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A review of modification of carbon electrode material in capacitive deionization

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Capacitive deionization (CDI) technology has attracted wide attention since its advent and is considered as one of the most promising technologies in the field of desalination and ion recycling. It is constructed with an electric field by applying a low voltage of direct-current to make ions migrate directionally in solution to achieve the purpose of ion separation and removal. The performance of CDI is heavily dependent on the electrode material. Carbon is widely used as CDI electrode material because of its lower price and better stability. To enhance the adsorption capacity, extensive research efforts have been made for the modification of carbon material. In this review, we enumerate and analyze four modification methods of carbon material including element doping, metal oxide modification, chemical treatment and surface coating. The influence of each modification method on CDI performance is concluded in the perspective mechanism and some constructive advice is put forward on how to effectively enhance the performance of CDI by the decoration of carbon materials.

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1. Introduction

Ions dissolved in water are very difficult to remove just by physical methods. Up to now, electro dialysis (ED),¹ multi-stage flash (MSF),² multi-effect distillation (MED)⁴ and reverse osmosis (RO)³ are the primary physical means for ion removal. However, due to the limitation of cost and scope of application, many existing energy-intensive technologies have not been used in a large scale in industry, *e.g.* microbial desalination,⁵ ion concentration polarization,⁶ capacitive deionization (CDI)^{7,8} and so on.

CDI is an emerging electrochemical desalination technology. When compared with other technologies, it has attracted much more attention because of its unique advantages such as environmental benignity, lower energy consumption, higher efficiency and simple operation. CDI devices work at a low direct-current voltage (usually lower than 2 V), which ensures their safety and lower energy consumption. Actually, in terms of economic efficiency, Welgemoed and Schutte conducted a simulation experiment and ran a 3785 m³ per day CDI desalination unit,⁹ while the desalination capacity of typical RO plant is 100 000 m³ per day. They found that under the circumstances of 2000 parts per million (ppm) feed solution, the desalination cost for CDI was \$0.11 per m³ against \$0.35 of

RO, indicating that economic efficiency of CDI is also applicable in high concentration solutions. By changing the preparation methods and experimental conditions, the adsorption capacity of different ions can also be changed and the goal of specific adsorption can be achieved. Seo *et al.* studied the adsorption force of CDI on divalent ion by preparing graphene nano-flakes.¹⁰ In terms of difficulty of ion removal, AlMarzooqi *et al.* proved that CDI can remove 85% of divalent ions in water, showing the potential for drinking water treatment.¹¹ CDI has been proven to be available in brackish water desalination.⁷⁻¹² In addition, CDI can also be used in medicine, chemistry, microbial fuel cells and so on.¹³⁻¹⁶

Theories and experimental results both showed that porous carbon materials generally possessed high specific surface areas and excellent capacitive properties, which led to better CDI performance. CDI theory is similar to that of supercapacitors.⁷ Up to now, the core of CDI technology is to seek the maximum adsorption capacity per unit mass and the minimum energy loss power from the perspective of theory and properties. In terms of experience and theory, the ideal CDI electrode materials require the following properties: (1) large specific surface area for adsorption; (2) the stability and applicability of pH, voltage and ion concentration in different areas; (3) high speed ion mobility, *i.e.* conductivity; (4) good plasticity and expandability; (5) suitable pore size distribution structure; (6) abundant and readily available natural or artificial low-cost materials; (7) stable and reusable physical and chemical properties.

With the rapid development of CDI electrode materials, research work emerges endlessly. In previous literature,

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researchers have analyzed their progress in terms of activated carbon,^{17–19} mesoporous carbon,^{20–22} carbon aerogels,^{23,24} carbon nanotubes,^{25–27} graphene,^{28,29} *etc.*, and reviewed the various carbon-based materials^{8,30} as well as graphene materials³¹ for CDI. However, there is still a review gap in the field of modification of carbon electrode materials. To fill this vacancy, we provide a comprehensive summary of common modification methods in the aspects of element doping, metal oxide doping and polymers, and put forward some prospects. This article summarizes the existing successful modification methods of carbon electrode materials in CDI.

2. History of CDI materials

Compared with other traditional desalination methods, CDI is a new effective technology.⁸ Studies on CDI can trace back to the 1960s and the conception was first reported by Murphy and Caudle. They designed a flow through mode whose electrode sheets consisted of porous activated carbon electrodes for water desalination. Then in 1970s, Johnson *et al.* studied the reversible process of CDI and proposed a reversible electrosorption model. In this theory, ions hidden in the electrode sheets could be released by removing electrical field.³² On the other hand, they gave a conclusion that the application of high-surface area electrodes would be beneficial to efficient desalination. However, there are still some limitations in carbon materials, such as poor stability, single composition, and too irregular structure. Scientists have tried abundant modification methods of carbon materials in the course of past few decades to seek more suitable modified materials for practical production. Thus, more and more materials such as carbon nanotubes, carbon aerogel, ordered mesoporous carbons, carbon nanofiber and graphene have been applied to CDI technology, which lead to a great boost in the field.¹⁸ Here some typical materials will be introduced with their properties and application examples.

Generally, activated carbon (AC) is the most widely used electrode material for CDI electrode because of its higher-capacity of environmental contaminants and cheaper price.³³ But lower conductivity and higher electrical transfer resistance limit its further development in CDI.³⁴ AC is usually combined with binders and conductive additives to fabricate CDI electrode. Hou *et al.* found that ten percent of PVDF (a kind of binder) could achieve a better result in CDI experiment.³⁵ Furthermore, some chemical modification methods by introducing hydrophilic reactive functional group can also greatly improve the electrosorption capacity of activated carbon electrodes.³⁶

Carbon Nanofiber (CNF) is a fibrous one-dimension carbon nanomaterial, which could be obtained by electrospinning. In recent years, more and more people have focused concerns on the application in the field of supercapacitors for its high surface area, high electrical conductivity and unique filamentous structure.³⁷ In 2014, Chen *et al.* fabricated phenolic resin-based CNF through electrospinning and one-step carbonization and applied the electrode materials to CDI.³⁸

Carbon aerogel (CA) possesses unique strengths such as good conductivity, continuous and favorable pore distribution,

as well as high surface area.³⁹ In 1990, Farmer *et al.* discovered carbon aerogel and applied this material to CDI electrodes. From then on, searching new carbon materials with a high specific surface area has become a focus of concern among scientist.⁴⁰ Although carbon aerogel has unique good properties, the complex processing technology and high cost limit its development.

Carbon nanotube (CNT), a kind of one-dimensional nanomaterial with hollow structure, is well known for its strong active functional groups and proper pore structure. In 2005, carbon nanotubes were first used for CDI according to Dai and his team's study.⁴¹ Arvind *et al.* found amazing adsorption effect by opening the termination of carbon nanotubes involve oxidation to get more organic functional groups.⁴²

Graphene is a new two-dimensional carbon nanomaterial with single atomic layer, possessing great development potentiality. Graphene was first gained by Andre *et al.* in 2004 and was used into CDI in 2009. It has some derivatives such as graphene oxide and reduced graphene oxide which are usually loaded on other electrode materials. Yu *et al.* incorporated graphene oxide with carbon nanofibers. The desalination performance was improved for the enhancement of electrical conductivity and mesoposity ratio.⁴³

Ordered mesoporous carbons (OMCs), possessing ordered pore size and less ion transfer resistance, are promising candidate for CDI electrode. Zou *et al.* first adopted OMCs to CDI. They found this electrode showed better ion removal performance and faster desorption speed compared with activated carbon.⁴⁴ Li *et al.* prepared mesoporous carbon electrode with larger surface area by adding $\text{NiSO}_4 \cdot 6\text{H}_2\text{O}$, which performed a better electrosorption performance.⁴⁵

As the research continues, researchers found that excellent CDI electrode materials require the properties of large specific surface area, suitable pore size, high hydrophilicity and high conductivity.^{34,46} To improve those properties, various modification methods have been developed. The timeline is shown in Table 1.

3. Modification methods

3.1. Heteroatom doping

Carbon materials, with the advantages of low cost, light weight, large specific surface area, corrosion resistance, oxidation resistance, excellent ductility and porous structure, occupy a dominant position in CDI electrodes. However, due to their super high stability and structural framework, carbon materials often have a single element type. That makes it difficult for CDI electrode materials to break through in terms of adsorption capacity, rate and types. Fortunately, the known organic species are far more than inorganic species, and carbon is the most basic element of the organic structure skeleton, that is to say, it has strong plasticity and compatibility. That makes it possible for scientists and researchers to dope other ions or elements into carbon materials to improve their electrosorption performance. Actually, heteroatom doping is one of the most used modification methods for carbon materials. It is worth mentioning that experiments showed that not all elements are



Table 1 History of CDI materials

Time	Event
1960s	Murphy and Caudle first proposed the concept of CDI and made CDI electrode sheets from activated carbon
1970s	Johnson <i>et al.</i> proposed a reversible electrosorption model
1990	Farmer <i>et al.</i> discovered carbon aerogel and applied this material to CDI electrodes
2005	Dai <i>et al.</i> applied carbon nanotubes into CDI for the first time
2006	Han-jun <i>et al.</i> introduced hydrophilic reactive functional group in activated carbon electrodes to improve the adsorbing capacity
2008	Zou <i>et al.</i> first adopted OMCs to CDI and found better ion removal performance and faster desorption speed compared with activated carbon
2009	Li <i>et al.</i> obtained the mesoporous structure with a larger surface area by adding NiSO ₄ ·6H ₂ O to ordered mesoporous carbons
2009	Graphene was first used into CDI
2010	Li <i>et al.</i> used graphite to produce a GO and applied it into CDI
2014	Chen <i>et al.</i> fabricated phenolic resin-based CNF and applied the electrode materials to CDI
2014	Yu <i>et al.</i> incorporated graphene oxide into carbon nanofiber to improve the conductivity and the mesoporosity of electrode materials
2015	Arvind <i>et al.</i> opened the termination of carbon nanotubes involve oxidation and found amazing adsorption effect

suitable for element doping. Non-metallic elements, such as N, P, S and Cl are commonly selected and calcined with carbon materials at high temperatures to prepare element doped electrodes. Several common heteroatom doping methods are as followed.

3.1.1. N-doped carbon material. Owing to the unique valence electron layer structure, nitrogen can form ionic bonds or covalent bonds with other atoms. This property makes it widely used in the field of CDI electrode modification. The existent forms of nitrogen in carbon materials can be divided into chemical nitrogen and structural nitrogen. Chemical nitrogen exists in the form of surface functional groups, such as amino and nitrite, while structural nitrogen atoms enter the framework of carbon materials, such as pyridine nitrogen and quaternary nitrogen. Nitrogen atoms in carbon skeleton not only optimize the pore structure, but also increase the chemical reactivity and electron transport properties of carbon atoms.⁴⁷ Chemical nitrogen doping is able to improve the hydrophilicity of carbon materials, thus enhancing their catalytic and adsorption capacities.⁴⁸ Moreover, nitrogen doping can increase the active sites of carbon materials, make metal particles evenly dispersed on the surface and reduce the amount of precious metals. It can also increase the adsorption capacity of porous carbon materials for CO₂ and H₂.^{49,50} In conclusion, nitrogen doping can improve the comprehensive properties of carbon material and expand its application field.

