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Facile synthesis of functionalized porous carbon from bitumite mixed with waste powder char for excellent wastewater purification

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Waste powder char (WPC) is harmful to the environment and abundant in the production process of the coal chemical industry, but produces relatively low heat energy and releases large amounts of coal ash if it is burned directly. To efficiently solve this issue and create high value materials, three-dimensional hierarchical porous carbons (HPCs) with abundant oxygen-containing groups were prepared from Yulin bitumite (YL) mixed with WPC by combined pyrolysis carbonization and steam activation. The effects of preparation parameters, including WPC doping amount, activation temperature, and activation time, on pore structure development and adsorption capacity of methylene blue from wastewater were explored in detail. The obtained HPCs are microporous structures with additional mesopores and their largest specific surface area (SSA) reached 1299.75 m² g⁻¹ (8-YLHPC-950-3). Furthermore, 8-YLHPC-900-3 presents a high oxygen content of 15.92 wt%. The SSA and total pore volume (1081.67 m² g⁻¹ and 0.685 cm³ g⁻¹) of the HPC with 8 wt% WPC doping are larger than those without WPC doping (1155.36 m² g⁻¹ and 0.755 cm³ g⁻¹). Moreover, these HPCs show superior adsorption of methylene blue in aqueous solution (301 mg g⁻¹ for 8-YLHPC-900-3 compared with 198 mg g⁻¹ for 0-YLHPC-900-3), especially when the HPC was prepared with a WPC doping amount of 8 wt%, activation temperature of 900 °C, steam flow of 0.8 mL g⁻¹ h⁻¹, and activation time of 3 h. These conditions give an optimal adsorption capacity of 301 mg g⁻¹ methylene blue adsorption value (MBV). We introduce a value-added utilization approach for coal-based solid waste and a potential feedstock for adsorption material in the purification of wastewater. The results achieved in this study provide a green and environmentally friendly method for the preparation of HPCs from waste powder char that not only is an industrial reference for the removal of methylene blue from wastewater by HPCs but also contributes to sustainable development in the coal chemical industry.

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Sustainability spotlight

Methylene blue in wastewater is extremely difficult to biodegrade due to its stability and flexibility; this endangers biological safety and has become one of the most challenging environmental problems in the international community. We innovatively prepared oxygen-rich hierarchical porous adsorption materials from low-rank coal and waste powder char by a synergistic carbonization and mild activation method. These materials can not only effectively remove methylene blue from wastewater but also realize the high value-added utilization of coal-based solid waste and solve the problem of green and sustainable development in the coal chemical industry. This research contributes to water purification and safety and aligns with *Goal 6: Clean Water and Sanitation* of the UN SDGs.

1. Introduction

Industrial development and human activities have resulted in increasingly severe water pollution, so the effective removal of methylene blue has attracted broad global attention.^{1,2} Methylene blue removal from textile industry wastewater is an urgent and serious problem. Synthetic methylene blue is difficult to

biodegrade owing to its stronger stability and flexibility compared to natural dyes. Thus, effective methylene blue removal from wastewater has become one of the most challenging environmental issues for international society.³ In this respect, multiple methods, such as membrane separation,⁴ chemical reaction binding⁵ and adsorption,⁶ have been applied to the adsorption of methylene blue. The commonly used adsorbents are porous carbons (PCs), zeolite, carbon aerogels, MOFs, carbon nanotubes and graphene sheets.^{7,8} Among these, PCs have become one of the most competitive wastewater treatment materials because of their high efficiency, simple operation, low cost, and stable physicochemical properties.^{9,10}

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PCs are primarily prepared from biomass wastes, petroleum, coal, and their derived products through a template-based method, chemical activation or physical activation.¹¹ While a variety of raw materials can be used, coal and its derivatives are still the main raw material for PCs due to their abundance, low cost, and high carbon content.

Hierarchical porous carbon, which has various proportions of micropores, mesopores and macropores, can achieve admirable adsorption behavior based on its particular pore size distribution, developed porosity, and abundant functional groups, as adsorbate molecules can be transferred rapidly in the pore channels of HPCs.¹² It is worth noting that micro/meso/macropores play different roles in the adsorption process. The micropores of HPCs mainly determine the adsorption capacity owing to their large proportion of the SSA and pore volume and can offer abundant adsorption sites for methylene blue molecules. Further, the mesopores are regarded as the transmission channel for molecules, while the effect of macropores is as a buffer which ensures that molecules can enter the pore channels quickly.¹³ Zhuang *et al.*¹⁴ adopted coal tar pitch as a precursor to prepare HPCs, which had excellent adsorption of methylene blue. Mbarki *et al.*¹⁵ prepared HPCs from corncobs by hydrothermal carbonization and CO₂ activation; the specific surface area reached 1111 m² g⁻¹ and the methylene blue adsorption reached 561.5 mg g⁻¹ at 25 °C. The surface of the HPC was found to be mainly basic, with a total amount of basic groups of 0.743 mmol g⁻¹ compared to 0.550 mmol g⁻¹ of acidic groups. The abundant functional groups are conducive to boosting the application of HPCs.¹⁶ Furthermore, chemical activation usually requires three to four times as many activators as the raw material, including KOH, ZnCl₂ and NaOH, which are more expensive, corrosive to the equipment and produce more corrosive gases during activation processes.^{9,17} Simultaneously, the resulting porous carbon requires an additional pickling step to achieve neutrality. In contrast, steam activation is a green and sustainable preparation process that allows the large-scale industrial production of HPCs due to its low cost, simple operation and non-corrosive properties.¹⁸

