RSC Advances

PAPER

Cite this: RSC Adv., 2019, 9, 23241

Received 15th May 2019 Accepted 15th July 2019 DOI: 10.1039/c9ra03659h

rsc.li/rsc-advances

Introduction

Currently, considering the continuous combustion of fossil fuels to meet the growing energy demand of the world, the everincreasing $CO₂$ emission has resulted in the quick rise of the atmospheric CO₂ concentration. Abnormal environmental problems have occurred in response to the massive $CO₂$ emission, such as global warming and its environmental effects, which include continuous rising sea levels, ocean acidification, increasing numbers of ocean storms and floods. $1-3$ However, $CO₂$ is known as an important carbon source with wide applications in industry and daily life. In order to mitigate $CO₂$ emission, CO₂ capture and sequestration technology is regard as one of the promising strategies for the reduction of $CO₂$ concentration and effective recycling of $CO₂$.⁴⁻⁶ According to the relevant statistical data, power plants are one of the largest

Biowaste-derived 3D honeycomb-like N and S dual-doped hierarchically porous carbons for highefficient CO₂ capture†

Weiwei Shi,^a Rongzhen Wang,^a Huili Liu,^a Binbin Chang, D^{*a} Baocheng Yang^{*a} and Zuling Zhang^b

Considering the characteristics of abundant narrow micropores of <1 nm, appropriate proportion of mesopores/macropores and suitable surface functionalization for a highly-efficient carbon-based CO₂ adsorbent, we proposed a facile and cost-effective strategy to prepare N and S dual-doped carbons with well-interconnected hierarchical pores. Benefiting from the unique structural features, the resultant optimal material showed a prominent CO_2 uptake of up to 7.76 and 5.19 mmol g^{-1} at 273 and 298 K under 1 bar, and importantly, a superb CO_2 uptake of 1.51 mmol g⁻¹ at 298 K and 0.15 bar was achieved, which was greatly significant for $CO₂$ capture from the post-combustion flue gases in practical application. A systematic study demonstrated that the synergetic effect of ultramicroporosity and surface functionalization determined the $CO₂$ capture properties of porous carbons, and the synergistic influence mechanism of nitrogen/sulfur dual-doping on CO₂ capture performance was also investigated in detail. Importantly, such as-prepared carbon-based $CO₂$ adsorbents also showed an outstanding recyclability and $CO₂/N₂$ selectivity. In view of cost-effective fabrication, the excellent adsorption capacity, high selectivity and simple regeneration, our developed strategy was valid and convenient to design a novel and highly-efficient carbonaceous adsorbent for large-scale CO₂ capture and separation from post-combustion flue gases. PAPER
 EXERCT AND AND SURFALL DEVALUE CONTINUES AND ASSESSED AND ALL CONTINUES AND ASSESSED AND ASSESSED AND A SURFACT AND

contributors for increasing atmospheric $CO₂$ concentration, which discharge about 60% of the global $CO₂$ emissions.⁷ Thus, it becomes an urgent need to capture and separate $CO₂$ from post-combustion flue gases. Although $CO₂$ adsorption from flue gases by liquid amine and ammonia solvents is an efficient strategy with satisfactory selectivity, this adsorption process suffers several tricky problems, which are high energy consumption, equipment corrosion, amine degradation, toxicity, and more severely the need for solvent regeneration.^{8,9}

In contrast, adsorption using porous solid materials as adsorbents for $CO₂$ capture has been recognized as the most promising alternative technology owing to its low energy consumption, low cost, relatively high efficiency and easy handling.^{10–12} The key of this technology is to develop new porous solid sorbents with superior properties for $CO₂$ capture. To this end, a number of porous solid materials, including zeolites, metal–organic frameworks, porous organic polymers, organic–inorganic hybrid adsorbents and porous carbon materials, have been extensively investigated.¹³⁻¹⁸ Among various types of adsorbent materials, porous carbons, as a special family of highly promising materials, have been extensively studied for high-efficiency $CO₂$ capture by virtue of their unique structural advantages, including high accessible surface area, low-cost preparation, facile regeneration, easy-todesign surface functionality and porosity, inertness to both

a Henan Key Laboratory of Nanocomposites and Applications, Institute of Nanostructured Functional Materials, Huanghe Science and Technology College, Zhengzhou, Henan 450006, China. E-mail: binbinchang@infm.hhstu.edu.cn; baochengyang@infm.hhstu.edu.cn

^bHenan Provincial Chemi-Industries Research Station Co., Ltd, Zhengzhou 450000, China

[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/c9ra03659h

bases and acids and moderate heat of adsorption. Nevertheless, most traditional activated carbons exhibit a relatively low $CO₂$ adsorption capacity of typically *ca.* 2–3 mmol g^{-1} under room temperature at 1 bar.^{19,20} In terms of $CO₂$ capture by porous carbons, it has been widely reported that $CO₂$ capture capacity at ambient pressure greatly depends on the proportion of narrow microporosity of smaller than 1 nm, and especially the ultramicroporosity of <0.7 nm, which plays a crucial role in determining $CO₂$ uptakes of porous carbons.²¹⁻²³ Thus, many efforts have been made to design microporous carbons with a large proportion of narrow micropores of <1 nm, especially to further tailor micropores to produce a more favorable ultramicroporous carbon for $CO₂$ capture. For example, Sevilla *et al.* prepared microporous biomass-based carbon materials via KOH chemical activation of hydrothermal carbons derived from mixtures of algae and glucose, which possessed a large number of narrow micropores (<1 nm) and exhibited a superior $CO₂$ adsorption capacity of 4.8 mmol g^{-1} at 298 K and 1 bar.²⁴ Our group synthesized cross-linked microporous carbon beads by air-assisted activated method using glucose-derived carbon microspheres as precursor, which developed a large proportion of ultramicropores with primary pore size of 0.5–0.9 nm and showed a satisfactory CO_2 uptake of 4.25 mmol g^{-1} at 298 K and 1 bar.²⁵ Another valid strategy to further improve $CO₂$ uptake of porous carbons is the incorporation of heteroatoms into carbon skeleton.²⁶⁻²⁸ Typically, nitrogen doping is the most attractive, which can provide more basic sites for enhanced interactions with acidic $CO₂$ molecule. Specifically, it has been testified that the pyrrolic and amine nitrogen functionalities have the strongest interactions with $CO₂$ molecules.²⁹ **Excharaces**

Some and access the controlled and moderate heating in the controlled substitute of the substitute o

In addition, it has been proposed that the enhancement of $CO₂$ uptake on porous carbons can be related to the improvement in polar interaction and hydrogen bonding interaction.^{30,31} Since hydrogen bonding or polar interactions of $CO₂$ within the carbon pore can more originate from other functional groups than nitrogen-based groups. Recently, $CO₂$ capture over sulfur-doped porous carbons has attracted much attention. For sulfur-doped porous carbons, the lone pair of electrons in a sulfur atom induces polarizability and interactions with oxygen.³² Xia et al. reported an improved $CO₂$ capture capacity of 2.4 mmol g^{-1} at 298 K and 1 bar on sulfur-doped porous carbon.³³ Seema et al. obtained a reduced-graphene oxide/polythiophene complex via chemical activation route, which exhibited an excellent CO₂ uptake of 4.5 mmol g^{-1} at 298 K and 1 bar.³⁴ Bandosz et al. reported the chemical interactions between sulfur doped carbons and $CO₂$.³⁵ These investigations manifested sulfur-containing functional groups in carbon adsorbents could facilitate $CO₂$ adsorption due to acid interactions of $CO₂$ with neutral sulfur, polar interactions of $CO₂$ with oxidized sulfur, and hydrogen bonding of $CO₂$ with sulfonic acids. However, there are relatively few studies about the synergistic effect of nitrogen and sulfur dual-doping on hierarchically porous carbons for $CO₂$ capture.^{36,37} Thus, it is significant for high-effective $CO₂$ capture to design a novel porous carbon adsorbent comprising multiscale pores including abundant narrow micropores of <1 nm (which favor high CO₂ uptake), appropriate proportion of mesopores/

macropores (which facilitate efficient $CO₂$ diffusion into and out of the adsorption sites) coupled with suitable surface nitrogen and sulfur functionalities (which further improve $CO₂$ uptake).