Usually, there are three methods to dope nitrogen into carbon materials. The first one is to use nitrogen-containing monomers, such as melamine, pyrrole, sucrose and polyaniline, as precursors to prepare nitrogen-containing polymers which are then heat-treated. Directly heating nitrogen-containing biomass materials or wastes is the second way. The last one is to use NH₃ or CO(NH₂)₂ as nitrogen source to post-treat the material, so that specific nitrogen-containing

groups will be introduced into the carbon material. The effects of different nitrogen doping methods on electrochemical properties and electroadsorption capacities of carbon materials will be described in detail below.

When nitrogen doping is used in capacitive deionization, it can effectively enlarge the surface pore size of carbon materials, *i.e.* enlarging the specific surface area. High specific surface area is one of the most remarkable features of carbon materials, which represents high adsorption capacity. Hence, nitrogen content is a major consideration of characterization. Zhao *et al.*⁵¹ used polystyrene (PS) as hard template, and dopamine hydrochloride as carbon and nitrogen sources to prepare nitrogen-doped porous carbon microspheres (N-PHCS). Its uniform, spherical and hollow carbon structure was observed in field emission scanning electron microscopy (SEM) and transmission electron microscopy (TEM) images (Fig. 1(a)–(c)). The nitrogen content, determined by X-ray photoelectron spectroscopy (XPS), was 2.92% (Fig. 1(d)). The result of the SAC curve and current transient of N-PHCS electrodes in 500 mg L⁻¹ NaCl solution also indicated that its electrochemical performance was fantastic (Fig. 1(e)). A 500 mg L⁻¹ NaCl aqueous solution was treated at a flow rate of 40 mL min⁻¹ at 1.4 V. The specific electrosorption capacity (SAC) was 12.95 mg g⁻¹ and its stability was perfect. Li *et al.*⁵² prepared similar nitrogen-doped materials. Nitrogen-doped cluster porous carbon materials (NCPC) with layered hollow nanostructures were fabricated under various temperatures by carbonizing dopamine/nano-ZnO composites. They possessed good wettability interconnection layer structure, enabling rapid ion binarization. Among them, the sample carbonized at 900 °C (NCPC-900) showed the best porous structure. Its specific surface area was 1357 m² g⁻¹ and the maximum adsorption capacity was 11.98 mg g⁻¹. In terms of nitrogen content, NCPC-700 reached the highest of 9.2%. Besides, Liu



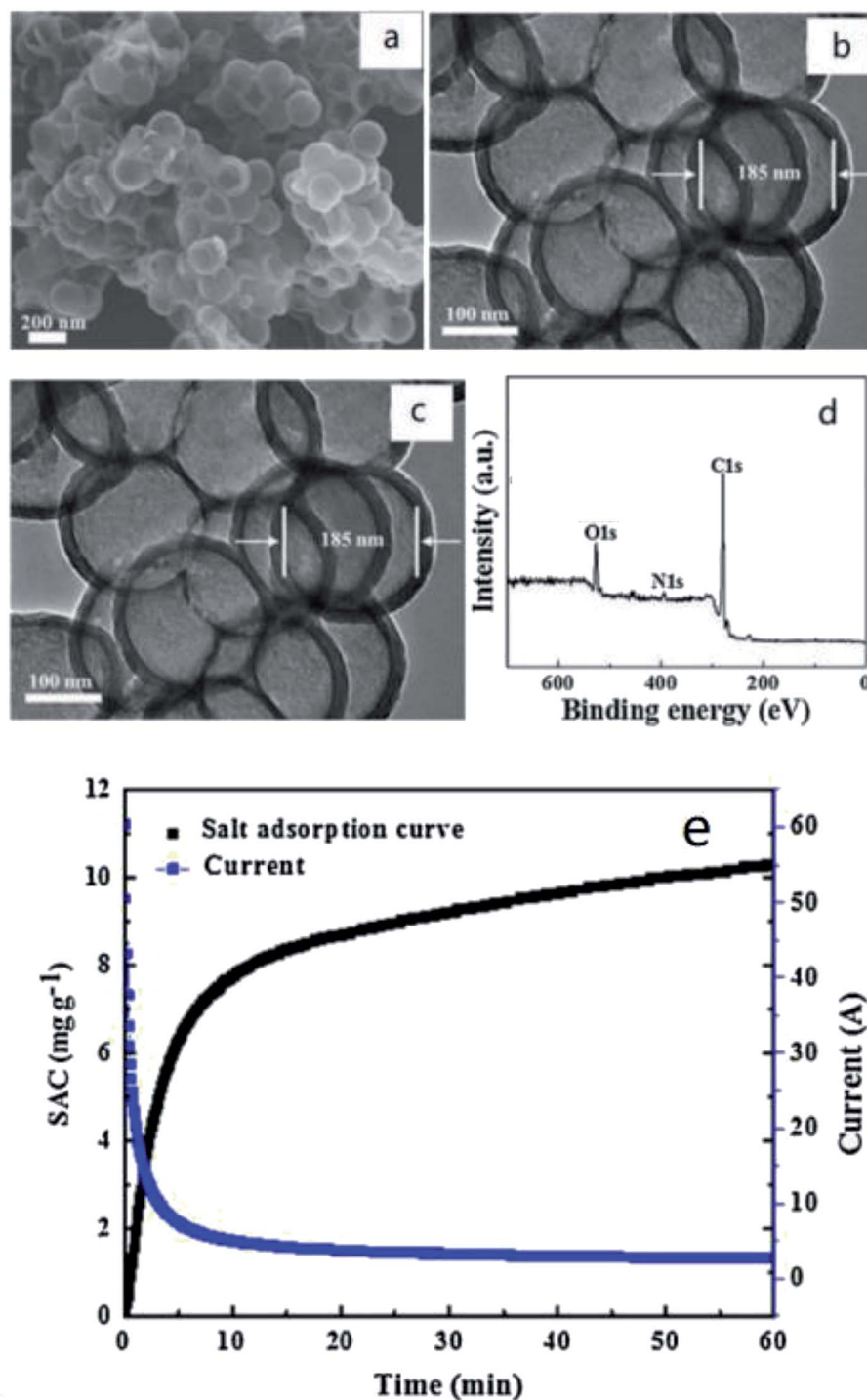


Fig. 1 (a) SEM (b) TEM and (c) HRTEM images of the N-PHCS; (d) XPS spectra of the N-PHCS sample; (e) the SAC curve and current transient of N-PHCS electrodes in 500 mg L^{-1} NaCl solution at the voltage of 1.2 V with a flow rate of 40 mL min^{-1} ; reproduced with permission from ref. 51.

*et al.*⁵³ prepared nitrogen-doped porous carbon microspheres (NPCS) by microblog-assisted rapid preparation method. In this process, sucrose was used as a precursor. Through a series of characterization and experimental measurements, it was found that NCPS treated at $1000 \text{ }^\circ\text{C}$ possessed the highest electroadsorption capacity of 14.91 mg g^{-1} in 1000 mg L^{-1} treated solution.

Gao *et al.*⁵⁴ synthesized graphene polyhedrons directly from pyrolytic imidazolite skeleton (ZIF-8). Characterization results showed that it had close SP-2 skeleton edges. When compared with carbon polyhedrons (CPs) and nitrogen doped carbon polyhedrons (NCPS), besides the benefits of $1134 \text{ m}^2 \text{ g}^{-1}$ remarkable specific surface area and considerable graphitization degree, the conductivity and cycling stability of nitrogen-



doped graphitic carbon polyhedrons (NGCP) were also distinguished. In CDI experiment, NGCP exhibited a desalination capacity of 17.73 mg g^{-1} with 500 mg L^{-1} NaCl solution at the voltage of 1.4 V. Duan *et al.*⁵⁵ and Liu *et al.*⁵⁶ also applied ZIF-8 to the preparation of analogous carbon materials and obtained good results (Fig. 2). The above experimental results proved that nitrogen doping was an effective method to improve the electroadsorption performance as well as recycling stability of carbon materials.

In addition, calcination of nitrogen-containing biomass at high temperature under pure nitrogen condition is a common method to prepare carbon materials. It can effectively determine the pore size of carbon materials and the distribution of nitrogen elements in the materials. Classically, Zhao *et al.*⁵⁷ fabricated nitrogen-doped porous carbon (NPC) by simple pyrolysis method, using soybean hull as raw material and KHCO_3 as catalyst. Then NPC sulfonate group was functionalized in aryl diazonium salt solution, leading to a final specific surface area of $844.0 \text{ m}^2 \text{ g}^{-1}$ and nitrogen content of 1.66 at%. The electroadsorption capacity of this material was 15.8 mg g^{-1} , and the adsorption rate was $0.37 \text{ mg (g min)}^{-1}$ due to the high surface negative charge density.

The last kind of nitrogen doping aims at introducing specific nitrogenous groups, such as NH_3 and NR_4 , onto the surface of carbon materials. Yang *et al.*⁵⁸ used aryl diazonium salt solution and acetone mixed with APTes to introduce $-\text{SO}_3^{2-}$ and $-\text{NH}_3^+$

groups into nanotubes to overcome co-ion effect. Characterizations and electroadsorption results indicate that the functionalized-CNTs electrodes enhanced the salt-removing performance. Later, Gao *et al.*⁵⁹ introduced ethylenediamine into the surface of microporous spectracarb carbon cloth and used it as the cathode electrode. Meanwhile, the anode carbon material was treated with nitric acid to increase its negative surface charge. The maximum adsorption capacity was 5.3 mg g^{-1} at 1.1 V working voltage and characterization results showed that the functional groups derived from ethylenediamine increased the microporosity of carbon materials. In the latest research, Ahmed G. *et al.*⁶⁰ modified activated graphene electrodes with polyatomic ions (NR_4^+). Carboxymethyl cellulose (CMC) was negatively charged and quaternary ammonium cellulose (QMC) was used as the disinfectant. The obtained materials asym-QC-3DAPGr (3DPGr: 3D porous graphene) were treated with 300 mg L^{-1} NaCl solution at 1.4 V cell voltage. And adsorption capacity per unit is up to 18.43 mg g^{-1} . In another aspect, the non-toxic water-soluble adhesive also reduces the interface resistance and improves the flexibility of the electrode.