Waste powder char (WPC) is a typical coal-based solid waste produced in large quantities by the coal chemical industry. WPC shows higher ash content and lower moisture content and volatile content than raw coal; it is a manual heat treatment by-product after high temperature carbonization. Currently, WPC is thrown away or directly buried, resulting in environmental pollution and the waste of coal resources. WPC exhibits low heat energy release and a high ignition point which makes it difficult to burn, although it can be recycled by combustion. However, WPC contains abundant carbon and oxygen contents which facilitate the preparation of HPCs for methylene blue removal from wastewater when mixed with bituminous coal. It is worth noting that this process could achieve the high value-added utilization of coal-based solid waste (WPC) and the environmentally friendly preparation of three-dimensional porous carbon. Coal is the primary energy resource in China and Yulin low metamorphism bitumite has abundant reserves, around 149 billion tons in Shaanxi, China. Although the energy density of low-rank bituminous coal is poor, its high reactivity

and oxygen-containing functional groups have attracted it increased attention as a precursor for HPCs. However, in liquid-phase adsorption applications, HPCs with abundant oxygen-containing functional groups obtained from low-grade bituminous coal mixed with coal-based solid waste by synergistic pyrolytic carbonization and steam activation have rarely been reported.

In this study, WPC and YL were adopted as the raw materials to prepare three-dimensional HPCs by a typical two-step method for the purification of wastewater containing methylene blue. First, the raw material was pyrolysis carbonized in a tube furnace and then the obtained coal char was activated by steam. The effects of WPC doping ratio, activation temperature, and activation time on the pore structure and adsorption performance of HPCs were explored. In addition, a semi-quantitative analysis of the surface oxygen-containing functional groups of the precursors and HPC was discussed. Herein, coal-based solid waste is used to prepare a high value-added porous adsorption material for wastewater renovation, which is always attractive and significant. The results achieved in this study not only exhibit the industrial significance of HPCs for the removal of methylene blue from wastewater, but also helpfully improve the high value-added utilization of coal-based solid waste and bitumite.

2. Experimental

2.1 Materials

Yulin bitumite collected from Shaanxi, China and waste powder char provided by Shaanxi Coal and Chemical Industry Group Co. Ltd. were used as raw materials in this research. The proximate and ultimate analyses of YL and WPC are listed in Table 1. Reagent grade chemicals were used in all tests, as shown in Table 2. The purity of the chemicals was determined by gas chromatography (GC). Methylene blue (MB) was provided by Xilong Chemical Co. Ltd. Sodium thiosulfate, hydrochloric acid and starch were purchased from Tianjin Chemical Reagent Research Institute Co. Ltd.

2.2 Preparation of HPCs

The HPCs were prepared by a representative two-stage process of carbonization and steam activation under N₂ flow at a heating rate of 10 °C min⁻¹. In step 1 (carbonization), the YL mixed with WPC was filled into a home-made mold and extruded into a cake block with a diameter of 25 mm and a thickness of 8 mm

Table 1 Proximate and ultimate analyses of YL and WPC^a

Sample	Proximate analysis (wt%)				Ultimate analysis (wt%)				
	M _{ad}	A _d	VM _{daf}	FC _{daf} ^b	C _{daf}	H _{daf}	O _{daf} ^b	N _{daf}	S _{daf}
YL	6.49	2.51	29.75	68.48	75.04	4.76	18.21	1.28	0.71
WPC	5.18	7.78	8.50	84.38	78.26	1.74	19.25	0.64	0.11

^a A: ash, M: moisture, FC: fixed carbon, VM: volatile matter, ad: air dry basis, d: dry basis, daf: dry ash-free, b: by difference.



Table 2 CAS registry number, mass fraction purity, and source of the chemicals

Component	CAS number	Supplier	Purity (mass fraction)	Analysis method
Methylene blue	61-73-4	Xilong Chemical Co. Ltd	0.99	GC
Sodium thiosulfate	2319-84-8	Tianjin Chemical Reagent Research Institute Co. Ltd	0.99	GC
Hydrochloric acid	7647-01-0	Tianjin Chemical Reagent Research Institute Co. Ltd	0.37	GC
Starch	9005-84-9	Tianjin Chemical Reagent Research Institute Co. Ltd	0.99	GC

under a pressure of 200 MPa. After natural air drying, it was broken into granular material with particle sizes of 3–10 mm. Subsequently, the mixture was carbonized at 600 °C for 45 min in a horizontal tube furnace. The obtained char is denoted as *x*-YLChar, where *x* = 0, 4, 8, and 12 for WPC weight percentages of 0%, 4%, 8%, and 12%. In step 2 (steam activation), the *x*-YLChar was activated at different temperatures (850, 900 and 950 °C), and times (2, 3 and 4 h) with 0.8 mL g⁻¹ h⁻¹ steam flow. Then, the prepared hierarchical porous carbons samples were dried at 150 °C for 3 h in a vacuum drying oven. The HPCs were denoted as *x*-YLHPC-*w*-*y* (*x* = 0, 4, 8 and 12, referring to the weight percentages of WPC of 0%, 4%, 8% and 12%, respectively; *w* = 850, 900, and 950, referring to the activation temperatures of 850, 900, and 950 °C, respectively; *y* = 2, 3, and 4, referring to the activation times of 2, 3, and 4 h, respectively). A schematic illustration of the preparation process of HPCs is shown in Fig. 1 and all experiments were carried out in parallel.