Considering the characteristics described above, herein, nitrogen and sulfur dual-doped porous carbons with wellinterconnected hierarchical pores were synthesized by a facile and cost-efficient strategy using bio-waste as carbon source and thiourea as nitriding and sulfurizing agent. Innovatively, the synergetic effect of ultramicropores (<0.7 nm) and nitrogen/ sulfur dual-doping on $CO₂$ capture was investigated in detail. Gratifyingly, the resultant optimal material exhibited a superior $CO₂$ capture capacity of up to 7.76 and 5.19 mmol $g⁻¹$ at 273 and 298 K under 1 bar, respectively. More importantly, such a $CO₂$ adsorbent also showed an outstanding recyclability and $CO₂/N₂$ selectivity adsorption property. Hence, such results suggested that we proposed a valid strategy to exploit novel porous carbon sorbents for the removal of $CO₂$ from post-combustion exhaust gases.

Experimental section

Material preparation

Preparation of hierarchically porous carbons. Hierarchically porous carbons were prepared by a facial chemical activation method using recycled waste paper as carbon source. In detail, 1 g of cleaned waste paper was dispersed in 30 mL of H_2O , and then the mixed solution was placed in a 100 mL of Teflon-sealed autoclave and maintained at 200 $^{\circ}$ C for 4 h. The puce products were obtained by filtration, washed repeatedly with distilled water and oven-dried at 80 \degree C for 6 h. Subsequently, 1 g of the dried puce products was impregnated in 20 mL of KOH/ZnCl2 solution and then stirred for 6 h. The mixed solution was dried at 110 °C for 8 h to obtain $KOH/ZnCl₂-impregnated carbon$ precursors, and then the precursors were activated and carbonized in a N_2 atmosphere at 800 °C for 1 h. After cooling down, the activated samples were repeated washed with HCl solution (1 M) and distilled water. Finally, these materials were dried under vacuum at 80 $^{\circ}$ C for 8 h to obtain the final products of hierarchically porous carbons, which were denoted as HPC_{K-x} and HPC_{Zn-x}, respectively ($x = 1, 2$ and 3, referring to the mass ratio of activating agent/puce product was 1 : 1, 2 : 1 and 3 : 1, respectively). For comparison, 1 g of the puce product was directly carbonized at 800 $^{\circ}$ C for 1 h without chemical activating agent to obtain hierarchically porous carbon, which was defined as HPC.

Synthesis of N and S dual-doped hierarchically porous carbons. Thiourea $(CN₂H₄S)$ was used as both N and S sources to synthesize the N and S dual-doped hierarchically porous carbons. In detail, 4.0 g of CN_2H_4S and 1 g of HPC_{K-1} (HPC_{Zn-1}) were mixed in 30 mL of deionized water and stirred for 6 h followed by drying at 90 \degree C overnight. And then the dried powders were heat-treated at 800 \degree C for 1 h with a heating rate of 5 °C min⁻¹ under a N₂ atmosphere. Following the wash with deionized water and dried at 90 \degree C, finally we obtained the N and S dual-doped hierarchically porous carbons, which were designed as N,S-HPC $_{K-1}$ and N,S-HPC $_{Zn-1}$, respectively.

X-ray diffraction (XRD) patterns were taken on a Bruker D8 diffractometer with Cu K α radiation ($\lambda = 0.15418$ nm). Nitrogen adsorption–desorption isotherms were acquired at 77 K using a Micromeritics ASAP 2020HD88 system. Before adsorption, the samples were out-gassed at 200 \degree C for 10 h. The specific surface area (S_{BET}) was evaluated according to the Brunauer–Emmett– Teller (BET) method at relative pressure of 0.05–0.25, and the pore size distribution was calculated according to the Density-Functional-Theory (DFT) method, and the micropores were analyzed using t-plot method, and the ultramicropores were analyzed by $CO₂$ adsorption at 273 K using DFT model. The morphology was imaged on a Quanta 250 FEG scanning electron microscope (SEM). Fourier transform infrared spectroscopy (FTIR) was performed on a Nicolet Avatar 370 spectrometer. The samples for the FTIR measurements were mixed together with KBr, following the standard method. X-ray photoelectron spectra (XPS) were obtained on a VG ESCALAB MK II spectrometer with an exciting source of Mg Ka at 1253.6 eV. Paper

Characterization (XR) patterns were taken on a Bruker Both constrained for the worst particle comparison in the common of the creative in the creative comparison in the comparison of the same pairs in the comparison

Gas adsorption measurements. The gas adsorption isotherms of $CO₂$ and $N₂$ were measured using a Micromeritics ASAP 2020HD88 system. Highly pure $CO₂$ (99.999%) and $N₂$ (99.999%) were employed for the measurements. The isotherms of $CO₂$ and N₂ at 273 and 298 K were acquired in an ice-water bath and a water bath, respectively. Prior to each gas uptake measurement, the samples were degassed at 200 $^{\circ}$ C for 10 h to remove adsorbed molecules from the pores.

Results and discussion

Materials characterization

Scheme 1 illustrates the schematic diagram for the fabrication of waste paper-derived N and S dual-doped hierarchically porous carbons via a convenient strategy combining chemical activation and post-modification methods. The waste paper towels were collected from a cardboard box in large quantity, and then thoroughly washed, cleaned and dried at 60 \degree C. These

dried waste paper towels were cut into pieces to be used as carbon source, and then were hydrothermal treatment at 200 $^{\circ}$ C for 4 h to obtain puce powders as precursor. These hydrothermally synthesized precursors were impregnated with activating agent solution for 4 h and then evaporated the solvent. Importantly, the resultant precursors not only contained abundant organic components, but also possessed an excellent adsorptive capacity to accommodate a large amount of activating agent, which favored generating well-developed porous structure, especially micropores. Followed, the dried activating-agentimpregnated precursors were carbonized and activated at 800 °C for 1 h in a nitrogen atmosphere, and washed by hydrochloric acid and water and dried to obtain hierarchically porous carbons. Subsequently, the resultant hierarchically porous carbons were further functionalized at 800 °C in a nitrogen atmosphere to prepare N and S dual-doped hierarchically porous carbons using thiourea as nitriding and sulfurizing agent.

To reveal the effect of carbonization and activation processes on the microcrystalline texture, the XRD patterns of all the resultant materials were shown in Fig. 1a. Two weak and broad peaks at approximately 22.1° and 43.6° can be observed in all samples, which are attributed to the (002) and (100) reflections of turbostratic carbon structure, suggesting an amorphous carbon texture with a low crystallinity. The diffraction intensity of the (002) and (100) peak in activated materials is lower than those in HPC under the same conditions of detection, resulting from the chemical activation to break down the hexagonal symmetry of the graphite lattice and lead to lattice defects in HPC_{K-1} and HPC_{Zn-1} samples.³⁸ In addition, the high intensity in the low angle region indicates the existence of abundant micropores in all samples.³⁹ The evolution of the chemical compositions of the waste paper precursor, activated samples and N and S dual-doped samples were characterized by FTIR (Fig. 1b). For hydrothermally synthesized precursor, it was similar to the chemical composition of the other plant fibers,⁴⁰ the main component of waste paper towel was lignocelluloses. The broad beak at about 3380 cm^{-1} was assigned to O–H

Scheme 1 Schematic illustrating the preparation of the waste paper-derived N and S dual-doped hierarchically porous carbons.

