    | 5398                    | [64]      |
| CNT/Carbon cloth                    | 5.73                     | 44                      | [65]      |
| Graphene/MnO₂/CNT                   | 3.2                      | 1280                    | [66]      |
| MnO₂/CNT/Graphene                   | 29                       | 1200                    | [67]      |

PSCs. Additionally, due to the non-volatility, high thermal stability of ionogel electrolyte, the assembled FSCs could run normally at a working temperature window from 0°C to 80°C. All of these reveal CNT-aerogel can be a favorable candidate for FSCs and the synthetic FSC is a favorable power source for flexible electronics.

Carbon foam (CF), a sponge-like rigid carbon material which possesses a large surface area with an open cell wall structure. Due to CF’s attractive properties like high chemical stability, excellent electrical conductivity, outstanding corrosion resistance and high mechanical strength, CF has been triggered in order to develop long-life and high-performance electrodes for responding to the increasing demand of energy storage shielding [57–61]. Dang et al. [62] fabricated a long-life and high-performance CF/CNT electrode material by incorporating CNTs with different length (chemical vapor deposition) on a hierarchically 3D CF, this assembly could collect current without adding any binders or conducting additives. Comparison of energy density as well as power density of different carbon materials was demonstrated in Table 1, which revealed CF/CNT materials were the best among pure CNT and CF materials. Superiority of this composite was correlated with the synergistic effect of combining CF and CNT together. This combination suggests a new method for producing electrode based on carbon material for energy storage.

Carbon cloth (CC) is such a substrate that can offer simple ionic intercalation of the electrolyte and a number of fast electro-transport pathways without adding polymer binder [68]. Owing to these favorable advantages, recently, Kadam et al. [69] grew crumpled sheet like RGO on CC through facile hydrothermal method without using any binder, which reduced contact resistance between current collector and active nanomaterials [68]. To investigated its performance, RGO was prepared within a wide working temperature window (120–200°C) with the step width of 20°C. Moreover, RGO exhibited good long-term cycling stability, and the capacitance loss after 1000 cycles was only 3%. Owing to these interesting structural, morphological and electrochemical properties, RGO grown on carbon cloth is considered as a promising electrode material for new generation of electrode based on carbon material for energy storage.

Zhang et al. [70] fabricated a 3D graphene-based nanostructure with graphene aerogel templating graphene nanosheets (GA-GNs) by an improved hydrothermal method and a microwave plasma chemical vapor deposition process. The schematic illustration of this 3D GA-GNs was shown in Figure 3. GA-GNs exhibited a prominent electrical conductivity of 1000 S/m, which were promising candidates for high-performance supercapacitor electrodes. Moreover, this resulting free-standing and binder-free GA-GN presented a specific capacitance as high as 245 F g⁻¹, outstanding rate capability and desirable long-term cycling stability (92% capacitance retention after 10000 cycles). Such an all-carbon electrode was used to fabricate a two-terminal symmetric solid supercapacitor, this supercapacitor shown favorable areal capacitance (1.2 F cm⁻²), low internal resistance and desirable long cycling stability (90% capacitance retention after 5000 cycles).
3 Metal-oxide composites capacitors

3.1 MnO$_2$-based composites capacitors

Patil et al. [71] demonstrated a developed coaxial fiber-shaped asymmetric supercapacitor (CFASC), which addressed the obstacles of traditional fiber-shaped supercapacitor’s low capacitance caused by its restricted surface area between two fiber electrodes and low energy density owing to the operating voltage range. Here, a gel electrolyte-coated core positive electrode was simply wrapped with the negative electrode, and MnO$_2$ with a little weight percentage of carbons, which is a new step of energy storage materials.

Jia et al. [72] reported a novel mesostructured CNT-on-MnO$_2$ nanosheet composite with a high weight percentage of MnO$_2$, which was constructed by vertically aligned MnO$_2$ nanosheets and in-situ formed oriented CNTs on MnO$_2$ nanosheets. The fabrication process was shown in Figure 4. Due to the synergy effect, this assembly exhibited a high specific capacitance (up to 1229 F·g$^{-1}$) and desirable long-term cycling stability (94.9% capacitance retention after 100000 cycles). The fabrication of this CNTs/MnO$_2$ opens the door for preparing MnO$_2$-CNT nanocomposites with a little weight percentage of carbons, which is a new step of energy storage materials.