2.3 Material characterization

The ultimate analyses of YL, WPC, and HPC were characterized by an elemental analyzer (Vario EL III, German). All samples which need to be characterized were pulverized below 200 mesh and dried at 150 °C for 12 h. The microcrystalline structures were characterized by X-ray diffraction (XRD) measurements on a Bruker D8 Advance X-ray diffractometer using Cu K α radiation (λ = 0.15418 nm) operating at 40 mA and 40 kV (10° min⁻¹ from 10 to 90°). Raman analysis of the YL, 8-YLChar and HPC samples was carried out on a Bruker Senterra Raman spectrometer by 532 nm laser excitation using a microscope. The Raman spectra were obtained in the range of 1000–2000 cm⁻¹ and all scans were performed three times. Field-emission

scanning electron microscopy (FESEM) images were taken by a Merlin Zeiss at 5.0 kV to observe the microscopic appearance of the samples. The surface chemical properties of the samples were characterized qualitatively using a Nicolet Magna IR560 Fourier-transform infrared (FTIR) spectrometer. X-ray photoelectron spectroscopy (XPS) data were collected using a Thermo ESCALAB 250Xi spectrometer to analyze the surface oxygen-containing functional groups. N₂ adsorption–desorption isotherms were measured at –196 °C with a Quantachrome Autosorb-iQ instrument. The SSAs were obtained by applying the Brunauer–Emmett–Teller (BET) method. Micropore surface area and micropore volume were calculated using the *t*-plot method. Total pore volume was determined at a relative pressure *P*/*P*₀ of 0.99. The pore size distribution was calculated from N₂ adsorption data using the density functional theory (DFT) model method. The concentrations of MB were measured by a UV-visible spectrophotometer at the maximum absorbance wavelength of 665 nm.

2.4 Adsorption experiments

HPCs were selected as adsorbents to evaluate their adsorption performance for MB. 10 mg HPCs were added to 12 mL MB solution (*C*₀ = 150 mg L⁻¹) and the mixture was shaken at 30 °C and 200 rpm for 30 min. After the completion of adsorption, the MB concentration was determined by spectrophotometry at 665 nm (TU-1900 double beam UV-vis spectrophotometer). The methylene blue adsorption capacity MBV (mg g⁻¹) was calculated based on the equation

$$MBV = (C_0 - C) \frac{V}{W}$$

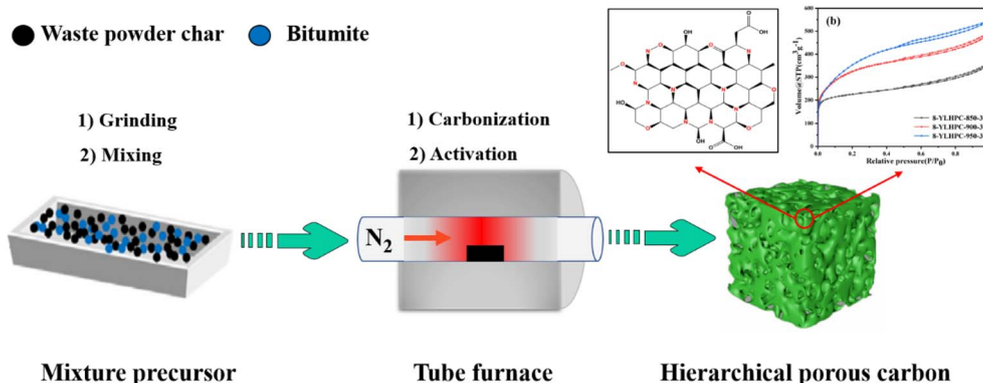


Fig. 1 Graphical schematic for the preparation of the HPCs.



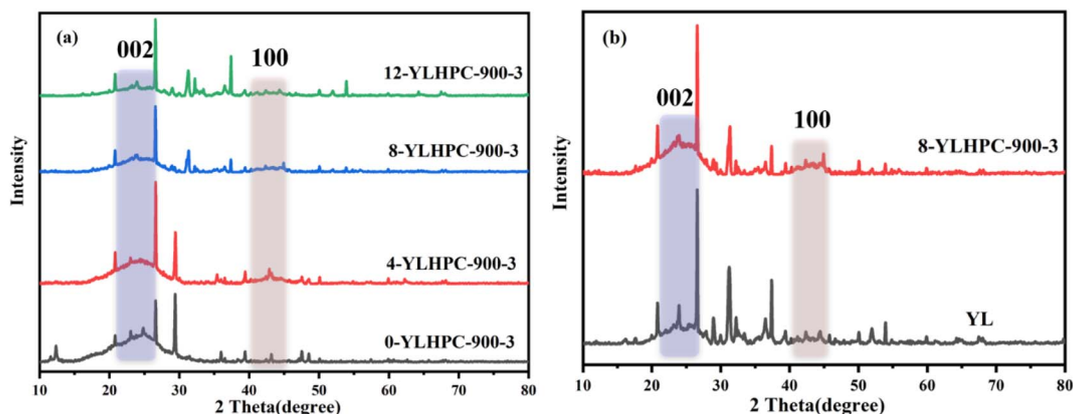


Fig. 2 XRD patterns of (a) x -YLHPC-900-3 and (b) YL and 8-YLHPC-900-3.

where C_0 and C (mg L^{-1}) are the methylene blue concentrations at the initial and final steps, respectively, V (L) is the volume of dye solution, and W (g) is the mass of HPCs.

3. Results and discussion

3.1 Morphology and microstructure characterization

As shown in Table 1, WPC has relatively low moisture content, high ash content and fixed carbon content compared with YL. Note that the higher carbon content makes WPC suitable for the preparation of porous carbon according to the ultimate analysis.

As a rapid and non-destructive technique, XRD is one of the most commonly used methods for revealing the microcrystalline structure of coal char and HPCs.¹⁹ As shown in Fig. 2, these spectra include two distinct peaks at 22° and 43° which are attributed to the interlayer reflections of the (002) and (100) planes, respectively.²⁰ The (002) peaks of x -YLHPC- w - y in Fig. 2(a) became weak and faintly broadened, indicating the decrease of graphitization degree with the percentage of WPC increasing from 0 wt% to 12 wt%. Compared with that of YL, the (100) peak intensity of 8-YLHPC-900-3 at about 43° was significantly enhanced, as seen in Fig. 2(b), which demonstrates that 8-YLHPC-900-3 possesses amorphous character and consists of small areas of ordered graphene sheets after steam activation.