Fig. 1 The XRD patterns (a) and FTIR spectra (b) of the resultant materials.

stretching vibration. The band at 2900 cm^{-1} was attributed to the C–H stretching vibrations in methyl and methylene groups, and the bands at 1425 and 1370 cm^{-1} were ascribed to aromatic skeletal vibrations combined the C–H deformation vibrations, while the band at 1320 cm^{-1} was due to the CH₂ rocking vibration. The bands at 1660 and 1635 cm^{-1} were ascribed to the C $=$ O stretching vibration and C $=$ C stretching vibration, respectively. The band at 1160 cm^{-1} was related to C-O-C asymmetric valence vibration, and the band at 1113 cm^{-1} was due to the C–C stretching or asymmetric in-phase ring stretching. The bands at 1060 and 1032 cm^{-1} were assigned to C-O valence vibration and C–O ether vibration, respectively. While, after carbonization and chemical activation, it was obvious that the bands in the resultant materials became broader and weaker, this could be related to the strong absorption of the

carbon skeleton. Only the $C=C$ stretching vibration could be found in other activated samples, and the other characteristic absorption bands became weaker and even disappeared. Such results should be resulted from the carbonization and chemical activation, resulting in the elimination of surface functional groups.

Scanning electron microscopy was conducted to reveal the evaluation of microstructure and morphology from paper towel to activated carbons and N, S dual-doped carbons material. Paper towel showed a long fiber structure with a diameter of range from \sim 5 to \sim 20 µm (Fig. S1a†), and the hydrothermally synthesized precursors completely inherited the original fiber morphology of paper towel (Fig. S1b†). As displayed in Fig. 2a, the fibrous texture could be obviously observed in HPC sample, which indicated the original structure could be retained during the high-temperature carbonization treatment. After KOH activation, HPC_{K-1} and HPC_{K-2} presented a three-dimensional interconnected honeycomb-like microstructure with numerous irregular holes (Fig. 2b and c), which was a unique structure that could effectively prevent the small and thin blocks from agglomerating on a large scale. Such prominent honeycomb-like porous structure was attributed to the KOH chemical activation and the followed $CO/CO₂$ physical activation in situ induced from the reactions of KOH and C.⁴¹ With an increase in the addition amount of KOH, the density of pores presented a developing trend. Whereas, as the mass ratio of KOH/precursor increased to 3, a harsh activation reaction would occur in carbon skeleton, which brought the pore **Exchanges**
 Commons Access Article. Published on 26 July 2019. Downloaded the control and the stress Article is article. The common access Article is a supported to the common access Article is likely and the stress Art

Fig. 2 SEM images. (a) HPC; (b) HPC_{K-1}; (c) HPC_{K-2}; (d) HPC_{K-3}; (e) HPC_{Zn-1}; (f) HPC_{Zn-2}; (g) HPC_{Zn-3}; (h) N,S-HPC_{K-1}; (i) N,S-HPC_{Zn-1}.

Fig. 3 $N₂$ adsorption–desorption isotherms (a and b) and pore size distributions (c and d) of all the resultant materials.

widening and the degradation of 3D honeycomb-like structure. As a result, carbon sheets was too thin to support the honeycomb-like structure, resulting in the collapse and restacking of partial sections to form the block with thick wall (Fig. 2d). Different from the morphology of HPC_{K-x} samples, after ZnCl₂ activation, all the resultant HPC_{Zn-x} samples exhibited a porous structure with some macropores (Fig. 2e–g). Such results could be ascribed to the activation mechanism of $ZnCl₂$ chemical activation, and $ZnCl₂$ often serves as a dehydrating agent to chemically activate carbonaceous precursors. Similarly, with the increasing of $ZnCl₂$ dosage, the density of generated pores was decreased, which could be related to pore coalescence. After further nitrogen and sulfur dual-doped, the N,S-HPC $_{K-1}$ retained the original honeycomb-like porous structure of HPC_{K-1} sample (Fig. 2h), and N,S-HPC_{Zn-1} also kept the initial porous structure of HPC_{Zn-1} (Fig. 2i), which testified that the further functionalized process at high temperature could not destroy the intrinsic morphology.

The porous structure features of the resultant carbonaceous sorbents were analyzed by N_2 adsorption-desorption at 77 K. In Fig. S2,† it can be clearly found that the hydrothermally synthesized precursors possess an only 1.79 m^2 g⁻¹ of BET total surface area and a 0.006 cm³ g^{-1} of pore volume, which indicated that such carbonaceous precursors obtained by sole hydrothermal treatment exhibited extremely poor porosity. After directly carbonized precursors at 800 $^{\circ}$ C, the HPC material depicted a typical type I curve (Fig. 3a), meaning a microporous characteristic pore structure, and these micropores could

mainly originate from the decomposition of volatile matter and the elimination of surface O- and H-groups during the carbonization process. But, the relatively small N_2 adsorbed amount manifested the low total surface area and pore volume (Table 1). With the KOH chemical activation, the HPC_{K-x} still presented a type I isotherm of microporous structure, and the sharply increased and highly adsorbed quantity at a low relative pressure $(P/P_0 < 0.01)$ owed to the capillary filling of micropores, which testified the pore structure of HPC_{K-x} mainly consisted of micropores. Comparatively, an abrupt increase of the isotherm occurred at the higher relative pressure $(P/P_0 > 0.9)$ for HPC_{K-2} and HPC_{K-3} samples, which was ascribed to the capillary condensation, implying that a larger pore size was generated when more KOH was employed as activating agent. Fig. 3b showed the pore size distribution of HPC and HPC_{K-x} samples. HPC owned a micropore size distribution of 1.17/1.86 nm, and the proportion of 1.17 nm micropores was larger. Apparently, with the use of different dosage of KOH activating agent, the pore size gradually extended from bi-modal distribution to multi-modal distribution, and larger pore size and proportion stepwise enlarged with the enhancement of KOH dosage. The pore size distribution of HPC_{K-1} was 0.47-0.86 and 1.27 nm, and the overall pore sizes were still in the micropore range. The majority of the pore sizes centered at 0.47–0.86 nm, while the minority of the pore sizes was 1.27 nm. With the increase of KOH activating agent concentration, some new and larger pore size distributions of 1.61/2.02 nm and 2.10 nm were produced in HPC $_{K-2}$ and HPC $_{K-3}$ sample, respectively. More noteworthy,

Table 1 Textural properties of all the resultant materials

^a BET surface area. ^b Micropore surface area calculated using the V–t plot method. ^c S_{meso} = S_{BET} – S_{micro}. ^d The cumulative ultramicropore <0.7 nm surface area measured by CO₂ adsorption at 273 K using DFT model. ^e The total pore volume calculated by single point adsorption at $P/P_0 = 0.9945$. f The micropore volume calculated using the V–t plot method. g The cumulative ultramicropore <0.7 nm volume measured by CO₂ adsorption at 273 K using DFT model.