Nitrogen doping is often combined with graphene. Graphene is a two-dimensional carbon nano-material with hexagonal honeycomb lattice composed of carbon atoms and sp^2 hybrid orbital. Its excellent conductivity and pore structure make it a new favorite of electrodes choosing in CDI, and it is often associated with nitrogen doping to form two-dimensional

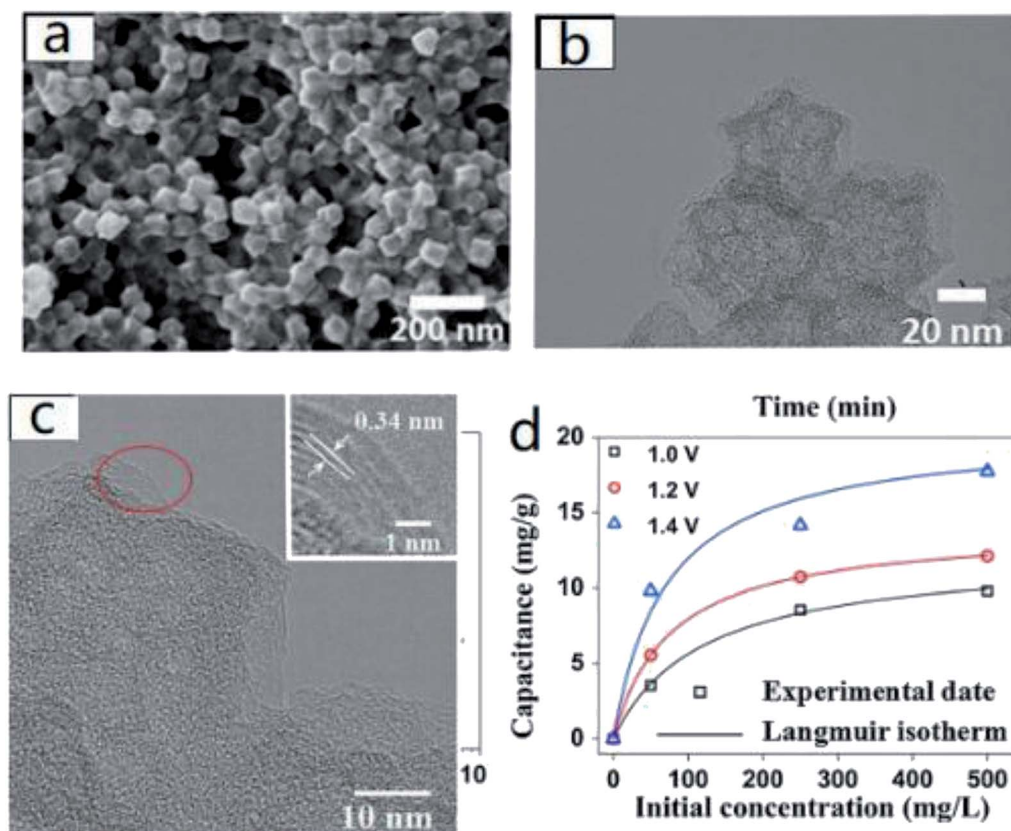


Fig. 2 (a) The SEM (b) TEM and (c) HRTEM image of NGCPs; (d) electroadsorption capacitance and Langmuir isotherm of NGCPs at various voltage in NaCl solutions with initial concentration ranging from 50 to 500 mg L^{-1} . Reproduced with permission from ref. 54.



carbon nanomaterials. In fact, as early as 2015, Xu *et al.*⁶¹ have prepared a novel nitrogen-doped graphene sponge (NGS), which has high specific surface area of 526.7 m² g⁻¹, and its designed structure can be also fabricated. Eventually, when treated with 500 mg L⁻¹ NaCl solution, the adsorption capacity attained 21 mg g⁻¹, which was also the highest at that time. In the same year, Amiri *et al.*⁶² published an article to demonstrate that highly-crumpled nitrogen-doped graphene and highly-crumple and few-layered graphene have enormous potentiality in industry desalination. Liu *et al.*⁶³ then applied this material to the rapid removal of heavy metal ions (Pb²⁺, Cd²⁺, Cu²⁺, Fe²⁺) in water, obtaining a removal rate of 90% in the range of 0.05–200 ppm and perfect recycling abilities. In the latest research, Zhou *et al.*⁶⁴ prepared two-dimensional sandwich graphene/nitrogen-doped carbon nanoparticles composites using graphene/porous SiO₂ composite as a template, ethylene diamine and carbon tetrachloride as carbon sources. It exhibits high capacitance up to 189 F g⁻¹ at a current density of 0.1 A g⁻¹. This further proves that nitrogen-doped graphene has broad application prospects in the field of CDI.

3.1.2. Multielement doping. In addition to N, the most commonly used element are S and P to modify carbon material. In the latest research, Ma *et al.*⁶⁵ introduced S atoms into the graphene skeleton and proved that this operation can effectively improve the capacitance performance of graphene. A few years early, Porada *et al.*⁶⁶ used salt template synthesis to prepare biomass precursors and sulfur-doped microporous carbon. Among their materials, NDC-Cs-900 (NDC: nitrogen doped carbon, Cs: Cs acetate) shew the best performance. The maximum specific surface area was 2830 m² g⁻¹ and the adsorption capacity could reach up to 15.0 mg g⁻¹ at the voltage of 1.2 V. In addition, they also compared it with the undoped P carbon material and found that heteroatom-doped carbon material was a promising CDI material. It also indicated that the high heteroatom content may be related to low charge efficiency. Meanwhile, they are often closely associated with nitrogen doping. Wang *et al.* introduced sulfur doping into carbon fibers and made pore and graphitized edges, which not only improved cycle stability, but also facilitated fast capacitive sodium storage.

Apart from sulfur, phosphorus is another doping element that is often used in carbon modification. Li *et al.*⁶⁷ compared the P and none element doped carbon nanofibers aerogels in 1000 mg L⁻¹ NaCl solution, and the adsorption capacity of undoped P was 12.81 mg g⁻¹ while that of (P)-doped carbon nanofiber aerogels (P-CNFA) was 16.20 mg g⁻¹. In addition, the nitrogen and phosphorus dual doped graphene aerogels were prepared to expand the capacitance and cycling stability of the materials. In this way, Li *et al.*⁶⁸ greatly improved the performance of carbon anodes used in the construction of electrochemical capacitors. Experiments show that the double doping of nitrogen and phosphorus makes graphene aerogel (NPGA) have high sodium storage capacity and wide application prospect in CDI field, and its cycling stability is excellent. On the level of practical application, Han *et al.*⁶⁹ firstly used phytic acid as P source, chitosan as N source, mixed with gallic acid to deal with graphene oxide and succeed in preparing highly

hydrophilic N, P co-doped three-dimensional hierarchical carbon architectures. Due to its three-dimensional structure, the material has a high adsorption capacity of 26.8 mg g⁻¹ at an applied voltage of 1.2 V.

To some extent, it seems that the more types of element were used to dope, the more remarkable the effect is. Of course, it is also inseparable from the reasonable element collocation and the selection of appropriate element sources. In 2018, Zhang *et al.*⁷⁰ succeeded in making N, P, S co-doped hollow carbon polyhedral. They chose poly(cyclotriphosphazene-co-4,4'-sulfonyldiphenol) as carbon sources and N, P, S co-doping sources, ZIF-8 as structural templates, which also acted as additional N-doping sources. TEM results showed that ZIF-8@PZS-C (poly(cyclotriphosphazene-co-4,4'-sulfonyldiphenol) coated zeolitic imidazolate framework-8) owned fabulous pore structure (Fig. 3(a) and (b)). Ultimately, the super-high adsorption capacity of 22.19 mg g⁻¹ was achieved in the treatment of 500 mg L⁻¹ NaCl solution (Fig. 3(c)). It is reasonable to believe that more and more attention will be paid to multielement doping.

3.1.3. Ion-doped carbon material. Aside from the common doping of non-metallic elements, ion doping refers to dope metal elements such as Li⁺, Fe³⁺ and Cl⁻ into carbon materials. Generally speaking, ion doping can effectively improve the specific capacitance of material, reduce the charging resistance of the composite electrode and improve its stability and regeneration efficiency, and enhance the electrochemical performance of the composite. Those functions indicate ion doping an effective method to improve the electroadsorption capacity of CDI electrode materials. Recently, Zhang *et al.*⁷¹

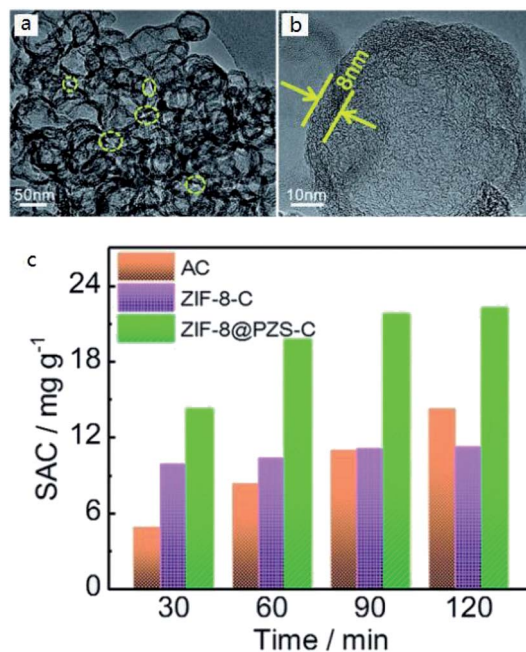


Fig. 3 (a) TEM images and (b) HRTEM image of ZIF-8@PZS-C; (c) plots of SAC vs. deionization time of various electrodes in 500 mg L⁻¹ NaCl solution at 1.2 V with a flow rate of 50 mL min⁻¹. Reproduced with permission from ref. 70.



prepared large-scale lithium ion-doped graphene/carbon nanofiber porous by electrospinning homogeneous LiNO_3 /graphene oxide/polyacrylonitrile nanofibers and carbonize it at 900°C . Its porous structure provides high electroadsorption capacity. The maximum efficiency can be as high as 87% and this material has broad application prospects in capacitive deionization.

In addition, ion-doped methods can be also associated with polypyrrole (PPy) and CNT. Chloride (Cl) and dodecylbenzene sulfonate (DBS) are both commonly used ion doping reagents. Normally, ion-doped CDI electrodes are affected by ion species, acidity and basicity in solution and experimental conditions. Wang *et al.*⁷² investigated the fouling characteristics of ion-doped PPy/CNT composite electrodes in the process of CDI by a half cycle running mode. It was found that the addition of Ca^{2+} or Fe^{3+} in the solution would alkalize the regeneration process during desalination, thus affecting the adsorption performance of CDI. In the aspect of organic pollution, the addition of sodium humate significantly reduced the desalination performance of CDI batteries, because the high concentration of sodium humate would destroy the structure of the electrode layer and further destroy the stability of the electrode. Ma *et al.*⁷³ explored the effects of temperature and pH on ion-doped PPy/CNT composite electrodes. Under the condition of $\text{pH} = 7$, $T = 15^\circ\text{C}$, the adsorption capacity was the largest, reaching 75% or more. Meanwhile, strong acid and alkali will aggravate the oxidation reaction on the surface of the electrode, and the increase of temperature will lead to the hydrophilic-hydrophobic transition on the surface of the electrode, which results in the decrease of the adsorption capacity. In 2016, Cai *et al.*⁷⁴ measured the electroadsorption capacity of PPy-DBS/CNT and PPy-Cl/CNT composite electrode. The electroadsorption capacity is $40.8\text{--}72.36\text{ mg g}^{-1}$, which directly proves the strong electroadsorption of ion-doped composite carbon materials. In another aspect, Cai inferred that the composite electrodes may have the good ability of ion selective adsorption for cations or anions.

3.2. Metal oxide-modified carbon material

For the reason of synergistic effects, the combination of carbon and metal oxides improves the specific capacitance, making them potential materials for capacitive deionization.⁷⁵ Moreover, the addition of metal oxide can prominently change some physicochemical properties of carbon material, like wettability, surface area and zeta potential, which may contribute to the improvement of capacitive deionization performance.