Raman spectroscopy was also used to measure the defective, disordered structure and crystallinity of the as-prepared samples. Thus, it was used to characterize the structural evolution of the 8-YLChar and 8-YLHPC-900-3 samples derived from YL and WPC under synergistic carbonization and steam activation to analyze the relationship between the amorphous carbon structure and graphitized carbon. Two characteristic peaks located at about 1342 cm^{-1} and 1579 cm^{-1} are identified in the Raman spectra (Fig. 3) and are attributed to the D-band (disordered or defective structure of carbon materials) and G-band (the sp^2 carbon-bonded graphitic structure), respectively.^{21,22} The amorphous degree of carbon materials is reflected by the value of I_D/I_G (I_D and I_G are the intensities of the D- and G-bands, respectively). Here, the I_D/I_G values of YL and 8-YLChar are 0.42 and 0.62, respectively. A slight intensity ratio

increase is observed with the use of carbonization which is related to the formation of polycyclic aromatic units *via* the condensation of the macromolecular network in the char. During this stage, the chemical bonds between the macromolecular compounds are broken to form micromolecules. These micromolecules deposit on the surface of the char, resulting in the formation of a defect structure. The I_D/I_G ratio increased to 0.98 after the activation of 8-YLChar. This indicates that the structure of 8-YLHPC-900-3 has more defects, likely because its crystalline structure is destroyed by severe steam activation at the temperature of 900°C .

As illustrated in Fig. 4, the thermal stability of WPC is better than that of YL. Even at 900°C , the mass loss of WPC is still less than 10%, because it is a carbonization product and many side and branched chains of polycyclic aromatic hydrocarbons have already been lost during this process. However, the mass loss of YL presents a big difference and could be divided into three stages. At the early stage of pyrolysis ($<400^\circ\text{C}$), the mass loss is mainly the gasification of free water and combined water and the decomposition of unstable organic small molecules to produce CO , CO_2 , CH_4 , *etc.* The main mass loss (about 30%) occurs at 400 – 700°C . This stage is mainly the decomposition

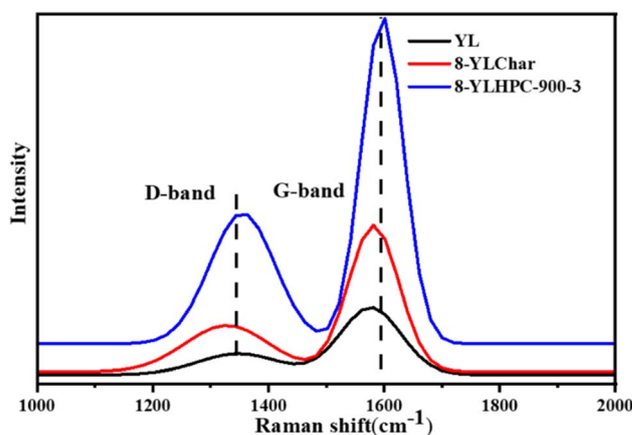


Fig. 3 Raman spectra of YL, 8-YLChar and 8-YLHPC-900-3.



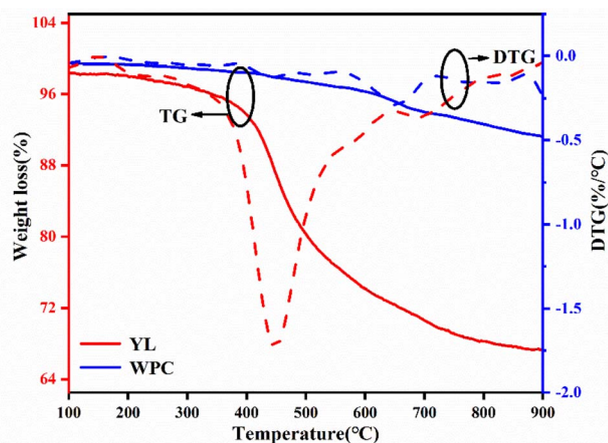


Fig. 4 TG and DTG curves of YL and WPC.

and integration of organic macromolecules, which produces large amounts of coal tar in addition to CO, CO₂ and CH₄. The third stage (>700 °C) has no obvious weight loss, indicating that the prepared porous carbon will show good thermal stability. Understanding the stability of the raw material provides a theoretical basis for the selection of the activation temperature.

0-YLChar and 8-YLChar were verified by FESEM to observe the microstructural differences of the samples after carbonization. According to the SEM graphs in Fig. 5(a) and (b), compared with the surface of 0-YLChar (Fig. 5(a)), the surface of 8-YLChar (Fig. 5(b)) is rougher and filled with many particles which are beneficial to the diffusion of the steam activator, thus promoting the development of a porous structure.²³ This also could be proved by the N₂ adsorption-desorption results of the *x*-YLHPC-900-3, as shown in Fig. 8(a). After steam activation, the surface of 8-YLHPC-900-3 presents a highly porous honeycomb structure with some irregular particles, seen in Fig. 5(c) and (d). 8-YLHPC-950-3, in Fig. 5(e) and (f), has a denser pore structure than 8-YLHPC-900-3 due to its complete activation at higher activation temperatures. Meanwhile, the prepared activated samples reveal a typical three-dimensional structure with, particularly in 8-YLHPC-950-3, a hierarchical porous structure with a pore size distribution made up of a combination of different pore types (macropores, mesopores and abundant micropores) after etching with an excess of steam. The three-dimensional hierarchical porous structure of the sample features a large number of active sites for subsequent adsorption. In addition, the prepared HPC shows well-developed porosity in its microcrystalline and amorphous structure. Overall, the FESEM results demonstrate the effectiveness of steam activation for pore structure development for the HPC from *x*-YLChar.