the majority of pore sizes distributed in 1.61 and 2.10 nm, and the minority of pore sizes was in micropores of 0.47–1.27 nm. When used $ZnCl₂$ as activating agent, the HPC_{Zn-1} sample exhibited a similar isotherm with HPC_{K-x} , meaning a characteristic micropore structure of HPC_{Zn-1} . Whereas, HPC_{Zn-2} and HPC_{Zn-3} presented a transitional isotherm from type I to type IV (Fig. 3c), and even a marked H3 type hysteresis loop in the relative pressure region between 0.4 and 0.7 could be clearly observed in HPC_{Zn-3} . Such results manifested the coexistence of micropores and slit-shaped mesopores. The pore size distributions of HPC_{Zn-x} samples were shown in Fig. 3d. The pore size of HPC_{Zn-1} centered at 0.68 and 1.27 nm, and overall pore size were in micropore, and the pore proportion was almost isometric. With the increase of $ZnCl₂$ dosage, some new mesopores of 2.34 and 3.07 nm were generated in HPC_{Zn-2} and HPC_{Zn-3} samples, respectively, and these mesopores were dominant in porosity. By comparing the pore size distributions of HPC_{K-x} and HPC_{Zn-x} samples, we could speculate such conclusions: (i) the hierarchical pore size distribution significantly depend on the activation of activating agents; (ii) KOH activating agent exhibits a more effective impact on tailoring multi-modal pore size distribution, especially in micropores; (iii) $ZnCl₂$ activating agent plays a high-effective role in developing mesopores. In addition, the BET surface area and other textural properties of all the resultant materials are summarized in Table 1. It can be clearly found that the activating agents have a vital influence on developing porosity of materials. With the increase of activating agent dosage, both surface area and pore volume were greatly improved owing to the further etch of activating agent on carbonaceous skeleton. However, as the activating agent dosage further increase to 3, because of the excessive activation, numerous micropores coalesced to form a mesopore, which resulted in the continuous enhancement of mesoporous surface area and mesopore volume and decrease of total surface area and micropore volume. Therefore, varying the dosages of the activating agent is an effective strategy to engineer the proportion of mesopore/micropore in porosity. Moreover, after N and S dual-doping, the specific surface area and pore volume changed slightly, which testified that surface functionalization process could not destroy the original porosity of materials.

CO₂ capture property

The $CO₂$ adsorption performance of the resultant HPC, HPC_{K-x} and HPC_{Zn-x} samples were investigated at two representative temperatures of 273 and 298 K under the ambient pressure, and the $CO₂$ adsorption isotherms are depicted in Fig. 4a and b. At the atmospheric pressure (1 bar), all the samples exhibit a higher $CO₂$ uptake at 273 K than that of at 298 K, which should be related to the exothermic feature of $CO₂$ adsorption.⁴² HPC sample exhibits a CO₂ uptake of 3.12 mmol g^{-1} at 273 K and 2.35 mmol g^{-1} at 298 K under 1 bar. After activated by KOH and $ZnCl₂$, all HPC_{K-x} and HPC_{Zn-x} (except HPC_{Zn-3}) samples present the improved CO_2 uptakes in the ranges of 3.82–7.26 mmol g^{-1} and 2.69-4.68 mmol g^{-1} at 273 K and 298 K under 1 bar, respectively. Obviously, HPC_{K-1} exhibits the highest $CO₂$ capture capacity of 7.26 and 4.68 mmol g^{-1} at 273 and 298 K in KOHactivated system, respectively. However, the further increase in activating agent dosage of KOH brings a negative effect on $CO₂$ adsorption capacity, and the declined $CO₂$ uptake values of 6.84 (6.25) and 4.25 (3.76) mmol g^{-1} are obtained over HPC_{K-2} (HPC_{K-3}) at 273 and 298 K, respectively. The similar variation tendency of $CO₂$ uptake can be found over HPC_{Zn-x} samples in $ZnCl₂$ -activated system. Noticeably, HPC_{K-x} samples exhibit the better CO_2 capture performance than HPC_{Zn-x} samples, which should be benefited from the more prominent porosity of HPC_{K-x} , especially the higher microporosity and ultramicroporosity. In addition, although HPC sample presents a low $CO₂$ uptake at the atmospheric pressure, and it shows a more superior $CO₂$ capture property than those of the other resultant samples below 0.15 bar at 273 K but 0.3 bar at 298 K (Fig. S3†), which suggest that $CO₂$ capture capacity is also related to the adsorption conditions. Under high adsorption temperature and low pressure, small micropores contribute more to $CO₂$ capture. But, with declining adsorption temperature and increasing adsorption pressure, $CO₂$ capture is more dependent on large micropores and even mesopores. Such result is ascribed to the developed multiscale micropore of <1 nm size distribution, especially its more prominently hierarchical ultramicropore size distribution (Fig. $S4, \dagger$ CO₂ micropore size of <1 nm distribution), and such behavior is closely related to the pore filling mechanism of $CO₂$ adsorption.²¹

Fig. 4 CO₂ adsorption isotherms of all the prepared materials. (a) at 273 K; (b) 298 K; (c) the comparison of HPC_{K-1}, HPC_{Zn-1}, N,S-HPC_{K-1} and N,S-HPC_{Zn-1} at 273 K; (d) the comparison of HPC_{K-1}, HPC_{Zn-1}, N,S-HPC_{K-1} and N,S-HPC_{Zn-1} at 298 K.

Furthermore, though HPC_{K-2} possesses the higher total surface area, micropore surface area and micropore volume than those of HPC_{K-1}, HPC_{K-2} sample presents a lower CO₂ uptake of 6.84 and 4.25 mmol g^{-1} at 273 and 298 K under 1 bar than that of HPC_{K-1} , and the similar result can be found over HPC_{Zn-2} and HPC_{Zn-1} samples. Such behavior should be attributed to their declined proportion of ultramicropores, giving rise to the decreased ultramicropore surface area and ultramicropore volume. This result testifies that the efficient $CO₂$ capture of porous carbons does not only depend on the micropore surface area/pore volume, but also more depend on ultramicropore surface area/volume, as ultramicropores are more beneficial to the $CO₂$ molecule filling.^{21,43} Thus, the inferior $CO₂$ capture property of HPC_{Zn-2} and HPC_{Zn-3} should also be due to the overactivated in micropore structure, which results in forming a larger proportion of mesopores but a poorer ultramicroporosity. After nitrogen and sulfur dual-doped, the obtained N,S-HPC $_{K-1}$ and N,S-HPC $_{Zn-1}$ samples both exhibit an enhanced $CO₂$ capture performance at 273 K and 298 K (Fig. 4c and d), manifesting that the nitrogen and sulfur functionalization of carbon skeleton brings an effective improvement for the $CO₂$ capture, which should be benefit from the increasing number of $CO₂$ adsorption sites on the pore surface. Especially, N,S-HPC $_{K-1}$ sample displays an excellent CO₂ uptake of 7.76 and 5.19 mmol g^{-1} at 273 and 298 K under 1 bar, respectively, which is much higher than other carbon-based sorbents and even some of the reported nitrogen-doped porous carbons and hierarchical porous carbons (Table S1†). Of particular importance is that the hierarchically porous carbons prepared in this work using cost-free biomass waste as carbon precursor, exhibits a comparable $CO₂$ uptake to those of N-doped carbonaceous adsorbents that are considered as good $CO₂$ adsorbents.