Zhang et al. [73] reported ternary rGO/MnO$_2$/CF nanocomposites, which had a 3D hierarchical multihole architecture. Owing to the synergistic effect of CF conductive skeleton, pseudocapacitive of MnO$_2$ and hierarchical porous structure jointly, specific capacitance of the composites reached up to 356.5 F·g$^{-1}$ at a scan rate of 10 mV·s$^{-1}$. Furthermore, the hybrid of rGO helped prevent the exfoliation of MnO$_2$ nanoparticles, so the resulting composites possessed a desirable long-term cycling stability (93.6% capacitance retention after 2000 cycles). The research firstly demonstrates the value of CF in potential application for high-performance supercapacitors. CF, derived from MP, is expected to composite with MnO$_2$ to apply on a broad range of emerging electrochemical supercapacitor.

Çakıcı et al. [74] firstly fabricated highly flexible carbon fiber fabric (CFF) filled with coral-like MnO$_2$ structures via a green hydrothermal process at different conditions. Here, green hydrothermal method realized the assembly of CFF and coral-like MnO$_2$ structures to improve the pseudocapacitance properties of the resulting MnO$_2$/CFF composites, and CFF acted as an perfect template because it could both support substrate and reductant under the hydrothermal condition. Subsequently, the composites were used as electrodes for electrochemical supercapacitors, Figure 5 shown the electrochemical properties of this device, this electrode material containing CFF, coated with coral-like MnO$_2$ structures, obtained excellent specific capacitance (463 F·g$^{-1}$ at 1 A·g$^{-1}$ in a 1.0 M Na$_2$SO$_4$ electrolyte) and remarkable capacitance retention of 99.7% even after 5000 cycles. Moreover, this device revealed excellent stability and energy density (20 W·h·kg$^{-1}$).

Zhao et al. [75] synthesized a state-of-the-art hierarchical hollow nanostructure consisting of δ-MnO$_2$ nanosheets deposited by in-situ growth on hollow carbon nanofibers (MnO$_2$/HCNFs) using the hydrothermal method and assembled them as an asymmetric supercapacitor (ASC) coin cell. Originating from its distinctive hollow structure, MnO$_2$/HCNFs obtained 293.6 F·g$^{-1}$ specific capacitance at 0.5 A·g$^{-1}$ in 1 M Na$_2$SO$_4$ electrolyte, which was higher than non-hollow structure. When the high operating voltage window was up to 2 V, this ASC cell held a 63.9 F·g$^{-1}$ specific capacitance. Moreover, the as-obtained ASC cell exhibited 35.1 Wh·kg$^{-1}$ high energy at 497.3 W·kg$^{-1}$ power density and maximum 8.78 kW·kg$^{-1}$ power density at 16.1 Wh·kg$^{-1}$ energy density with excellent stability. All of these demonstrated MnO$_2$/HCNFs electrodes have stronger competition than MnO$_2$/SCNFs elec-
Figure 5: The electrochemical properties of the device were tested using two-electrode system for the CFF/MnO$_2$ composite that synthesized with 4 h reaction time. (A) CV curves of the device collected in different scan voltage windows, (B) CV curves at different scan rate, and (C) Ragone plots of the fabricated device.

trodes in electrochemical properties, thus rational idea and manufacture of electrode materials with nanoarchitecture can help improve the use ratio of electro chemical active substances, thus greatly improving electrochemical performance.

3.2 TiO$_2$-based composites capacitors

Recent, hydrogenated TiO$_2$ has stimulated extensive research due to its low cost, high electrical conductivity, outstanding rate capability, and excellent stability. Herein, Pham et al. [76] fabricated hydrogenated TiO$_2$@reduced graphene oxide (HTG) sandwich-like nanosheets, which were applied for high voltage and symmetric supercapacitors. The sandwich-like nanostructure was created by utilizing a sol-gel method of growing ultrafine TiO$_2$ nanoparticles on the surface of GO sheets, even after hydrogenation, this special nanostructure remained. Nonetheless, with the increase of the hydrogenated TiO$_2$ nanoparticles' diameter, the interval between the GO sheets was significantly changed, notably at a hydrogenation temperature of 500℃. HTG revealed 51 F·g$^{-1}$ specific capacitance at 1 A·g$^{-1}$ and 82.5% capacitance retention. Furthermore, supercapacitors which were fabricated by HTG revealed desirable long-term cycling stability (80% capacitance retention after 10000 cycles). These favorable properties indicate that HTG is such a great electrode material that can realize high-performance of supercapacitors.