3.2.1. TiO_2 -modified carbon material. TiO_2 is an n-type semiconductor oxide, with qualities of high stability, high hydrophilicity, low cost as well as eco-friendliness. As early as 2003, TiO_2 was utilized to modify carbon electrodes for CDI.^{17,76} By introducing TiO_2 , the wettability and electrochemical properties of carbon electrodes was reported to increase, resulting in the enhancement of electroadsorption capacity.^{77,78}

High hydrophilicity of TiO_2 is due to abundant hydroxyl groups on its surface. Those Ti-OH groups can act as electroadsorption sites on the surface of carbon electrode, which is

beneficial to the CDI performance. Kim have *et al.*⁷⁷ fabricated one of the TiO_2 coated carbon electrodes. As shown in Fig. 4, the wettability of carbon electrode was highly enhanced by coating with TiO_2 . The TiO_2 coated carbon electrode showed an electroadsorption capacity of 17 mg g^{-1} which was nearly twice as high as that of carbon electrode. This improvement was resulted from the easier accessibility of ions and water from solution to the surface of electrode due to enhanced wettability. Recently, Kang *et al.*⁷⁹ synthesized TiO_2 -modified activated carbon fibers (ACF). From the Nyquist impedance plots, it can be calculated that the value of charge transfer resistance decreased with TiO_2 doping amount, implying that the introduction of TiO_2 promotes ion migration and diffusion on the surface of carbon electrode. Consequently, the highest electroadsorption capacity reached up to 10.6 mg g^{-1} , which was 71.9% higher than that of the unmodified ACF electrode.

The strong hydrophobicity of CNT dramatically limits its CDI performance. With high hydrophilicity, TiO_2 is promising for improving this situation. Li *et al.*⁸⁰ reported a TiO_2 /CNTs composite electrode for CDI. This electrode displayed an enhanced electroadsorption capacity of 4 mg g^{-1} which was twice as high as that of pristine CNTs. Feng *et al.*⁸¹ recently fabricated TiO_2 /CNTs membrane electrodes by atomic layer deposition method. The deposition of TiO_2 was reported to transform CNT from strongly hydrophobic to hydrophilic. This composite electrode showed an improved desalination capacity of 5.09 mg g^{-1} as well as good regeneration.

Cycling stability of CDI is challenged with peroxide formation and rapid oxidation of anodic electrode.⁸² To overcome this issue, Srimuk *et al.*⁸³ presented a strategy by hybridizing nano-carbon particles with sol-gel-derived TiO_2 . This carbon/metal oxide hybrid showed 90% of the initial electroadsorption capacity after 100 cycles, indicating an outstanding cycling stability. TiO_2 was confirmed to prevent the formation of local peroxide by modifying redox reactions. In their subsequent research, TiO_2 -modified activated carbon electrodes were fabricated.⁸⁴ The electrode loaded with 15 wt% TiO_2 displayed a considerable desalination capacity of 17.4 mg g^{-1} and a better cycling stability, attributed to favorable three-electron transfer reactions.

3.2.2. MnO_2 -modified electrode. Great specific capacitance (83.9 F g^{-1}) and electroadsorption capacity (5.4 mg g^{-1}), which illustrated the CDI electroadsorption efficiencies of CMK-3, the hybrid electrodes and the deionization-regeneration profile as well as deionization-regeneration cycles of the 15Mn@C-400 electrode, had been achieved by the use of MnO_2 nanoparticles decorated mesoporous carbon, reported by Zhao *et al.*,⁸⁵ who raised a substantial enhancement compared with the normal ordered mesoporous carbon due to its well coexistence. Yang *et al.* suggested that the suitable pore size distribution property and high surface area may be the main contribution of the improvement.⁸⁶ Besides the high capacitance (204.7 F g^{-1}), an effective cation intercalation and high surface adsorption capability were proven to be extra superiority. And the MnO_2 nanoparticles decorated anode made sure to avoid oxidation of carbon anodes, raised by Wu *et al.*, who chose a MnO_2 anode and an anion-selective quaternate poly(4-



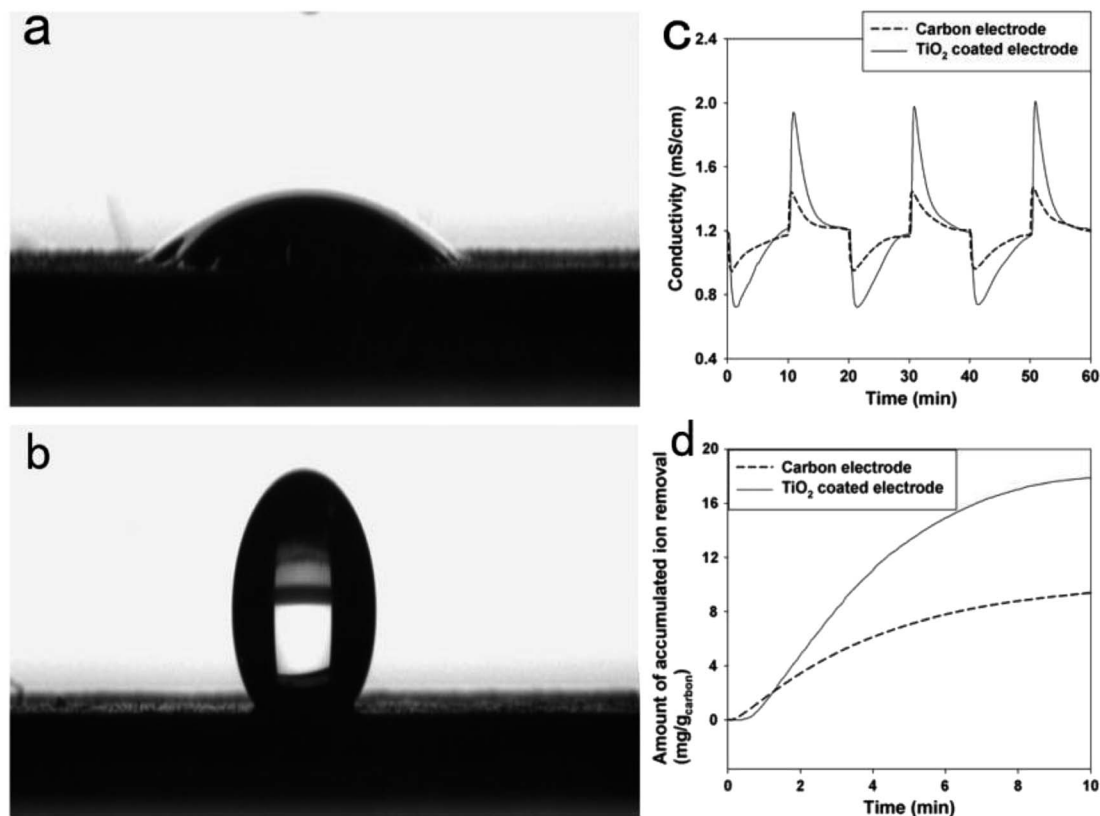


Fig. 4 Surface wetting property of (a) the TiO₂ coated electrode and (b) carbon electrode. (c) Ion removal performance. (d) Accumulated ion removal amount during the charging step (selected at the third cycle). Reproduced with permission from ref. 77.

vinylpyridine) coated activated carbon cathode in CDI, which led to a high SAC retention ratio of 95.4% after 350 adsorption-desorption cycles, comparing with normal AC cathode in Fig. 5.⁸⁷

Bryan *et al.* reported four tunnel crystal structures of manganese oxide nanowires as faradaic electrodes.⁸⁸ Two of them are square tunnel structured MnO₂ (α -MnO₂ and todorokite-MnO₂), and others have ordered and disordered unions of structural tunnels, which had been shown in SEM and cross-sectional HAADF-STEM images (Fig. 6). It was found that larger tunnels did a great favor to the removal of larger hydrated radii cations compared with the smaller ones. And the surface redox reactions as well as the intercalation of ions to the structural tunnels may be the main reason of ion removal.

In the study of Steven and Roland,⁸⁹ electroless deposition (ED) and cyclic voltammetry (CV) were different methods to modify the electrodes and fabricated the MnO₂ coated ones. Both techniques increased the electrode capacitance, however, only ED electrodes can improve the desalination performance. MnO₂ mass deposition contributed dispersedly and fully in ED modified aerogel and while creating a discrete crust in CV electrodes. Thanks to this, a great specific charge capacitance (77.6 F g⁻¹) was exhibited in a MnO₂/AC composite electrode and storage capacity in thin-film sodium manganese oxide (NMO) was increased to 170 times higher than the uncoated electrodes.⁹⁰ And according to the report of Yang *et al.*,⁹¹

Polystyrene Sodium Sulfonate (PSS) can facilitated the dispersion in composite electrodes and enhance the growth of MnO₂, resulting in a great salt removal efficiency (96.8%) and a high ion adsorption capacity (80.4 $\mu\text{mol g}^{-1}$) in MnO₂/PSS/CNTs electrodes.

3.2.3. ZnO-modified carbon material. Zinc oxide (ZnO) is an n-type broadband gap semiconductor and its unique properties enable it to grow in various forms on the surface of carbon materials.^{92,93} Myint *et al.* firstly applied ZnO nanomaterials to CDI technology and made continuous breakthroughs in this technology from 2012 to 2015.⁹⁴⁻⁹⁷ They focused on the application of zinc oxide nanocomposite electrodes grafted on activated carbon cloth in the removal of ions from brackish water by capacitive deionization. It was found that the polarization of ZnO nanorods increased the strength and penetration depth of electric field between the electrodes, thus improving the capacitance and charge efficiency to a certain extent. Besides, uniformly distributed electric field lines between two electrodes brought about faster ion adsorption rates as well as reduced electrode saturation time, resulting in $\sim 45\%$ increase in the average desalination efficiency at all applied potentials.