As is well known, the post-combustion flue gas streams produced in industrial processes $(e.g.,$ fossil fuel-fired power stations) contain a large proportion of $CO₂$. The partial pressure of $CO₂$ in the flue gas streams is typically around 0.15 bar, and it is still a challenge to capture $CO₂$ from post-combustion flue gases. In order to examine the possibility of the resultant samples to act as the $CO₂$ adsorbents for the flue gas streams, we also list the adsorbed $CO₂$ amounts at 298 K under 0.15 bar in Fig. 5. The $CO₂$ uptakes at 0.15 bar vary with the amount of the activating agent. The variation trend at 0.15 bar is slightly different from that at 1 bar. It can draw a conclusion that the $CO₂$ capture amount more rely on the proportion of ultramicropores and pore size distribution of these ultramicropores, and thus the high $CO₂$ uptake amount of 1.29 at 298 K at 0.15 bar for HPC $_{K-1}$ arises from its prominent ultramicroporosity. More importantly, after modification with nitrogen and sulfur, N,S-HPC $_{K-1}$ sample exhibits a more superior CO_2 capture

Fig. 5 The CO₂ uptakes of all materials at 298 K under 0.15 bar and the correlation with ultramicropore volume.

capacity of up to 1.51 mmol g^{-1} at 0.15 bar at 298 K, greatly surpassing the uptakes of previously reported carbon-based $CO₂$ adsorbents (Table S1†). Consequently, the currently designed nitrogen and sulfur dual-doped hierarchically porous carbons have excellent potential for capturing $CO₂$ from the post-combustion flue gases.

Effect of hierarchical porosity on $CO₂$ capture capacity

To better understand the $CO₂$ uptake behaviors of the resultant samples, we further analyzed the effect of the hierarchical porosity and structure on the $CO₂$ adsorption capacity. A comparison of the porosity characteristics and the $CO₂$ uptake amounts shows that the $CO₂$ uptake amount of the resultant samples is not directly determined by the BET total surface area and total pore volume (shown in Fig. 6a and b).

But, the $CO₂$ capture capacities of the resultant materials are strongly correlated with their microporosity of pore size <1 nm, especially the ultramicropore volume between 0.3 and 0.7 nm. Especially, it can be clearly found that the $CO₂$ uptakes at 0.15 bar greatly depend on the ultramicropore volume, and the $CO₂$ capture capacity enhances with the increase of ultramicropore volume (Fig. 6c). Such result suggests that the micro and narrow mesopores were mainly responsible for the high $CO₂$ sorption performance of the materials at low pressure, which is in agreement with previous findings.^{3,44} By analyzed the data in Fig. 6d, it can be concluded that the best $CO₂$ capture capacity of HPC_{K-1} should be owed to the largest ultramicropore volume. In addition, HPC_{Zn-x} samples exhibit the lower $CO₂$ uptakes

compared to HPC_{K-x} samples although they possess the comparative BET surface area, which should be correlated with the lower ultramicropore volumes of HPC_{Zn-x} . Thus, to achieve an outstanding $CO₂$ capture capability, it is more important to tailor the ultramicropore size than to have a high surface area for porous carbon-based $CO₂$ adsorbents. But, the $CO₂$ uptakes at 1 bar are also still affected by the BET total surface area, as revealed by Fig. 6a, which should be related to the important impact large micropores and even mesopores on $CO₂$ capture capacity at high pressure region. Therefore, the large proportion of microporosity and suitable ultramicroporosity combined with the largest surface area might be an essential factor for the outstanding $CO₂$ uptake at 273 and 298 K under ambient pressure. RSC Advances Compariso of up to the most of the comparison of the published on 26 Kullet are the published on 26 ML and 2018. The comparison of the common access Article is licensed on 26 ML and the common access Article

Effect of surface functionalization on $CO₂$ capture capacity

In general, surface chemical modification, especially nitrogen or sulfur doping in carbon skeleton, is another non-negligible factor for determining the $CO₂$ capture capacity of porous carbon-based sorbents. In this work, the resultant $CO₂$ adsorbents should consist mainly of nitrogen, sulfur and oxygen atoms on the surface of carbon, and therefore we investigated the effects of N, S and O on the $CO₂$ capture capacity. Fig. S5[†] depicts the elemental mapping of C, N, S and O collected by scanning N,S-HPC $_{K-1}$ porous microstructure. The homogeneous distribution of N, S and O element in N,S-HPC $_{K-1}$ carbon framework indicates that the successful modification of N and S on the surface of carbon skeleton in N,S-HPC $_{K-1}$ sample. To

Fig. 6 (a) Dependence of the CO_2 uptake amount at 273 K and 1 bar on the total and microporous surface areas. (b) Dependence of the CO_2 uptake amount at 273 K and 1 bar on the total pore and microporous volumes. (c) The correlation of CO₂ uptake at 273 and 298 K under 0.15 bar with ultramicropore volume. (d) The cumulative ultramicropore volume measured by CO_2 adsorption at 273 K using DFT model.

better understand the contribution of the chemical interactions towards the high $CO₂$ sorption, X-ray photoelectron spectroscopy (XPS) analysis was conducted on the optimally performing materials to study the nature of the surface groups. Fig. 7a presents the full XPS spectrum of the N,S-HPC $_{K-1}$ sample with four main peaks at binding energies of 163.6, 284.1, 398.2 and 532.5 eV, which corresponds to S, C, N and O element, respectively. The C, N, O and S atomic percentages of $N, S-HPC_{K-1}$ determined by XPS are 89.95%, 4.32%, 4.57% and 1.16%, respectively.

N 1s core spectrum contributes to a further understanding of the types and local environment of the nitrogen atoms modified on the surface of carbon matrix. The deconvolution of N 1s spectrum reveals the bonding of N with C (Fig. 7b), and the N 1s signal can be resolved into four individual peaks centered at binding energies of 398.5, 400.1, 400.9, 403.6 eV, which are related to four different types of nitrogen functional groups, corresponding to pyridinic-N ($=N-H$), amine ($-NH₂$), pyrrolic/ pyridone-N $(-N-H)$ and quaternary-N $(=N^{+}-H)$, respectively. Apparently, the pyridinic-N and amine nitrogen groups are the most stable and abundant, which has been identified the pyridinic-N or amine nitrogen groups to have stronger interactions with CO_2 molecules,^{29,45} resulting in bringing the more contribution for $CO₂$ capture. In addition, it was also revealed that the introduction of nitrogen into carbon matrix facilitated hydrogen-bonding interactions between the carbon surface and $CO₂$ molecules,³⁰ favoring the improvement of $CO₂$ uptake.

The deconvolution of S 2p core level peak of N,S-HPC_{K-1} sample displays the peaks located at 163.8 and 164.9 eV corresponding to neutral S, and the peak centered at 168.5 eV assigns to oxidized S (Fig. 7c). Sulfur-containing functional groups in carbon skeleton could facilitate $CO₂$ adsorption owing to the acid interactions of $CO₂$ with neutral sulfur (163.8 and 164.9 eV) and polar interactions of $CO₂$ with oxidized sulfur (168.5 eV).³⁷ As a result, N,S-HPC $_{K-1}$ exhibits an enhanced CO₂ capture capacity than that of HPC_{K-1} . Additionally, to examine the synergistic effect of N and S dual-doping on $CO₂$ capture capacity, a conjecture is made to ascertain whether there is an optimum N/S ratio, which furthest promoting the $CO₂$ capture capacity. As presented in Fig. S6,† unexpectedly, there is no obvious correlation between N/S ratio and $CO₂$ uptake, which could be related with the heterogeneous distribution of nitrogen and sulfur in the surface of carbon matrix and the limited sites for the doping of nitrogen and sulfur groups.