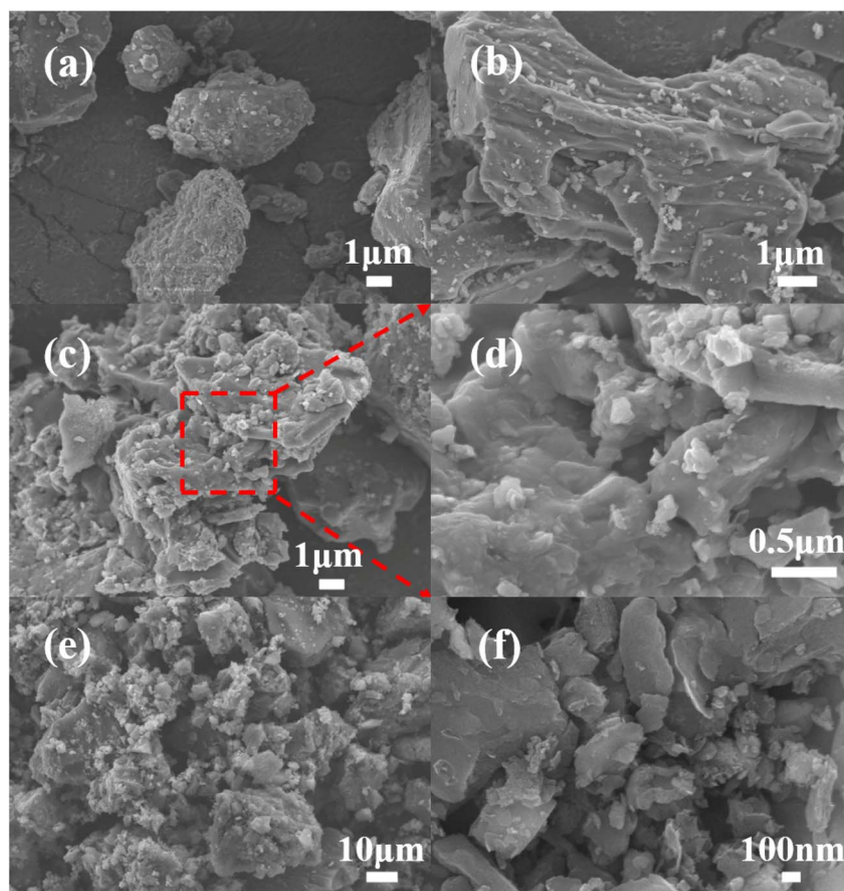


Fig. 5 SEM images of (a) 0-YLChar, (b) 8-YLChar, (c and d) 8-YLHPC-900-3 and (e and f) 8-YLHPC-950-3.



Table 3 Ultimate and XPS analyses of YL and 8-YLHPC-900-3

Sample	Ultimate analysis (wt%)							XPS analysis ^c (%)		
	O/C	H/C	C _{daf}	H _{daf}	O _{daf} ^a	N _{daf}	S _{daf} ^b	C 1s	O 1s	N 1s
YL	0.243	0.063	75.04	4.76	18.21	1.28	0.71	81.00	17.68	1.33
8-YLHPC-900-3	0.135	0.071	81.57	5.78	11.03	0.96	0.66	83.74	15.92	0.33

^a Calculated by difference. ^b Total sulfur in dried basis. ^c Atomic ratio.

3.2 Surface chemistry analysis

Ultimate and XPS analysis results of YL and 8-YLHPC-900-3 are shown in Table 3. The oxygen content of YL exhibits an obvious decrease from 18.21 wt% to 11.03 wt% after carbonization and activation treatment, whereas the carbon content increased significantly, indicating the partial removal of oxygen-containing functional groups and an improvement of the properties of YL. Additionally, the atomic ratio of O/C was obviously reduced, illustrating that dehydration, decarboxylation, aromatization, and integration reactions occurred during the preparation of HPCs.

FTIR spectroscopy was employed to detect the surface chemical properties of YL, 8-YLChar and 8-YLHPC-900-3 derived from the different thermal conversion processes. As shown in Fig. 6, it is obvious that the oxygen-containing functional groups of 8-YLHPC-900-3 are different from those of YL and 8-YLChar based on the spectral patterns of 8-YLHPC-900-3, 8-YLChar and YL over the range of 4000–400 cm⁻¹. In general, the side chains in YL are mainly composed of methoxyl, phenolic hydroxyl and aldehyde groups. 8-YLHPC-900-3 contains an abundance of oxygen-containing groups, such as phenolic hydroxyl, carboxyl, carbonyl, and aromatic ether groups, according to its FTIR spectrum (Fig. 6). In addition, condensation polymerization of small side chains and branched organic molecules also occurs to form the coal char polymer framework of HPC, which may be due to the synergistic carbonization and integrated activation reactions of YL and WPC. The carbon polymer framework of

HPC is composed of dense aromatic hydrocarbons with many stable oxygen-containing functional groups. Based on Fig. 6, the functional groups of 8-YLHPC-900-3 are –OH at 3432 cm⁻¹, –CH in arenes/epoxide at 3250–3000 cm⁻¹, C=O at 1625 cm⁻¹, –OH/C=C in arenes at 1410 cm⁻¹, and aryl ether at 1200–1005 cm⁻¹.²¹ Compared with YL and 8-YLChar, the –COOH (1620 cm⁻¹) and –OH (1350–1000 cm⁻¹) contents of 8-YLHPC-900-3 are relatively low based on its weaker peak intensities; this is attributed to intramolecular and intermolecular decarboxylation, dehydration and aromatization during the preparation process.^{24,25} However, the intensities of the –OH (3600–3200 cm⁻¹) peak and C–O (1300–1100 cm⁻¹) of ether peak present a slight increase. This might be due to thermochemical reaction processes causing structural damage to the organic macromolecules of 8-YLHPC-900-3, thus generating new functional groups. Some scholars have reported that steam works not only as a dehydrating agent but also as an oxidizing agent during the activation process. Furthermore, these oxygen-containing species improve the wastewater purification performance of the HPCs *via* boosted wettability and the formation of chemisorption active sites.