Rice-like particle, which is one member of 1D nanostructure, can help enhance electron transport in the electrodes due to the short transport path. Liu et al. [77] successfully fabricated rice-like titanium oxide (TiO$_2$)/graphene hydrogel (RTGH) using a facile one-pot hydrothermal self-assembly method. Here, sodium citrate (SC), an environment-friendly coordination agent, was employed as a structure-directing agent. Owing to the covalent chemical bonding, there is a strong interaction between rice-like TiO$_2$ nanoparticles and GNs, accordingly, in contrast to pristine graphene hydrogel (GH), P25/GH, and RTGH-blank, RTGH revealed excellent physicochemical properties, such as better adsorption capacities, specific capacitance and stability. For instance, RTGH obtained a high 372.3 F·g$^{-1}$ specific capacitance in the three-electrode system and 332.6 F·g$^{-1}$ specific capacitance in
the two-electrode system at 0.2 A g\(^{-1}\) in 1M \(\text{H}_2\text{SO}_4\). Additionally, in the two-electrode mode, this RTGH had 91.4% capacitance retention after 2000 cycles and favorable stability of 75.4% after 120 hours of floating test both at 4 A g\(^{-1}\), consequently, this synthesis method can be further utilized to fabricate other metal oxide/carbon hybrid materials for supercapacitors.

Yue et al. [78] successfully assembled TiO\(_2\) nanowires-reduced GO (TiO\(_2\) NWs-rGO) nanocomposite via one-step hydrothermal synthesis. Initial mass ratio of TiO\(_2\) NPs and GO, which tremendously affected the morphologies and electrochemical properties of the synthesis, was investigated in detail. CV curves were shown in Figure 6, which revealed that when initial mass ratio of GO and TiO\(_2\) NPs (with a length of approximately 10 µm as well as a diameter of approximately 50 nm) was 4:1, TiO\(_2\) NWs-rGO could obtain the best performance: this nanocomposite possessed 572 F g\(^{-1}\) specific capacitance at a current density of 1 A g\(^{-1}\) using 6 M KOH aqueous solution as electrolyte and a high rate capability with a capacitance retention ratio of approximately 84% after 5000 charge/discharge cycles at 10 A g\(^{-1}\) (Figure 7). This research effort reveals that rational coupling some promising materials such TiO\(_2\) and rGO mentioned above can further enhance electrochemical performance of supercapacitor electrodes.

4 Conducting polymers-carbon capacitors

4.1 Composite capacitors with CNTs

Wang et al. [80] developed the flexible PANI-CNT@ZIF-67-CC as supercapacitor electrode. Here, PANI was electrodeposited on CNT@ZIF-67/CC for the first time. The active materials were the PANI-CNT@ZIF-67 and flexible collector electrode was the flexible CC. After characterizing the obtained electrode, Figure 8 demonstrated that porous ZIF-67 was enclosed by a mess of CNT and partial CNT went through the ZIF-67, which highly enhanced the electroconductivity of the electrode. Owing to the synergistic effect of PANI (including prominent electrical activity, desirable pseudo capacitance and favorable chemical doping/undoping) as well as CNT@ZIF-67-CC (covering large
Figure 7: Electrochemical performance. A: GCD curves of the TiO$_2$ NWs-rGO nanocomposites at 1 A g$^{-1}$. B: Specific capacitance of the TiO$_2$ NWs-rGO nanocomposites at 1 A g$^{-1}$. C: GCD curves of the TG-1:4 at various current densities. D: specific capacitances of the TG-1:4 at various current densities

Figure 8: SEM images of (A) PANI-CC, (B) PANI-CNT@ZIF-67-CC, (C) ZIF-67-CC and (D) PANI-ZIF-67-CC

specific surface area, hierarchical porous nanostructures and outstanding electroconductivity), this synthetic PANI-CNT@ZIF-67-CC supercapacitor electrode held satisfying properties. The capacitance retention of this electrode could remain 83% even after 1000 cycles at 0.5 mA cm$^{-2}$, and specific capacitance was about 3511 mF cm$^{-2}$.