The surface oxygen functionalities on the carbon matrix was also been revealed to contribute to the enhancement of $CO₂$ adsorption.⁴⁶ The O 1s spectrum shown in Fig. 7d exposes the bonding of N with C, which can be fitted into three deconvoluted peaks located at ca. 531.4, 532.6 and 533.8 eV, respectively. The peak at about 531.4 eV ascribes to the contribution of oxygen in carboxyl groups and the peak at ca. 532.6 eV assigns to the $-C=O$ band, and the one centered at ca. 533.8 eV should be related to the band of –C–O–C. The oxygen content detected from XPS is correlated with the $CO₂$ capture capacity of the as-obtained materials (Fig. S7†), and there is no obvious trend between the oxygen content and the $CO₂$ uptake. Moreover, a guess is made to ascertain whether there is an optimum N/O ratio, which favors improving the $CO₂$ uptake. As displayed in Fig. S8, \dagger no correlation is observed with N/O ratio, which could be ascribed to the heterogeneous distribution of oxygen in the surface of carbon framework. Similarly, such no correlation relationship between $CO₂$ capture capacity and the oxygen content was found in porous carbon sorbents.⁴⁷ Paper

Very method on 2019. Download the chemical interactions and interactions of O₂ with reading the total control

compute the big Co₂ supplies Article is a compute the published on 26 pM. This article is and

one p

Fig. 7 The XPS spectrum of N,S-HPC_{K-1}. (a) The survey; (b) N 1s; (c) S 2p; (d) O 1s.

Fig. 8 Isosteric heat of $CO₂$ adsorption on all the as-obtained samples calculated from the adsorption isotherms at 273 and 298 K. (a) HPC, HPC_{K-x} and N,S-HPC_{K-1}; (b) HPC_{Zn-x} and N,S-HPC_{Zn-1}.

$CO₂$ isosteric heat of adsorption (Q_{st})

In order to reveal the combination interaction strength between the $CO₂$ molecule and adsorbents, and evaluate the energetic heterogeneity of surfaces of adsorbents, the isosteric heat of

adsorption (Q_{st}) was calculated from the $CO₂$ adsorption isotherms measured at 273 and 298 K based on Clausius–Clapeyron equation. As depicted in Fig. 8, the Q_{st} values of the resultant HPC_{K-x} and HPC_{Zn-x} adsorbents at low surface coverage are in the range of 30-35 and $18-27$ kJ mol⁻¹, respectively. It can be clearly found that the Q_{st} gradually declines at the low $CO₂$ uptake, and then reaches a near plateau as the continuous occupation of adsorption active sites with the increasing $CO₂$ uptake, manifesting the heterogeneity of interaction between $CO₂$ molecules and the surface of adsorbents, which could be related to the heterogeneity of the surface chemistry and pore sizes. Obviously, the Q_{st} values of HPC_{K-x} are much higher than those of HPC_{Zn-x} samples, which should be benefit from their more prominent microporosity, especially the developed micropores of <1 nm. Moreover, it is worth of notice that the Q_{st} values of HPC_{K-x} samples gradually decline with the increase of activating agent dosage, suggesting the weaker and weaker interactions between adsorbent surfaces and $CO₂$ molecule. Such result should be related to the poorer and poorer microporosity, especially the decreased proportion of ultramicropores with the rise of activating agent dosage, resulting in the weaker interaction. The similar trend can be observed in HPC_{Zn-x} samples, and such result further confirms the crucial role of micropore proportion and size in capturing CO₂. Even though HPC_{K-3} has a higher Q_{st} value at low CO₂ coverage, its Q_{st} value continuously declines with the enhanced $CO₂$ uptake, indicative of the nonuniform and unstable $CO₂$ adsorption on HPC_{K-3} adsorbent surface, resulting in the poor regeneration and reversibility, which seriously restricts it to be an efficient $CO₂$ adsorbent. In addition, it is clearly observed that the samples functionalized by nitrogen and sulfur groups **EXADEMEDIES ARTIFE ACCESS ARTIFE ACCESS ARTIFE ARTIF**

Fig. 9 Adsorption selectivities. (a) CO₂ and N₂ adsorption isotherms of HPC_{K-1} at 273 and 298 K. (b) CO₂ and N₂ adsorption isotherms of N,S-HPC_{K-1} at 273 and 298 K. (c) IAST-calculated selectivities of CO₂/N₂ on HPC_{K-1} and N,S-HPC_{K-1} at 273 K. (d) IAST-calculated selectivities of CO₂/ N_2 on HPC $_{K-1}$ and N,S-HPC $_{K-1}$ at 298 K.

show an enhanced Q_{st} values, meaning an improved interaction, which should be ascribed to the more adsorption sites contributed by nitrogen and sulfur doping. Importantly, the initial $Q_{\rm st}$ of N,S-HPC_{K-1} is up to 39 kJ mol⁻¹, is higher than those of porous carbons and is comparable to that of nitrogen doped carbons.⁴⁸⁻⁵² For sulfur doped carbons, since the radius of sulfur is larger than that of carbon, sulfur atoms tend to protrude out of the carbon layer.⁵³ The disruption of the carbon structure will induce strain and defects, and facilitate charge localization creating favorable conditions for the adsorption of CO2. Furthermore, due to the large and polarizable d-orbitals of sulfur, the lone pair of electrons in doped sulfur atom can interact easily with the oxygen.⁵⁴ In the current work, the doped sulfur interacts with the oxygen of $CO₂$ molecule and thus strengthens the adsorption of $CO₂$, as reflected by the high initial Q_{st} value. The high Q_{st} at low CO_2 loading is helpful for $CO₂$ capture at a low pressure, and the moderate Q_{st} is beneficial for easy desorption to regenerate the sorbents. Paper

Move an enhanced Q_k values, meaning an improved interact

com, which shock the median on the more assembles to the under published in the signal matrix and its are the endanged on the signal matrix and its articl

$CO₂/N₂$ selectivity and cycling stability

Highly efficient $CO₂$ adsorbents need to meet the following characteristics: (i) large $CO₂$ uptake; (ii) fast adsorption kinetics; (iii) moderate heat of adsorption; (iv) good selectivity against other gas molecules; (v) easy regeneration. To evaluate the $CO₂$ separation performance of the selected sample of HPC_{K-1} and N,S-HPC $_{K-1}$ samples, their CO₂ and N₂ isotherms at 273 and 298 K are presented in Fig. 9a and b. Obviously, the adsorption capacity of N_2 is much lower than that of CO_2 at the same condition for both HPC_{K-1} and N,S-HPC_{K-1} samples. At 298 K and 1 bar, N,S-HPC $_{K-1}$ sample has a CO₂ uptake of 5.19 mmol $\rm g^{-1}$ and a N₂ adsorption capacity of 0.67 mmol $\rm g^{-1}.$ The equilibrium CO_2/N_2 adsorption ratio of 8 is higher than the value of 7 for HPC_{K-1} sample, indicative of the better selectivity of N,S- HPC_{K-1} for $CO₂$ from N₂.