Additionally, Liu *et al.* used polytetrafluoroethylene emulsion to mix AC particles and nano ZnO to prepare ZnO/AC composite electrodes.⁹⁸ It was reported that -ZnO/AC||AC capacitor showed a very stable CDI performance with significant electrosorption capacity of 9.4 mg g⁻¹ and charge



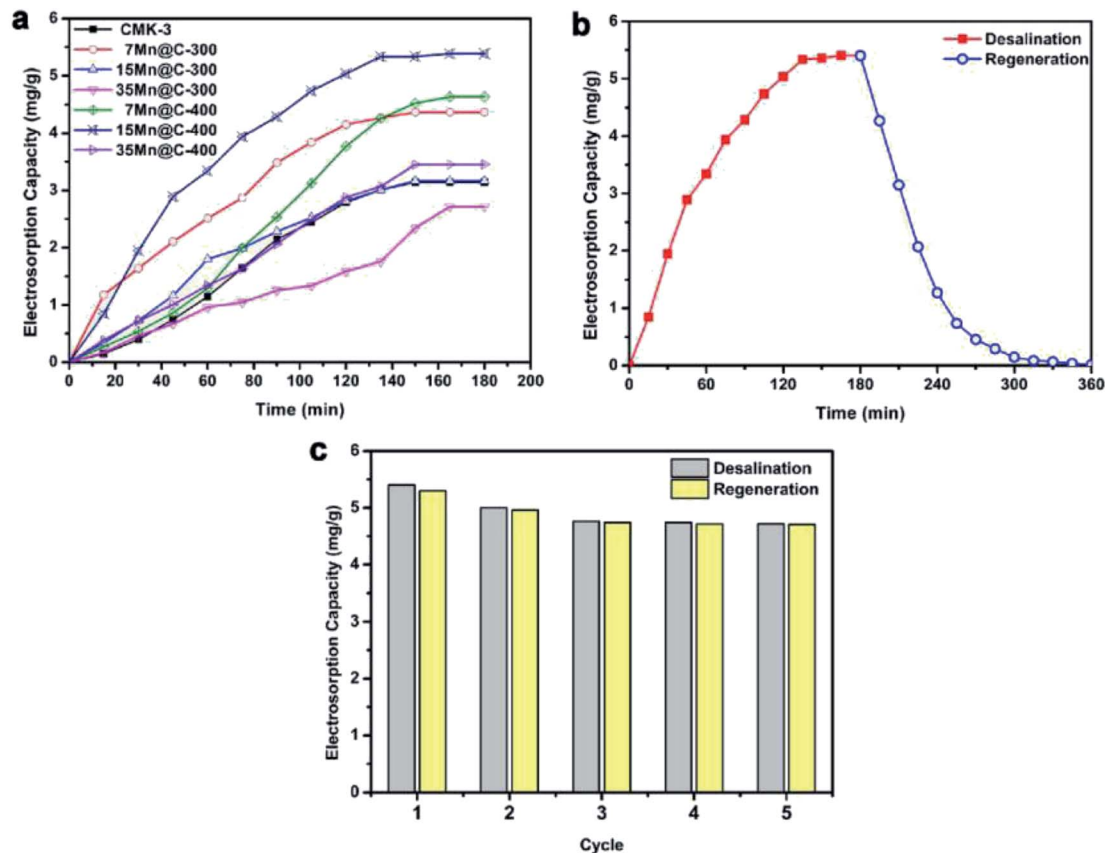


Fig. 5 (a) CDI profiles of CMK-3 and $x\text{Mn@C-T}$ hybrids under 1.2 V; (b) deionization-regeneration profile and (c) deionization-regeneration cycles of the 15Mn@C-400 electrode. Reproduced with permission from ref. 87.

efficiency of 80.5%, while $+\text{ZnO/AC}||\text{AC}$ capacitor showed declining desalination behavior after several charge-discharge cycles, as evident from Fig. 7. The different CDI properties on negative and positive electrodes were due to the changes of pH during charging process, which led to zeta potential variation of ZnO.

3.2.4. Other metal oxide-modified carbon material. Wouters *et al.*⁹⁹ synthesized carbon fiber sheets coated with thin-films of SiO_2 and $\gamma\text{-Al}_2\text{O}_3$. It showed a desalination efficiency of Ca^{2+} greater than 90%, which was 30% higher than that of uncoated carbon. This promotion could be attributed to the increase of specific surface area (SSA) and wettability of the composite, as well as the decrease of the resistance to ion transport process. It was also found that $\gamma\text{-Al}_2\text{O}_3$ exhibited a positive charge while SiO_2 a negative charge. When applied potential had the same polarity as that of the coatings, the electrodesorption capacity for counter-ions could get enhanced.

Additionally, Deen *et al.*¹⁰⁰ reported graphene/tin dioxide nanoparticles composites (Gr/SnO_2) fabricated by microwave irradiation. The incorporation of SnO_2 led to a high specific capacitance of 323 F g^{-1} , and prevented graphene sheets from aggregating, which brought about a significant increase in surface area. In CDI experiments, electrode with 15 wt% SnO_2 showed excellent cycling stability as well as considerable desalination efficiency of 83%.

Fe_3O_4 is a chemically stable and environmentally friendly material, with high pseudocapacitance. $\text{Fe}_3\text{O}_4/\text{RGO}$ nanocomposites synthesized *via* hydrothermal method were reported by Li *et al.*¹⁰¹ Cyclic voltammetry curve of this composite revealed a typical electrical double layer capacitive behavior and an enhanced specific capacitance, probably attributed to the surface hydroxylation of Fe_3O_4 . The maximum electrodesorption capacity of $\text{Fe}_3\text{O}_4/\text{RGO}$ nanocomposites calculated by Langmuir model was 8.33 mg g^{-1} , which was twice as high as that of RGO (4.63 mg g^{-1}).

3.3. Chemical treatment of carbon material

3.3.1. Acid treatment. Generally, acid treatments are associated with surface oxidation of carbon materials, which lead to huge improvement in oxygen-containing functional surface groups, such as carboxyl, carbonyl, hydroxyl, quinone, lactone, *etc.*¹⁰² These functional groups are vital for the adsorption of alkaline-earth metals due to chelating effect. Besides, electrochemical properties and wettability of carbon materials will also benefit from these groups.

Acidic oxidizing agents such as concentrated nitric acid and concentrated sulfuric acid have been extensively used for the acid treatment of carbon materials. Huang *et al.*¹⁰³ reported nitric acid-modified AC electrodes for capacitive deionization.



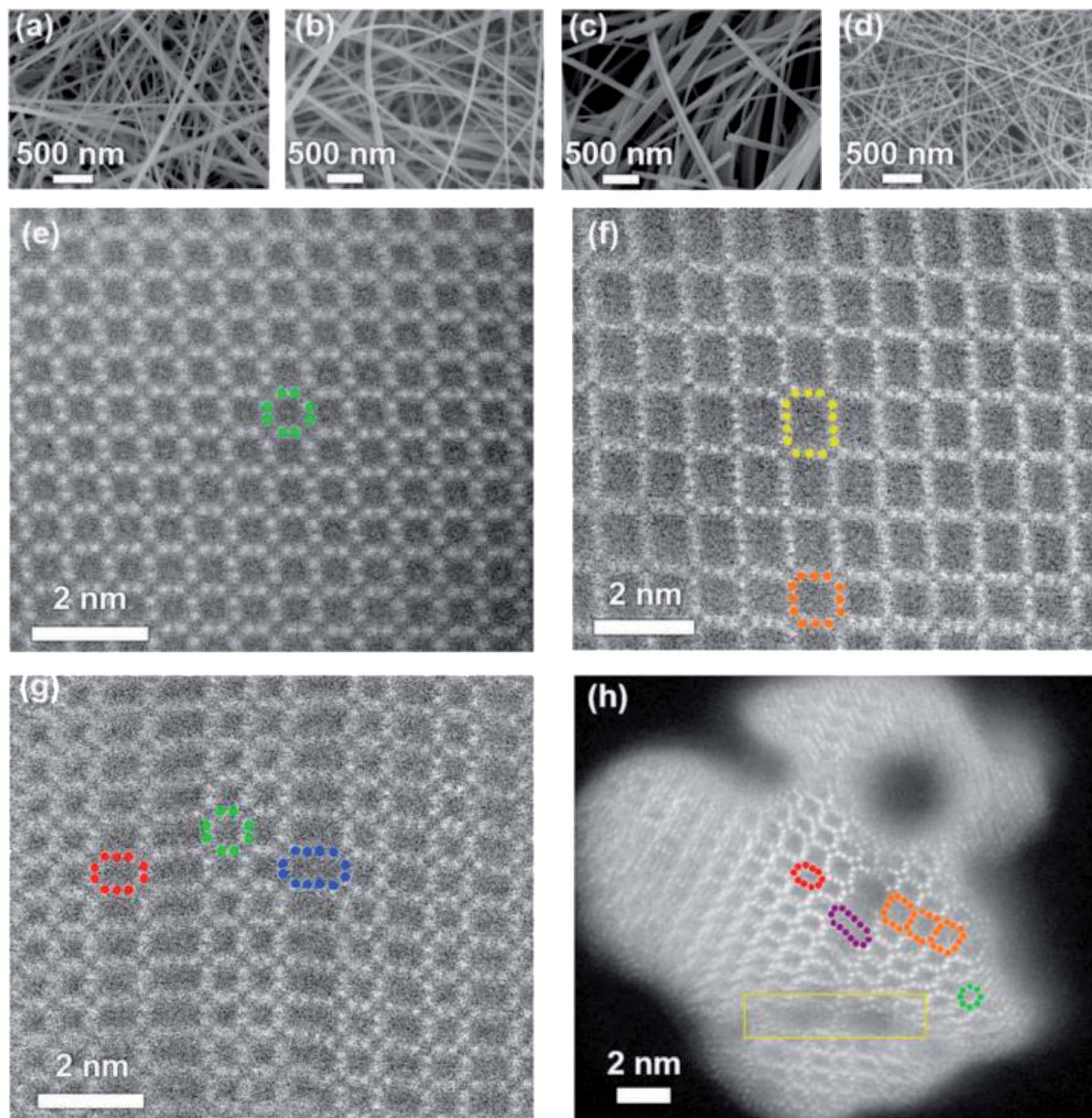


Fig. 6 SEM (a–d) and cross-sectional HAADF-STEM (e–h) images of α -MnO₂, Tod-MnO₂, $2 \times n$ -MnO₂, and hybrid-MnO₂: (a and e) α -MnO₂, (b and f) Tod-MnO₂, (c and g) $2 \times n$ -MnO₂, and (d and h) hybrid MnO₂. Reproduced with permission from ref. 88.

Attributed to improved capacitance of AC caused by the increase of oxygen-containing functional surface groups, CDI experiments showed that desalination efficiency had increased by 15%. To prepare sulfuric acid functionalized AC (FAC), Niu *et al.*¹⁰⁴ modified activated carbon with 98% sulfuric acid *via* hydrothermal method. Compared to pure AC, FAC exhibited an improved electroadsorption capacity of 3.54 mg g⁻¹. Wu *et al.*¹⁰⁵ treated activated carbon fiber with 6 M nitric acid to reduce co-ion expulsion effect. Electroadsorption capacity of this ACF-HNO₃ could reach up to 12.8 mg g⁻¹. In this work, carboxylic groups were grafted onto the ACF surface, which positively shifted the potential of zero charge (E_{pzc}) of ACF. It could be observed that ACF-HNO₃ is unable to adsorb many anions even at a positive voltage, indicating its excellent cation selectivity. Thus, the co-ion expulsion effect which decreases adsorption capacity was limited.

Though traditional oxidation treatment of carbon materials can promote electroadsorption capacity, it is destructive to the surface area and porous structure. In the research of Maroto-Valer *et al.*,¹⁰⁶ nitric acid treatment of activated carbon was reported to decrease the total pore volume and BET surface area by 8.8% and 9.2% respectively. A promising method to improve this status is oxygen plasma treatment, which can oxidize the carbon external surface without modifying the internal structure effectively. Talemi *et al.*¹⁰⁷ reported oxygen plasma modified carbon electrodes. In their study, electroadsorption capacity was enhanced but there was no obvious specific surface area change. Also, the SEM images of electrodes at different levels of magnification shows no distinct variation on the surface after oxygen plasma treatment.