Wang et al. [81] demonstrated three-dimensional nitrogen-doped porous activated carbon monoliths (3D-NDP-ACMs) with regulative macro-mesopores and controllable morphologies through a two-step template-free route, which exhibited high capacitance properties as electrode materials. The preparation was shown in Figure 9, during thermal decomposition, the CNTs formed 3D-NDP-ACMs by shrinking the outer layer of PAN, thus providing additional pores. The overall characteristics of the 3D ordered porous structure and the high wettability of nitrogen functional groups could improve the surface utilization rate of carbon materials and facilitate the diffusion of ions in the electrolyte. The significant capacitive properties of this novel bulk material were due to the synergistic effect between conducting CNTs and the pseudo-capacitive behavior of pan derived nitrogen functional groups. These methods have universality and extensibility, and are suitable for the production of novel activated carbon monolithic materials extracted from polyimide, polyvinyl alcohol, cellulose and other polymers.
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**4.2 Composite capacitors with graphene**

Li *et al.* [84] prepared a novel 3D RGO/CNS/PANI ternary nanocomposite and employed them as supercapacitor electrode materials. Here, ammonium persulfate ((NH₄)_2S₂O₈, APS) was added to a mixture of GO, aniline and CNS in an ice bath, and subsequently Zn powder coated on the surface of CNTs, the polyaniline/carbon nanotubes/graphene/polyester textile electrode revealed high electrochemical properties. At the current density of 1.5 mA cm⁻², the capacitance of the maximum area was 791 mF cm⁻². Even after 3000 charge-discharge cycles, the capacitance retention rate of this composite electrode could still reach up to 76% in contrast to previously reported PANI electrodes with fast decay. In addition, the special electrode structure revealed strong stability under the mechanical bending and stretching conditions.

Recent developments in high-performance electrodes have created an interest in the development of various renewable energy storage systems. Malik *et al.* [83] utilized plasma enhanced chemical vapor deposition to synthesize vertically aligned CNT arrays on horizontally aligned CNT sheet for the first time. This innovative design combined excellent electrical conductivity and flexibility of CNT sheets with high surface area of CNT arrays. Electrodeposition of freestanding N-doped CNT(NCNT) sheets controlled deposition of polyaniline, thus enabled rapid charge transfer. In addition, the CNT core could offer reinforcement to the PANI coating, which obviously improved conductive polymer’s long-term cycling stability. Figure 10a was the SEM image of NCNT sheet coated with polyaniline (30 cycles), and Figure 10b and 10c demonstrated NCNT maintained high capacitance and desirable long-term cycling stability, respectively.

[Jin *et al.* [82] constructed a three-dimension (3D) conductive network by CNTs and GNs on the polyester fabric via a “dipping-drying” process and electrophoretic deposition method, which significantly increased the electron transportation rate and reduced the electrolyte ion-diffusion path. The resulting composite fabric provided a promising substrate for flexible supercapacitor’s textile-based electrode preparation. When PANI was further coated on the surface of CNTs, the polyaniline/carbon nanotubes/graphene/polyester textile electrode revealed high electrochemical properties. At the current density of 1.5 mA cm⁻², the capacitance of the maximum area was 791 mF cm⁻². Even after 3000 charge-discharge cycles, the capacitance retention rate of this composite electrode could still reach up to 76% in contrast to previously reported PANI electrodes with fast decay. In addition, the special electrode structure revealed strong stability under the mechanical bending and stretching conditions.

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was added to RGO to acquire RGO/CNS/PANI composites. During the reduction process from GO to RGO, the concomitant CNS was embedded between RGO nanosheets at the same time, effectively preventing RGO nanosheets from agglomerating as a "spacer". Specific capacitances of RGO, RGO/CNS, PANI, RGO/PANI and RGO/CNS/PANI at different current densities were shown in Figure 11, due to the excellent rate capability of EDL capacitors as well as high capacitance of pseudocapacitors from PANI [85, 86], the resulting RGO/CNS/PANI revealed the largest capacitance of all the tested composites. In addition, the capacity retention was 72% at 10 A·g⁻¹ and after 1000 charge/discharge cycles, the capacity was 86%. The preparation and capacitance application of ternary nanocomposites were reported for the first time, and this work reveal RGO/CNS/PANI ternary nanocomposites are one of the most promising materials for supercapacitor electrodes.