The ideal adsorbed solution theory (IAST) model was employed to evaluate the selectivity for $CO₂$ adsorption from simulated post-combustion flue gas, which has been widely used to predict the selectivity of adsorbents for any two gases in a binary gas mixture by using the isotherms of pure gas. The calculated CO_2/N_2 selectivity of HPC_{K-1} and N,S-HPC_{K-1} samples with IAST model are displayed in Fig. 9c and d. The $CO₂/N₂$ ratio is 15/85 in the calculation, representing the typical composition of flue gas. Apparently, the selectivity of HPC_{K-1} and N,S-HPC_{K-1} samples significantly decrease in the low pressure region, finally reaching a plateau despite the increase in pressure. The highest selectivity of N,S-HPC $_{K-1}$ in the low pressure are 72 and 73 at 273 and 298 K, respectively, and the corresponding values are 70 and 69 for HPC_{K-1} sample. Evidently, the selectivity of N,S-HPC_{K-1} is superior than that of HPC_{K-1} under the same conditions, manifesting that the introduction of nitrogen and sulfur groups into the HPC_{K-1} framework brings an improvement in CO_2 capture and CO_2/N_2 selectivity. Such results should be attributed to the strengthened affinity of $CO₂$ molecule and N,S-HPC $_{K-1}$ skeleton owing to the more adsorption sites derived from the doping of nitrogen and sulfur. The same result can be found in N,S-HPC $_{Zn-1}$ and HPC $_{Zn-1}$ samples (Fig. S9†),

Fig. 10 Recyclability test on CO_2 capture for N,S-HPC_{K-1}. (a) at 273 K; (b) at 298 K.

which further confirms the improved effect of nitrogen/sulfur doping in carbon matrix for enhancing $CO₂/N₂$ selectivity. Importantly, these values are even greatly higher than most of reported porous carbons and nitrogen-rich carbon materials.⁵⁵–⁵⁸ Moreover, it is noticeable that the selectivity has no decrease with the increase of adsorption temperature.

Besides prominent separation performance, the recyclability of an adsorbent is a critical property determining the potential of practical utilization. After the adsorbent was saturated with $CO₂$ up to 1 bar, the adsorbent was recycled. Then, the recycled adsorbent was degassed at the room temperature for 10 min to apply for the next adsorption. The regeneration test of N,S- HPC_{K-1} sample was conducted for ten consecutive cycles at 273 and 298 K. As presented in Fig. 10a and b, the $CO₂$ uptakes are almost similar without noticeable loss, indicative of the outstanding recyclability of N,S-HPC $_{K-1}$ with relatively low energy requirement for regeneration. Moreover, such recyclability test suggests $CO₂$ can be completely desorbed from the adsorbent by only changing the pressure. It offers a chance for such adsorbents to be applied in the pressure swing adsorption technology.

Conclusions

In this work, a series of hierarchically porous carbons with wellinterconnected porosity were successfully prepared by one-step carbonization and activation route using no-cost biomass waste as carbon source. Followed by a facile nitrogen and sulfur binary doping using thiourea as nitriding and sulfurizing agent, N,S-co-doped porous carbons were obtained with more surface active sites for efficient $CO₂$ capture. By further investigating activation mechanism of KOH/ZnCl₂ activating agents, it can be drawn that KOH activating agent mainly produce micropores with different pore sizes and $ZnCl₂$ activating agent tends to develop mesopores. As expected, the resultant N,S- HPC_{K-1} material exhibits an excellent CO_2 capture capacity of up to 7.76 and 5.19 mmol g^{-1} at 273 and 298 K under 1 bar, respectively, which are much higher than those of reported carbon-based $CO₂$ adsorbents. By analyzing the $CO₂$ capture performance in detail, it can be found that $CO₂$ capture at atmosphere pressure is a combined action of the ultramicropore structure and the surface chemistry, in which the ultramicropore plays a determined role. Further investigations shows that the synergetic effect of nitrogen and sulfur dual-

doping has an improvement in $CO₂$ capture, but there is no an obvious connection between nitrogen/sulfur doping content and $CO₂$ uptake. More importantly, N,S-HPC_{K-1} material also exhibits a moderate isosteric heat of adsorption, a satisfactory recyclability and high selectivity for $CO₂$ capture from the gas mixture of $CO₂$ and N₂. Considering the attractive features for $CO₂$ capture, the low-cost preparation as well as the aspect of environmental friendliness, our prepared N/S dual-doped hierarchically porous carbons are believed to be a promising candidate for capturing $CO₂$ from flue gas. **BSC Articles.** Complement article is the street is the street in the street is are the street in the street i

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the National Natural Science Foundation of China (51702114), the National Natural Science Foundation of China (51872110), Natural Science Foundation of Henan Province (172102210381), Natural Science Foundation of Education Department of Henan Province (18A150011 and 19A150032).