To further confirm the evolution of the surface composition and chemical state during the preparation process, semi-quantitative investigations of YL and 8-YLHPC-900-3 were carried out with an XPS analyzer.^{26,27} The high-resolution C 1s and O 1s peaks were curve-fitted using the XPSPeak software in accordance with relevant literature and the test results are similar to the FTIR results. As shown in Fig. 7 and Table 4, the contents of C=C/C–C and C=O increased from 53.40% to 60.27% and 16.81% to 46.87%, respectively, while the contents of C–OH and COO exhibited a significant decline. The C–OH and –COOH might be transformed to C=O and C=C/C–C, respectively, which agrees with the foregoing analysis of the FTIR spectra. This phenomenon may be caused by the condensation polymerization and aromatization of organic molecules at high temperatures, where some of the oxygen-containing functional groups are released as water, carbon dioxide, and small molecules of coal tar. Although the oxygen content of the prepared HPCs was reduced, the figure is still as high as 15.92%, as seen in the XPS analysis in Table 4. The obtained oxygen-rich hierarchical porous carbon should show superior performance in adsorption applications due to its high wettability and number of active sites. The presumed surface oxygen-containing functional groups of 8-YLHPC-900-3 are shown in Fig. 7(f), based on the ultimate analysis, FTIR and XPS characterization analysis results.

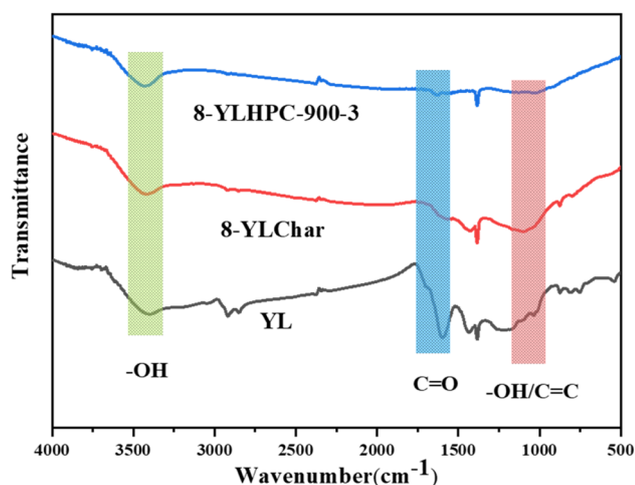


Fig. 6 FTIR spectra of YL, 8-YLChar and 8-YLHPC-900-3.



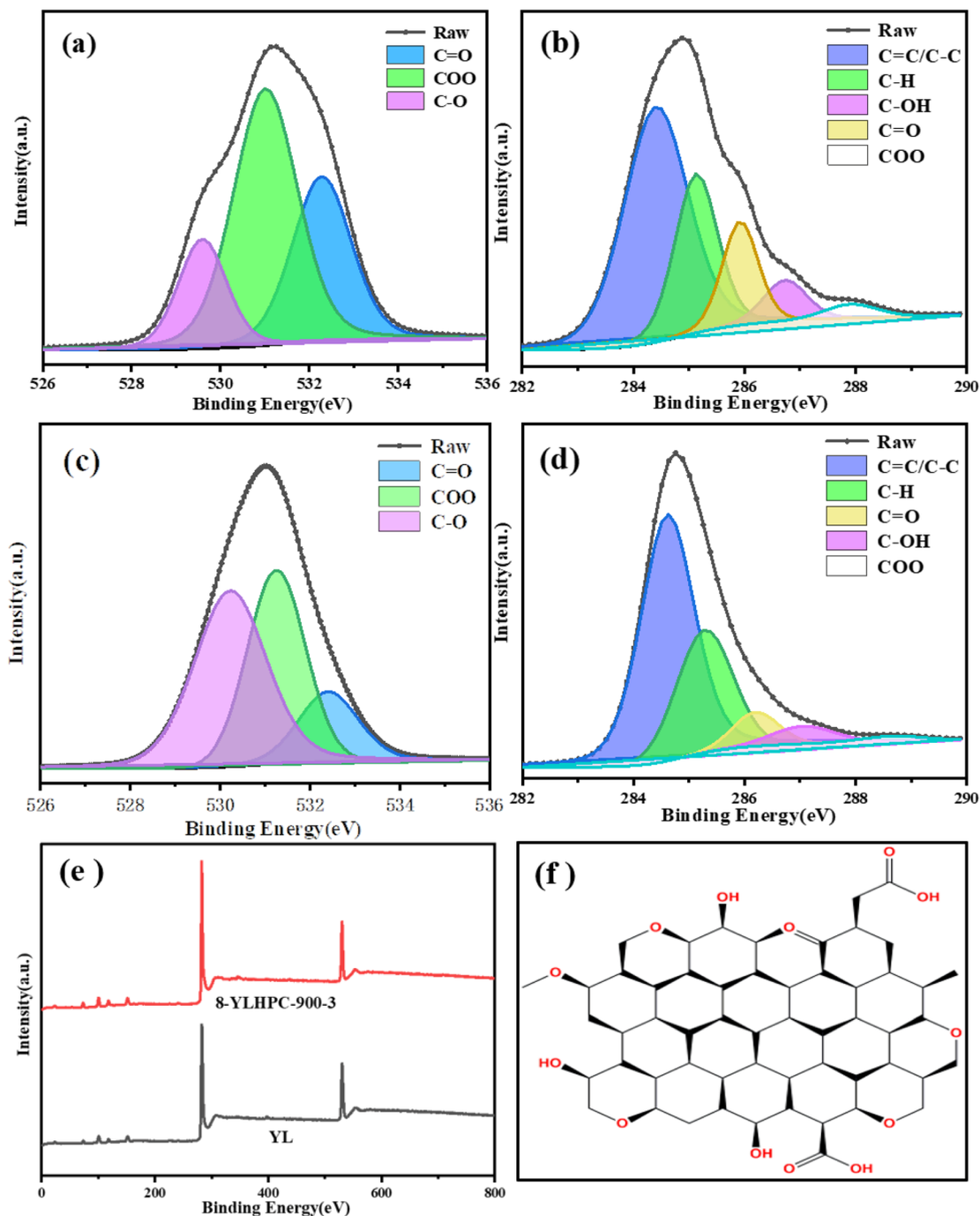


Fig. 7 High-resolution XPS spectra of (a and c) O 1s and (b and d) C 1s of YL and 8-YLHPC-900-3; (e) XPS spectra of YL and 8-YLHPC-900-3; (f) schematic of oxygen-containing functional groups in the carbon lattice.