3.3.2. Alkali treatment. Alkali represented by potassium hydroxide usually has the function of chemical activation. As



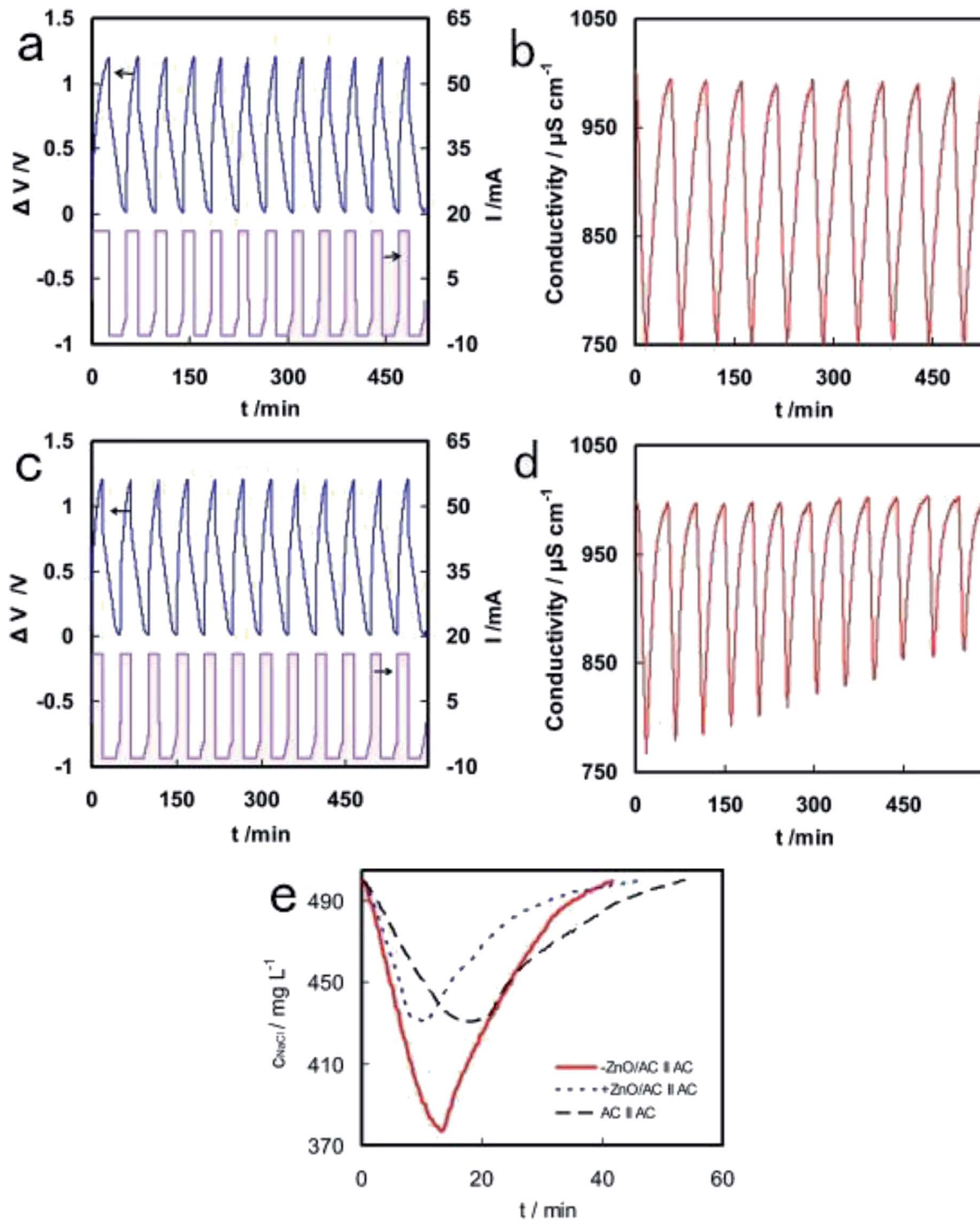


Fig. 7 Cell voltage/current curves (a and c) and conductivity variation (b and d) with time on $-ZnO/AC||AC$ (a and b) and $+ZnO/AC||AC$ (c and d) in the continuous charge–discharge process. (e) Conductivity variations of $-ZnO/AC||AC$, $+ZnO/AC||AC$ and $AC||AC$ in CDI experiments. Reproduced with permission from ref. 98.

a highly efficient activator, potassium hydroxide has been widely used in the preparations of porous carbon materials with abundant pore structure and high specific surface area. Yeh *et al.*¹⁰⁸ fabricated coconut shell-based activated carbons with various mesoporous ratio by manipulating the impregnation ratio of KOH, which was shown in Fig. 8(a). From Fig. 8(b), it can

be observed that in CDI experiments, the electrode with 70.7% ratio of mesoporosity (AC-1-2.0) showed a considerable electroadsorption capacity of $9.72 mg g^{-1}$, higher than that of micropore-dominant electrodes, which indicated the association between mesoporosity ratio and electroadsorption capacity.



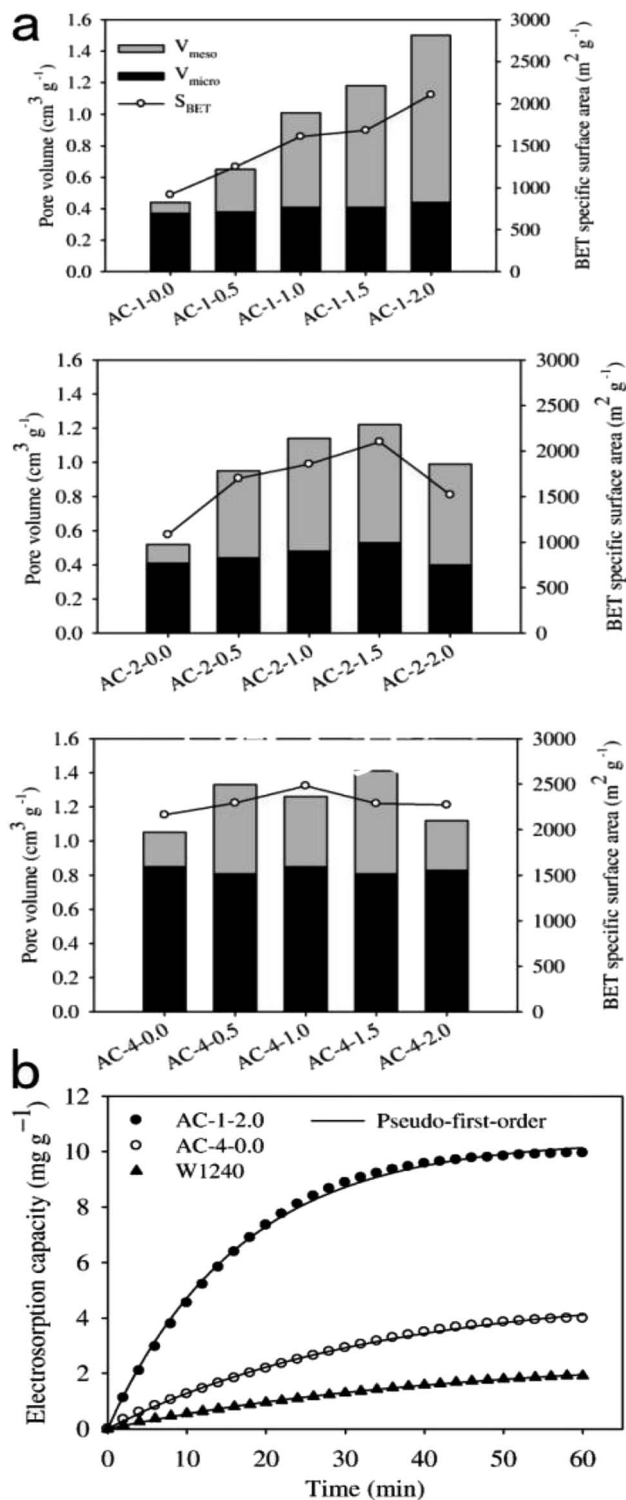


Fig. 8 (a) Pore characterization of the activated carbons prepared with different impregnation ratios of KOH. (b) The electrodesorption kinetics of AC-1-2.0, AC-4-0.0, and W1240 electrodes. Reproduced with permission from ref. 108.

Dehkhoda *et al.*¹⁰⁹ revealed a novel way to tailor porous activated biochar. By controlling drying conditions and carbonization temperature of KOH activation, biochars with predominantly mesoporous, microporous or combined porous

structure could be obtained. Electrochemical analysis showed that electrodes with majorly microporous structure had high total capacitance ($222\text{--}245\text{ F g}^{-1}$) but large electrode resistance, while mesopore-dominated electrodes (>90% of total pore volume) possessed ideal capacitive behavior. CDI experiments suggest different ranges of desalination applications, with small hydrated ions (Na^+ , K^+ , Cl^- , Mg^{2+} , Cu^{2+}) utilizing microporous electrodes and larger ions (Cr^{3+} , Zn^{2+} , Cd^{3+}) utilizing mesoporous electrodes.

Besides optimizing pore structures, KOH also plays an important role in introducing hydrophilic functional groups. Ji *et al.*¹¹⁰ reported KOH-modified activated carbons with higher surface area and increased surface hydrophilic functional groups, indicating the potential of KOH for improving wettability. Wang *et al.*¹¹¹ modified activated carbon with different mass concentrations of KOH solution. The electrode modified with 5% KOH exhibited a maximum specific capacitance of 108.8 F g^{-1} as well as 23% increase in desalination efficiency comparing to non-modified AC. The contact angle analysis and FTIR spectra image indicated that this promotion resulted from the introduction of hydroxyl functional groups on the electrode surface. Lee *et al.*¹¹² studied the surface modification of carbon felts by KOH and HNO_3 . It was observed that KOH increased the BET surface area of the electrode, while HNO_3 brought damages on the pore structure. Thus KOH-modified carbon felts displayed higher desalination efficiency of 83.3% compared with HNO_3 -modified carbon felts (65.4%), although the oxygen-containing functional groups formatted by HNO_3 was much greater.

3.3.3. Other chemical treatment. As a hydrophilic group, sulfonic group ($-\text{SO}_3\text{H}$) is expected to improve the wettability of carbon materials. In the work of Niu *et al.*¹⁰⁴ mentioned above, the introduction of $-\text{SO}_3\text{H}$ was confirmed, which also resulted in the promotion of CDI performance. Qian *et al.*¹¹³ prepared sulfonated graphene (SG)-coated CNFs as MCDI electrodes. SG was abundant in negatively charged functional groups. By introducing it, CNFs were tuned from hydrophobic to hydrophilic. Thus in batch CDI experiments, this SG-CNF electrode exhibited an electroadsorption capacity of 9.54 mg g^{-1} and increased by 22.0% in charge efficiency. Moreover, by introducing sulfonic groups, the co-ion repulsion effect can be reduced. Min *et al.*¹¹⁴ recently reported activated carbon materials modified by TiO_2 coating and Tiron sulfonating (AC-Ti-S). The introduction of negatively charged sulfonic groups positively shifted the Epzc, as shown in Fig. 9. When AC-Ti-S was used as negative electrode, the co-ion repulsion effect could be reduced. Consequently, this AC-Ti-S electrode showed an electroadsorption capacity of 7.6 mg g^{-1} , which was 1.5 times as high as pristine AC.