References

- 1 S. Mane, Z. Y. Gao, Y. X. Li, D. M. Xue, X. Q. Liu and L. B. Sun, J. Mater. Chem. A, 2017, 5, 23310–23318.
- 2 L. M. Yue, L. L. Rao, L. L. Wang, L. Y. An, C. Y. Hou, C. D. Ma, H. DaCosta and X. Hu, Energy Fuels, 2018, 32, 6955–6963.
- 3 B. B. Chang, W. W. Shi, H. Yin, S. R. Zhang and B. C. Yang, Chem. Eng. J., 2019, 358, 1507–1518.
- 4 M. Idrees, V. Rangari and S. Jeelani, *J. CO₂ Util.*, 2018, 26, 380–387.
- 5 W. J. Tian, H. Y. Zhang, H. Q. Sun, A. Suvorova, M. Saunders, M. Tade and S. B. Wang, Adv. Funct. Mater., 2016, 26, 8651– 8661.
- 6 Y. Lin, C. Kong, Q. Zhang and L. Chen, Adv. Energy Mater., 2017, 7, 1601296.
- 7 Y. W. Chen, Z. W. Qiao, J. L. Huang, H. X. Wu, J. Xiao, Q. B. Xia, H. X. Xi, J. Hu, J. Zhou and Z. Li, ACS Appl. Mater. Interfaces, 2018, 10, 38638–38647.
- 8 M. Sevilla, P. Valle-Vigón and A. B. Fuertes, Adv. Funct. Mater., 2011, 21, 2781–2787.
- 9 Y. H. Abdelmoaty, T. D. Tessema, N. Norouzi, O. M. El-Kadri, J. B. M. G. Turner and H. M. El-Kaderi, ACS Appl. Mater. Interfaces, 2017, 9, 35802–35810.
- 10 A. M. Fracaroli, H. Furukawa, M. Suzuki, M. Dodd, S. Okajima, F. Gandara, J. A. Reimer and O. M. Yaghi, J. Am. Chem. Soc., 2014, 136, 8863–8866.
- 11 G. K. Parshetti, S. Chowdhury and R. Balasubramanian, RSC Adv., 2014, 4, 44634–44643.
- 12 J. Gong, M. Antonietti and J. Y. Yuan, Angew. Chem., Int. Ed., 2017, 56, 7557–7563.
- 13 H. Li, M. M. Sadiq, K. Suzuki, R. Ricco, C. Doblin, A. J. Hill, S. Lim, P. Falcaro and M. R. Hill, Adv. Mater., 2016, 28, 1839– 1844.
- 14 Z. Xiang, R. Mercado, J. M. Huck, H. Wang, Z. Guo, W. Wang, D. Cao, M. Haranczyk and B. Smit, J. Am. Chem. Soc., 2015, 137, 13301–13307.
- 15 J. Chen, J. Yang, G. S. Hu, X. Hu, Z. M. Li, S. Wshen, M. Radosz and M. H. Fan, ACS Sustainable Chem. Eng., 2016, 4, 1439–1445.
- 16 L. H. Zhang, W. C. Li, L. Tang, Q. G. Wang, Q. T. Hu, Y. Zhang and A. H. Lu, J. Mater. Chem. A, 2018, 6, 24285– 24290.
- 17 L. W. Wang, L. L. Rao, B. B. Xia, L. L. Wang, L. M. Yue, Y. Q. Liang, H. Dacosta and X. Hu, Carbon, 2018, 130, 31–40.
- 18 J. Wang, P. Zhang, L. Liu, Y. Zhang, J. Yang, Z. Zeng and S. Deng, Chem. Eng. J., 2018, 348, 57–66.
- 19 D. Qian, C. Lei, E. M. Wang, W. C. Li and A. H. Lu, ChemSusChem, 2014, 7, 291–298.
- 20 D. Qian, C. Lei, G. P. Hao, W. C. Li and A. H. Lu, ACS Appl. Mater. Interfaces, 2012, 4, 6125–6132.
- 21 Z. Zhang, J. Zhou, W. Xing, Q. Xue, Z. Yan, S. Zhuo and S. Z. Qiao, Phys. Chem. Chem. Phys., 2013, 15, 2523–2529.
- 22 V. Presser, J. Mcdonough, S. H. Yeon and Y. Gogotsi, Energy Environ. Sci., 2011, 4, 3059–3066.
- 23 N. Wickramaratne and M. Jaroniec, J. Mater. Chem. A, 2013, 1, 112–116.
- 24 M. Sevilla, C. Falco, M. Titirici and A. Fuertes, RSC Adv., 2012, 2, 12792–12797.
- 25 B. B. Chang, L. Sun, W. W. Shi, S. R. Zhang and B. C. Yang, ACS Omega, 2018, 3, 5563–5573.
- 26 F. Q. Yang, J. Wang, L. Liu, P. X. Zhang, W. K. Yu, Q. Deng, Z. L. Zeng and S. G. Deng, ACS Sustainable Chem. Eng., 2018, 6, 15550–15559.
- 27 P. Zhang, Y. Zhong, J. Ding, J. Wang, M. Xu, Q. Deng, Z. Zeng and S. Deng, Chem. Eng. J., 2019, 355, 963–973.
- 28 Y. H. Abdelmoaty, T. D. Tessema, N. Norouzi, O. M. EI-Kadri, J. B. Mcgee Turner and H. M. EI-Kaderi, ACS Appl. Mater. Interfaces, 2017, 9, 35802–35810.
- 29 A. Sanchez-Sanchez, F. Suarez-Garcia, A. Martinez-Alonso and J. M. Tascon, ACS Appl. Mater. Interfaces, 2014, 6, 21237–21247.
- 30 W. Xing, C. Liu, Z. Zhou, L. Zhang, J. Zhou, S. Zhuo, Z. Yan, H. Gao, G. Wang and S. Z. Qiao, Energy Environ. Sci., 2012, 5, 7323–7327.
- 31 J. Wang, Y. Lin, Q. Yue, K. Tao, C. Kong and L. Chen, RSC Adv., 2016, 6, 53017–53024.
- 32 J. P. Paraknowitsch and A. Thomas, Energy Environ. Sci., 2013, 6, 2839–2855.
- 33 Y. Xia, Y. Zhu and Y. Tang, Carbon, 2012, 50, 5543–5553.
- 34 H. Seema, K. C. Kemp, N. H. Le, S. W. Park, V. Chandra, J. W. Lee, et al., Carbon, 2014, 66, 320–326.
- 35 M. Seredych, J. Jagiello and T. J. Bandosz, Carbon, 2014, 74, 207–217.
- 36 T. J. Bandosz, M. Seredych, E. Rodríguez-Castellon, Y. Cheng, L. L. Daemen and A. J. Ramírez-Cuesta, Carbon, 2016, 96, 856–863.
- 37 Y. H. Sun, K. X. Li, J. H. Wang, N. Tang, D. Zhang, T. T. Guan and Z. Jin, J. Colloid Interface Sci., 2018, 526, 174–183.
- 38 G. Y. Xu, J. P. Han, B. Ding, P. Nie, J. Pan, H. Dou, H. S. Li and X. G. Zhang, Green Chem., 2015, 17, 1668–1674.
- 39 L. J. Xie, G. H. Sun, F. Y. Su, X. Q. Guo, Q. Q. Kong, X. M. Li, et al., J. Mater. Chem. A, 2016, 4, 1637–1646.
- 40 L. C. Zhang, X. Sun, Z. Hu, C. C. Yuan and C. H. Chen, J. Power Sources, 2012, 204, 149–154.
- 41 Y. X. Liu, Z. C. Xiao, Y. C. Liu and L. Z. Fan, J. Mater. Chem. A, 2018, 6, 160–166.
- 42 C. Ge, J. Song, Z. F. Qin, J. G. Wang and W. B. Fan, ACS Appl. Mater. Interfaces, 2016, 8, 18849–18859.
- 43 H. Wei, S. Deng, B. Hu, Z. Chen, B. Wang, J. Huang and G. Yu, ChemSusChem, 2012, 5, 2354–2360.
- 44 S. Ello, L. K. C. de Souza, A. Trokourey and M. Jaroniec, Microporous Mesoporous Mater., 2013, 180, 280–283.
- 45 E. M. Kutorglo, F. Hassouna, A. Beltzung, D. Kopecky, I. Sedlařova and M. Šooš, Chem. Eng. J., 2019, 360, 1199– 1212.
- 46 W. Xing, C. Liu, Z. Zhou, J. Zhou, G. Wang, S. Zhuo, Q. Xue, L. Song and Z. Yan, Nanoscale Res. Lett., 2014, 9, 1–8.
- 47 J. He, J. W. F. To, P. C. Psarras, H. Yan, T. Atkinson, R. T. Holmes, D. Nordlund, Z. Bao and J. Wilcox, Adv. Energy Mater., 2016, 6, 1–11.
- 48 J. C. Wang and Q. Liu, Nanoscale, 2014, 6, 4148–4156.
- 49 Z. Liu, Z. Zhang, Z. J. Jia, L. Zhao, T. T. Zhang, W. Xing, S. Komarneni, F. Subhan and Z. F. Yan, Chem. Eng. J., 2018, 337, 290–299. Paper

39 L.J.Xie, G.H. San, F. V. San, X. Q. Gao, Q. Q. Kang, X. M. Li_s 49 Z. Lis 2. Zamg, Z. J. Jih, L. Zhan, T. T. Zhang, W. Xing, C. C. C. Nan, C. C. Nan, C. C. Nan, C. C. N. (2019. Downloaded on 26 July 2019. Downlo
	- 50 S. Ghosh and S. Ramaprabhu, Carbon, 2019, 141, 692–703.
	- 51 X. Li, Z. Y. Sui, Y. N. Sun, P. W. Xiao, X. Y. Wang and B. H. Han, Microporous Mesoporous Mater., 2018, 257, 85–91.
	- 52 L. S. Shao, M. Q. Liu, J. H. Huang and Y. N. Liu, J. Colloid Interface Sci., 2018, 513, 304–313.
	- 53 W. Kiciński, M. Szala and M. Bystrzejewski, Carbon, 2014, 68, 1–32.
	- 54 S. A. Wohlgemuth, R. J. White, M. G. Willinger, M. M. Titirici and M. Antonietti, Green Chem., 2012, 14, 1515–1523.
	- 55 X. M. Ren, H. Li, J. Chen, L. J. Wei, A. Modak, H. Q. Yang and Q. H. Yang, Carbon, 2017, 114, 473–481.
	- 56 L. S. Shao, Y. Li, J. H. Huang and Y. N. Liu, Ind. Eng. Chem. Res., 2018, 57, 2856–2865.
	- 57 J. C. Geng, D. M. Xue, X. Q. Liu, Y. Q. Shi and L. B. Sun, AIChE J., 2017, 63, 1648–1658.
	- 58 M. Q. Liu, L. S. Shao, J. H. Huang and Y. N. Liu, Microporous Mesoporous Mater., 2018, 264, 104–111.