Table 4 Relative contents of the functional groups of YL and 8-YLHPC-900-3 in the XPS spectra

Sample	C 1s						O 1s		
	O/C ^a	C=C/C (%)	C-H (%)	C-OH (%)	C=O (%)	COO (%)	C=O (%)	COO (%)	C-O (%)
YL	0.218	53.40	24.12	13.25	6.42	2.81	16.81	52.35	30.84
8-YLHPC-900-3	0.190	60.27	27.76	6.28	4.42	1.26	46.89	37.74	15.36

^a Atomic ratio.



Table 5 Textural characteristics of *x*-YLHPC-*w*-*y* under different preparation conditions

Sample	S_{BET}^a ($\text{m}^2 \text{g}^{-1}$)	V_t^b ($\text{cm}^3 \text{g}^{-1}$)	V_{micro}^c ($\text{cm}^3 \text{g}^{-1}$)	V_{meso}^d ($\text{cm}^3 \text{g}^{-1}$)	D_{ave}^e (nm)	Yield (wt%)
0-YLHPC-900-3	1081.67	0.685	0.148	0.452	2.47	25.2
4-YLHPC-900-3	1165.07	0.722	0.127	0.471	2.48	18.0
8-YLHPC-900-3	1155.36	0.755	0.077	0.620	2.61	15.7
12-YLHPC-900-3	1105.30	0.769	0.097	0.608	2.78	15.6
8-YLHPC-850-3	845.92	0.552	0.188	0.328	2.41	31.7
8-YLHPC-950-3	1299.75	0.853	0.210	0.643	2.86	5.9
8-YLHPC-900-2	926.56	0.564	0.189	0.315	2.44	28.7
8-YLHPC-900-4	916.71	0.676	0.047	0.637	2.95	10.7

^a S_{BET} : specific surface area from multiple BET method. ^b V_t : total pore volume at $P/P_0 = 0.99$. ^c V_{mic} : *t*-method micropore surface area and micropore volume. ^d V_{meso} : difference of V_t and V_{mic} . ^e D_{ave} : average pore diameter.

3.3 Textural properties and adsorption performance

The effects of WPC doping ratio, activation temperature, and activation time on the yield and porous structure of HPCs are shown in Table 5. The pore structures of *x*-YLHPC-*w*-*y* were investigated by N_2 adsorption-desorption experiments, in accordance with previous reports.^{28,29} The adsorption performance was evaluated by MBV.

3.2.1 Effect of WPC doping ratio. The WPC doping ratio has a great influence on the adsorption performance of HPCs and the main purpose of doping WPC is to simultaneously recycle coal-based solid waste and promote porous structural development. As shown in Fig. 8(a) and (b), the isotherm of *x*-YLHPC-900-3 is a combination of type I and IV with an H4 hysteresis loop, according to the IUPAC classification.^{20,30} The pore structure of *x*-YLHPC-900-3 includes micropores and mesopores, with micropores occupying a large proportion. In addition, small tails at the P/P_0 near 1.0 reveal the presence of macropores. The adsorption-desorption isotherm curve of 8-YLHPC-900-3 is at the top while that of 0-YLHPC-900-3 is at the bottom, indicating that the pore structure development of HPCs prepared by adding 8 wt% WPC is the largest. The SSA, pore volume, and average pore size parameters of the HPCs are given in Table 5. It can be seen that the SSA, micropore volume, micropore percentage and total pore volume of *x*-YLHPC-900-3 first increased and then reduced with the rise of the proportion of WPC. Moreover, the SSAs and total volumes of 4-YLHPC-900-3, 8-YLHPC-900-3 and 12-YLHPC-900-3 are larger than those of 0-YLHPC-900-3. This is because the WPC has a lower volatile content and higher fixed carbon content and is able to undergo complete oxidation with steam after the second carbonization, which is conducive to promoting the development of pore structure. Due to the appropriate WPC doping ratio, 8-YLHPC-900-3 reaches the largest SSA of $1155.36 \text{ m}^2 \text{g}^{-1}$ and pore volume of $0.755 \text{ cm}^3 \text{g}^{-1}$, indicating that the optimal WPC doping ratio is 8 wt%.

Porous carbon is limited by the diameter of adsorbate molecules when adsorbing substances. Studies have shown that the pore diameter of porous carbon should be 1.7 times the diameter of adsorbate molecules for effective adsorption.^{31,32} Generally, the MBV is used to reflect the development degree of pores with a diameter of 1.5–10 nm.^{20,33} As shown in Fig. 9, the MBVs of *x*-YLHPC-900-3 significantly differ at different WPC

doping ratios. With the WPC doping ratios from 0 wt% to 12 wt%, the MBVs are 198, 230, 301 and 271 mg g^{-1} , showing a trend of first increasing and then decreasing with the rise of WPC doping ratio. The trend shows that WPC doping is advantageous to achieving a more intense activation reaction due to the mixture containing more carbon matrix. However, excessive amounts of WPC lead to forming fewer initial micropores, which means obtaining fewer mesopores under the same activation reaction conditions. The adsorption performance is consistent with the nitrogen adsorption and desorption characterization results, demonstrating that an 8 wt% WPC doping ratio is optimal.