Additionally, working voltage of CDI is dependent on the potential difference between the Epzc of cathode and anode. More negative Epzc of the cathode and more positive Epzc of the anode are required to enlarge the potential difference. Frequent oxygen-containing functional groups are negatively charged, which can positively shift the Epzc. Therefore, to further enlarge the potential difference, positively charged groups like $-\text{NH}_3^+$ are of great importance. Gao *et al.*⁹⁹ synthesized amine-modified



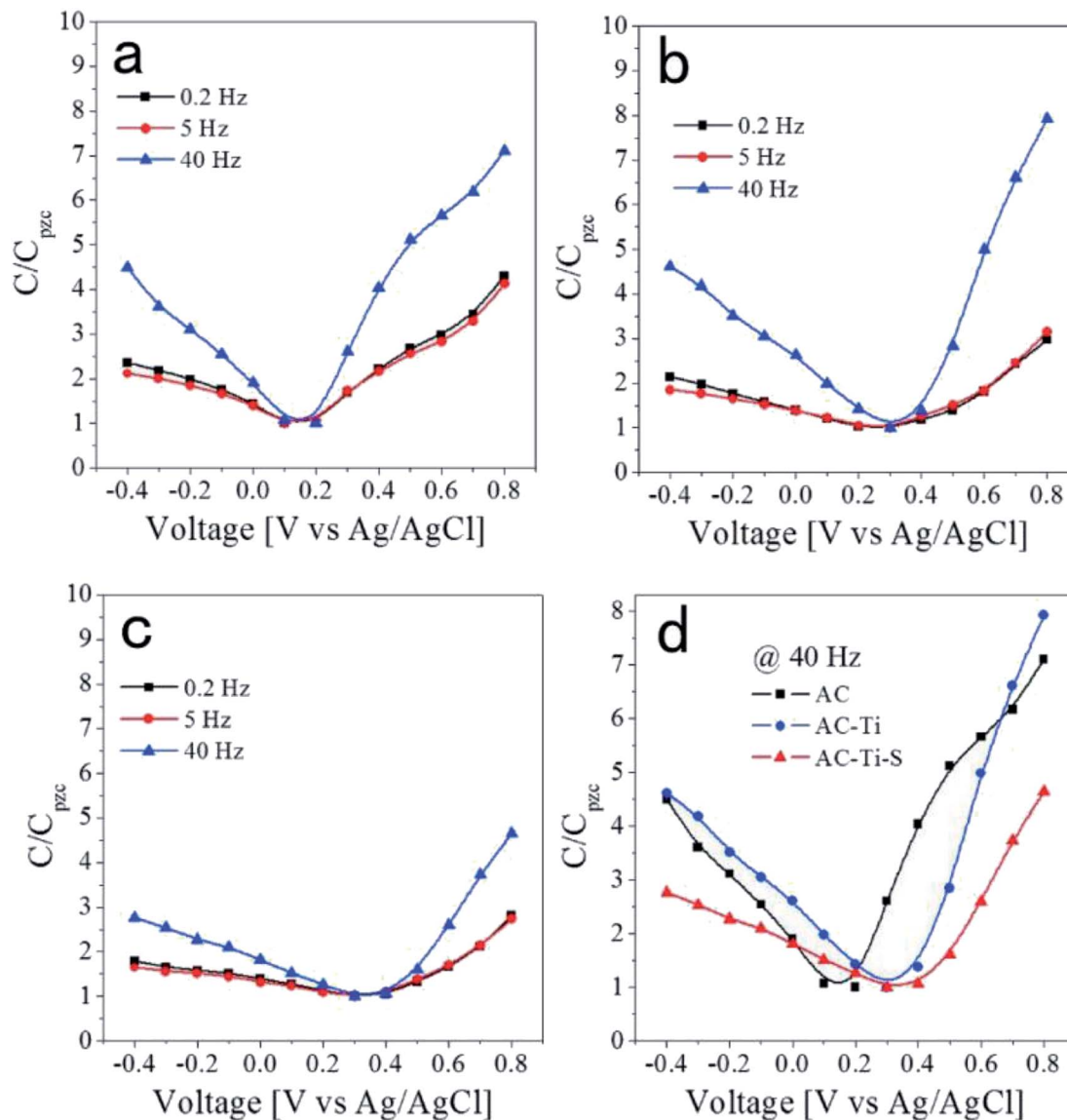


Fig. 9 Relative capacitance at various frequencies as a function of the applied voltage for the (a) AC, (b) AC-Ti and (c) AC-Ti-S electrodes. (d) The comparison of the C/C_{pzc} values at 40 Hz. Reproduced with permission from ref. 114.

microporous carbon cloths. To attach $-NH_2$ functional groups onto the surface, nitric acid-treated carbon cloth was further modified by ethylenediamine. In neutral solution such as NaCl, $-NH_2$ was protonated to $-NH_3^+$, resulting in the increase of surface positive charges which led to promoted desalination capacity of 5.3 mg g^{-1} . Additionally, ion selectivity of $-NH_3^+$ was utilized to promote CDI performance, reported by Liu *et al.*¹¹⁵ They synthesized a novel 3D graphene electrode grafted with amine and sulfonic functional groups. Those charged groups could serve as ion-selective functional coatings to reduce the co-ion effect. As a consequence, this electrode displayed an excellent electrosorption capacity of 13.72 mg g^{-1} and a high charge efficiency of 85%.

3.4. Electrode coated in CDI

Working voltage of CDI depends on the potential difference between the E_{pzc} of cathode and anode. Cation-exchange

polymer and metallic oxide account for a large part of materials used for coating the electrodes. The latter has been introduced in the above article, so we will focus on different cation-exchange polymers coating on electrodes in this part.

The cation-exchange polymer coated electrode, widely used in membrane capacitive deionization (MCDI), is found more efficient than most of commercial carbon electrodes.¹¹⁶ Among a large amount of efforts over the years to prolong the service life of a CDI device and decrease the interfacial resistance of the membrane capacitive deionization, coating layers of cation-exchange polymer on the electrodes seems perfectly solving the problems.¹¹⁷

Kim and Choi¹¹⁸ reported a method to fabricate polyvinyl alcohol (PVA) and sulfosuccinic acid (SSA) coated electrodes, which was considered as a milestone in the synthesize of a cation-exchange polymer coated electrode. The salt-removal efficiencies were observed to increase from 50–67% to 75–



85%. The similar material, reported by Kim *et al.*,¹¹⁹ was fabricated by blending PVA with SSA and poly(styrene sulfonic acid-co-maleic acid) (PSSA_MA) (PVA/PSSA_MA/SSA), in which the removal efficiency of these electrodes were different due to disparate crosslinking synthesis temperatures and SSA contents. The PVA/SSA coating showed high adhesion to the electrodes, and the interfacial resistance of membrane capacitive deionization was much lower than the commercial ones, so the removal efficiency increased to 88% in a mixed solution with 300 ppm. According to the images of the surfaces and cross-sections of the electrode and ion exchange polymer-coated electrodes, the carbon electrode surfaces were better coated with both cation and anion exchange polymers.

A new composite, reported by Yan *et al.*, was fabricated *in situ* polymerization with polymer polyaniline and activated carbon. The combination of materials provided a considerable improvement of conductivity, in the meantime decreasing the number of micropores. And the promotion came from the conducting chains which linked microactivated carbon granules together and blocked a large amount of micropores.

Liu *et al.* reported m-MCDI electrodes fabricated by leading the anion exchange polymer (dimethyl diallyl ammonium chloride) and cation exchange (polyethyleneimine) polymer into CNT electrodes. What illustrated detailly in the article was that due to the reduction of co-ion expulsion effect while introducing ion exchange polymers and enhancing adhesion between ion exchange polymers and electrodes, it exhibited a relatively high NaCl removal of 93%.¹²⁰ And the “co-in” effect could also be reduced by coating the sulfonated poly(phenylene oxide) and aminated polysulfone on the surface of traditional carbon electrodes.

With a new electrode, a salt removal efficiency of nearly 100% was obtained, which was illustrated. This electrode was fabricated with synthesized sulfonated poly(phenylene oxide) (SPPO) and aminated polysulfone (APSf), which realized a leap of the performance of membrane capacitive deionization (MCDI). Thanks to its coatings of sufficient thickness at 4.89 μm for SPPO and 5.16 μm for APSf, adequate anchoring of the polymers made sure there wasn't any delamination.¹²¹

Moreover, the ratio of selectivity to the degree of resistance enhancement was found to have positive correlation with the degree of promotion in MCDI performance.¹²² The negative electrode was coated with Ca-alginate in MCDI for hardness control and showed great ion removal efficient for hardness species like calcium ions, which were 44% more than that of the commercial CDI, as shown in Fig. 10.

In recent years, CNTs has been widely used in electrode coated for its special property. Wang *et al.* synthesized carbon nanotube and polypyrrole (PPy) composite electrode with the method of chemical oxidation and used sodium dodecyl benzene sulfonate as the dopant. Their experiment showed the PPy/CNT composites enhanced the capacitance of electrodes three times than traditional carbon nanotubes and considerably improved their adsorption capacity. Compared to CNT, the TEM image of PPy/CNT composites indicated that PPy/CNT composites were nanotube form with the CNT wrapped by the uniform PPy layer, as shown in Fig. 11.¹²³ Mechanism of the PPy/

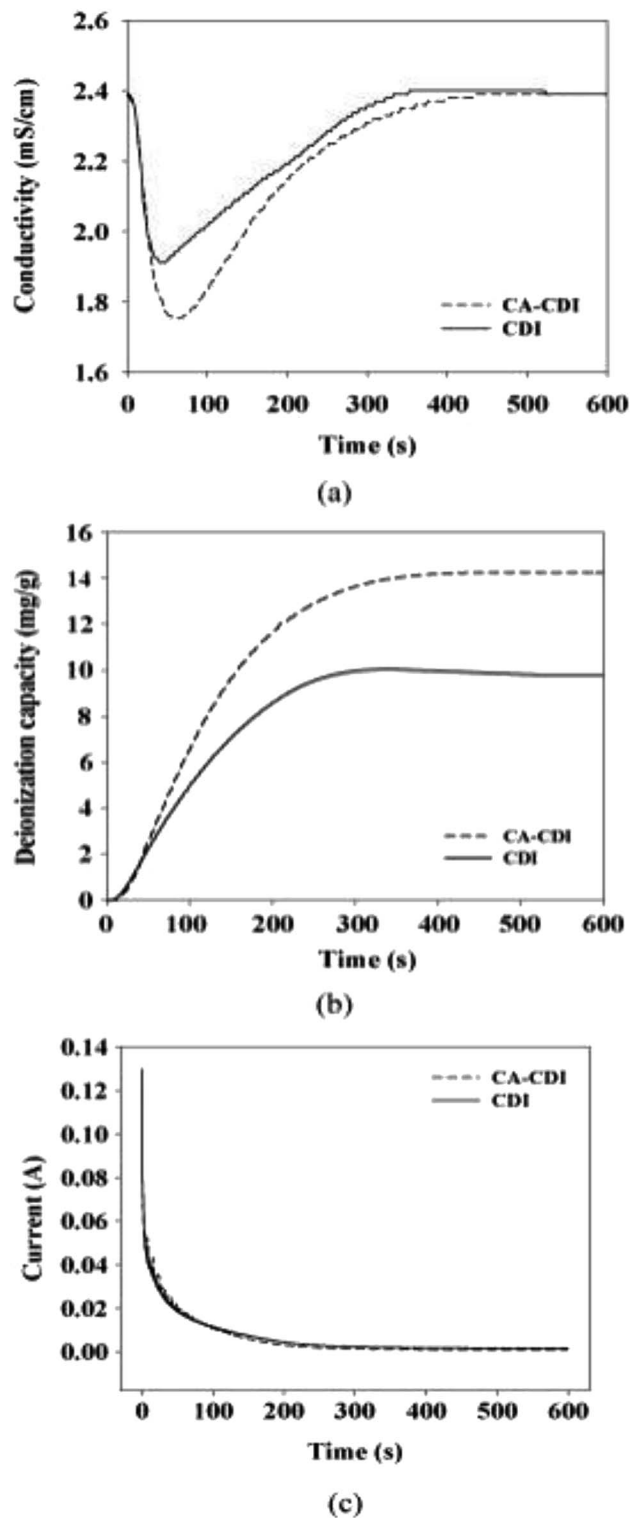


Fig. 10 (a) Conductivity profile, (b) deionization capacity and (c) consumed current of Ca-alginate coated CDI (CA-CDI) compared with the control CDI with time. Reproduced with permission from ref. 122.