![Figure 11: Specific capacitances of RGO, RGO/CNS, PANI, RGO/PANI and RGO/CNS/PANI at different current densities](image)

Yu et al. [87] demonstrated a novel method to resultant graphene/polyaniline paper (GPp) and employed it as freestanding supercapacitor electrodes. The preparation process of GPp was demonstrated in Figure 12. Aniline pretreatment of GO in the first step was the key to grow high-density PANI nanoarrays onto GO nanosheets. During the process of sequent polymerization, GO nanosheets heterogeneously grew on surfaces and homogeneous grew in the solution, as a result, PANI nanoarrays and PANI nanofibers were fabricated, respectively. In the resultant GPp, PANI nanoarrays created a 3D network to guarantee GPp’s self-supported, and PANI nanofibers constructed important pores and channels. This unique composite structure not only optimizes the structure of PANI, but also highlights electron transfer and diffusion of electrolyte, which realized the excellent electrochemical performance of GPp electrode.

Wan et al. [88] demonstrated a facile scale-up process to fabricate a novel type of RGO/polypyrrole/cellulose (RPC). This RPC paper electrode obtained a low sheet resistance of 1.7 Ω·s·q⁻¹, a high areal capacitance of 1.20 F·cm⁻² at a discharge current of 2 mA·cm⁻², desirable long-term cycling stability with a capacitance retain 89.5% after 5000 times cycles and prominent mechanical flexibility. In addition, an all-solid flexible layered SSC device prepared by RPC-2.5 and H₃PO₄/PVA gel electrolyte realized a high areal capacitance of 0.51 F·cm⁻² and a superior energy density of 1.18 mWh·cm⁻³ under the condition of 0.1 mA·cm⁻². All of these results indicate that RPC-2.5 is suitable substance for the fabrication of multifarious high-performance supercapacitor with a high value of specific capacitance compared to several pure RGO.

Ensafi et al. [89] fabricated a thermally reduced graphene oxide/polymermelamine formaldehyde nanocomposite (TRGO/PMF) by a polymerization method which was much simpler than previous methods for polymer synthesis [90, 91]. During the synthesis process, the added GO dispersed in the structure of the polymer and transformed to RGO, which resulted in an increase in the capacity and conductivity of PFM. Then, the performance of TRGO/PMF was evaluated, galvanostatic charge–discharge curves were shown in Figure 13, TRGO/PMF electrode obtained a specific capacitance of 2270 F·g⁻¹ at 1.0 A·g⁻¹. Additionally, Figure 14 manifested that the Bode phase angle plot was about 80° at the tails, which was nearly an ideal capacitor. All of these results revealed TRGO/PMF holds great potential as a candidate electrode material.

5 Conclusion

This review summarizes the latest research on supercapacitor electrodes based on carbon materials. Carbon materials have been widely employed for fabricating SC electrodes owing to their satisfactory corrosion resistance, low density, excellent stability and low cost, and all-carbon composite materials can hold larger specific surface area and higher electrical conductivity. However, all-carbon materials reveal low power density and low cycling life. To overcome these defects, metal oxides are introduced, metal oxides are easy to synthesize and hold outstanding electrochemical properties. Hence, carbon materials are utilized as the scaffolds, metal oxides are deposited in situ
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Figure 12: Preparation process of GPP

Figure 13: The GCD at current densities of (A) 1.0 A·g⁻¹, (B) 1.5 A·g⁻¹, (C) 2.0 A·g⁻¹, and (D) 4.0 A·g⁻¹ were obtained in the presence of various amounts of GO; (E)–(G) show the specific capacitance vs. the current density curves obtained from (A)–(D)
on the surfaces of carbon materials. Furthermore, conducting polymer is another promising candidate for supercapacitor electrode, when compositing with carbon materials, drawbacks of pure PANI are solved, at the same time, electroconductivity, thermal stability and specific capacitance are significantly improved. All of these results demonstrate that composite materials can improve the performance of supercapacitor electrodes that one single material never could.

However, there are still a number of riddles that need to be solved. It’s no doubt that compositing carbon materials with other active basic will result in better performance of supercapacitors. In the future, carbon materials can be combined with other materials such as metal sulfides to find more application areas. We do hope this review will inspire more and more researchers to throw themselves into the fabrication of high performance supercapacitors to face the energy dilemma.

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