CNT's satisfying performance was that the uniform and loose PPy layer resulted in the lower charging resistance of the coated electrode than that of commercial CNTs.¹²⁴

Besides the cation-exchange polymer coated electrodes, electrodes coated with other materials were also studied by



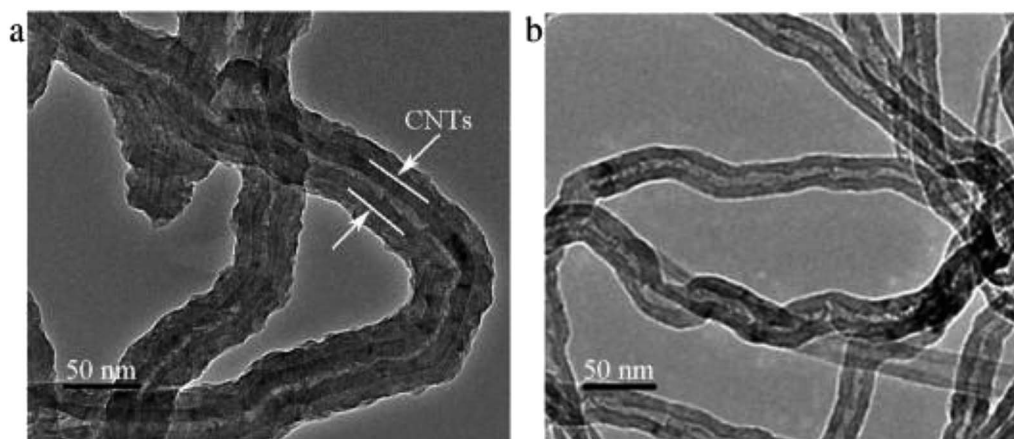


Fig. 11 SEM images of (a) PPy/CNT composites and (b) CNT. Reproduced with permission from ref. 123.

many researchers. Yoon *et al.* tested the Ag coated hybrid CDI system.¹²⁵ The enhanced specific capacity uniting the capacitance in AC electrode with the Ag mediated charge transfer reaction, leading to a significant improvement – the charge efficiency (116%), the CDI deionization capacity (188%) and rate (139%).

4. Summary

As a novel water treatment technology, capacitive deionization has drawn wide attention for the advantages of lower energy consumption and cost, higher efficiency and better regeneration. Selection and modification of electrode materials is one of the most essential factors affecting desalination performance, which has become the focus of CDI research in recent years. In this article, we summarize the modification methods of carbon materials including heteroatom doping, metal oxide modification, chemical treatment and surface coating, and analyze their mechanism for enhancing CDI performance. The electrochemical and electrosorption performances of various carbon electrodes modified by different methods are summarized in Table 2.

Generally speaking, the purpose of modification is to improve the properties of carbon electrodes, such as the porous structure, specific surface area, wettability and conductivity. Specifically, element doping is mainly used to optimize the pore structure and increase the specific surface area of the carbon materials. Nitrogen doping can also introduce nitrogen-containing functional groups to enhance hydrophilicity and conductivity. Metal oxide modification, represented by TiO₂, can considerably improve the wettability of carbon electrode by introducing hydroxyl groups on the surface. Additionally, the combination of carbon and metal oxides is able to increase the specific capacitance for the reason of synergistic effects. Acid treatment generally leads to significant increase in oxygen-containing functional surface groups. Apart from improving wettability and hydrophilicity, these negatively charged functional groups can positively shift the E_{pzc} of carbon materials, thus reducing the co-ion expulsion effect and promoting

electrosorption capacity. Alkali treatment is used to tailor porous carbon materials with different ratios of mesoporosity, due to the chemical activation function of alkali. Besides, cation-exchange polymer coating layers can reduce the interfacial resistance of MCDI which leads to the enhancement of electrochemical performance.

Although these methods can lead to an increase in CDI performance, there still exist some shortcomings. Element doping usually needs high temperature, but the specific temperatures are unpredictable thus requiring large numbers of experiment. In addition, acid treatment is destructive to the surface area and porous structure of carbon electrode due to the oxidation process. Talemi *et al.* proposed a solution of oxygen plasma modification, but the complicated operations limited its further promotion.¹⁰⁷ Moreover, the introduction of surface functional groups can accelerate the faradaic reactions which reduce the charging efficiency. By taking such issues, CDI performance will be further improved.

5. Prospects

As a new type of desalination technology developed rapidly in the middle and late 19th century, CDI has attracted the attention of scholars with its high economical value and perfect stability. However, there are still several aspects that limit the development of CDI in practical applications. The biggest problem remains that the electrosorption performance of conventional electrode materials limits the upper limit of CDI material adsorption. High specific area materials such as activated carbon, graphene, and carbon aerogel have good pore structure and electrochemical properties, but for decades, the space for increasing the unit adsorption capacity is still very limited. This is the basic reason for various modification methods proposed to increase the unit adsorption amount. In fact, in addition to the various modification methods discussed in this paper, there is another new idea that can effectively improve the adsorption efficiency of CDI units. That is to replace the traditional electrosorption theory with a reversible chemical reaction. This theory is based on the fact that the



Table 2 Different carbon electrodes and their CDI performances

Electrode	Specific capacitance (F g ⁻¹)/scan rate (mV s ⁻¹)	Surface area (m ² g ⁻¹)	Voltage (V)	Solution (NaCl mg L ⁻¹)	Electrosorption capacity (mg g ⁻¹)	Ref.
N-PHCS	152/5	512	1.4	500	12.95	51
NCPC-900	199.0/1	1357	1.2	100	11.98	52
NPCS	290.74/1	1640.5	1.2	1000	14.91	53
NGCP	307.4/10	1134	1.4	500	17.73	54
NPC	165.2/6	1036.2	1.28	40	15.8	57
N-SC	—	1421	1.1	250	5.3	59
asym-QC-3DAPGr	98.9/5	2680	1.4	300	13.83	60
NGS	286.86/5	526.7	1.2	500	13.83	61
NDC-Cs-900	—	2830	1.2	—	15.0	66
P-CNFA	295.1/5	728.2	1.2	1000	16.20	67
N, P co-doped 3D hierarchical carbon	221/5	451	1.2	500	26.8	69
TiO ₂ /AC	107/2	2300	1.2	585	17	77
TiO ₂ /ACF	182/5	886.1	1.2	500	10.6	79
TiO ₂ /CNTs	122/1	—	1.2	500	4.3	80
TiO ₂ /AC	94/5	1749	1.2	293	17.4	84
MnO ₂ nanoparticles decorated ordered mesoporous carbon	83.9/10	1092	1.2	—	5.4	85
MnO ₂ /nanowires	—	—	1.2	878	27.8	88
MnO ₂ /PSS/CNTs	77.2/10	191.8	1.2	128	—	91
ZnO/AC	66/2	—	1.2	500	9.4	98
SiO ₂ and γ-Al ₂ O ₃ coated electrode	—	35.3	1.2	111 (CaCl ₂)	—	99
Gr/SnO ₂	323/5	—	1.4	—	1.49	100
Fe ₃ O ₄ /RGO	—	150.8	1.2	500	8.33	101
FAC	232/1	1244.7	1.2	—	3.54	104
HNO ₃ /ACF	—	1545	1.2	500	12.8	105
KOH/AC	181.8/5	2105	1.0	29	9.72	108
SG-CNF	117/2	—	1.2	400	9.54	113
AC-Ti-S	238/20	—	1.0	500	7.6	114
3DNGR-3DSGR	108.3/10	133	1.4	500	13.72	115
PVA and SSA coated carbon	97.4/5	1260	1.5	200	—	118
Dimethyl diallyl ammonium chloride and polyethyleneimine coated carbon	51.51/5	—	1.2	500	9.5	120
Ca-alginate coated carbon	—	—	1.2	1110	14.2	122
PPy coated CNT	180.61/5	—	1.4	58 440	43.99	124
Ag coated carbon	110.2/2	—	0.7	584.4	23.2	125

Faraday capacitance is much larger than the double layer capacity of the material. Tantalum capacitance refers to the underpotential deposition of electroactive materials in the two-dimensional or quasi-two-dimensional space on the surface or in the bulk phase, resulting in highly reversible chemisorption, desorption or redox reactions, which are related to the charge potential of the electrode capacitance. It can be produced not only on the surface of the electrode but also inside the entire electrode, so its development potential is enormous. For example, a Prussian blue analogue has recently been proposed as an electrode material for high-efficiency capacitive deionization. Its unique open-frame structure of large ion channels makes it an excellent material for Na⁺ intercalation and deintercalation. It is reported that the unit adsorption capacity can be up to 59.9 mg g⁻¹, this result is difficult to achieve with traditional carbon-based materials, and it indicates that this may be the key to CDI's future use in actual water desalination.

Another major factor limiting CDI to the theoretical level is the general adsorption of various types of ions. The simple electrochemical principle of CDI indicates that the highly charged ions with appropriate pore size are more easily

absorbed, and the low-charge materials whose pore size does not conform to the pore structure of the material are not easily absorbed. In other words, the selective adsorption capacity of CDI is weak. This is also a major technical vacancy in the field of CDI. Specific adsorption is of great significance for the recycling of various ions and the enrichment of heavy metals. Related studies have shown that from the perspective of pore structure, the unit adsorption capacity of traditional carbon-based materials can be effectively improved, while positive and negative charge modification can achieve the effect of ion selection to some extent. Once a breakthrough is made, it will definitely be a major advancement in the application of CDI to actual water treatment.

In conclusion, capacitive deionization is a promising technology. Through continuous research, CDI will play an important role in sewage disposal and seawater desalination in the future.

Conflicts of interest

There are no conflicts to declare.



Acknowledgements

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