3.2.2 Effect of activation temperature. Activation temperature is one of the most important factors for controlling activation extent.³⁴ As shown in Fig. 8(c) and (d), the nitrogen adsorption-desorption isotherms and pore size distributions of 8-YLHPC-*w*-3 changed significantly with the increase of activation temperature (850, 900 and $950 \text{ }^\circ\text{C}$). The total pore volume of 8-YLHPC-950-3 is quite large compared with that of 8-YLHPC-850-3, suggesting that 8-YLHPC-950-3 has more pores. Moreover, the isotherm of 8-YLHPC-950-3 exhibits a combination of type I and IV with a larger hysteresis loop,³⁵ demonstrating that 8-YLHPC-950-3 possesses a further proportion of mesopores and macropores. However, the curve of 8-YLHPC-850-3 shows a different trend, with the N_2 adsorption mainly occurring at a relative pressure less than 0.2 and a horizontal plateau at high relative pressure, suggesting that 8-YLHPC-850-3 is mainly micropores. Furthermore, the SSA and total volume increased to $1299.75 \text{ m}^2 \text{g}^{-1}$ and $0.853 \text{ cm}^3 \text{g}^{-1}$, respectively, with the increase of the activation temperature from 850 to $950 \text{ }^\circ\text{C}$, as seen in Table 5. It is worth noting that the higher activation temperature is not only conducive to the formation of micropores but also can transform the micropores into mesopores and macropores owing to the more intense steam activation reaction. Therefore, the average pore size sharply increased from 2.41 nm in 8-YLHPC-850-3 to 2.86 nm in 8-YLHPC-950-3.

The MBVs of 8-YLHPC-*w*-3 prepared at different steam activation temperatures present significant differences, as seen in Fig. 9. The MBVs of 8-YLHPC-900-3 (301 mg g^{-1}) and 8-YLHPC-950-3 (295 mg g^{-1}) are significantly larger than that of 8-YLHPC-850-3 (160 mg g^{-1}). This is because the pore structure development of 8-YLHPC-900-3 and 8-YLHPC-950-3 is excellent



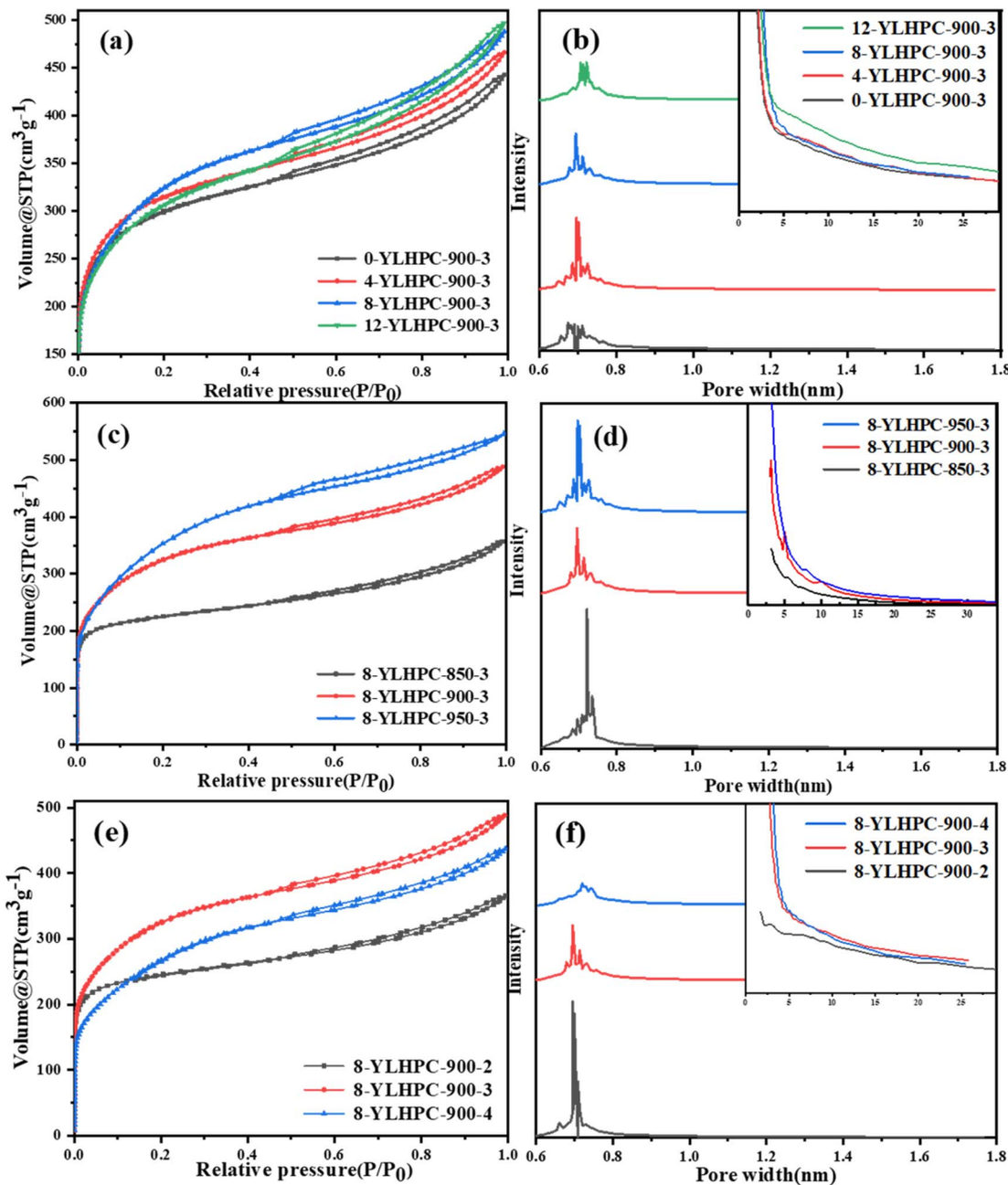


Fig. 8 Nitrogen adsorption isotherms and pore size distribution patterns of *x*-YLHPC-*w*-*y* with different WPC doping ratios (a and b), activation temperatures (c and d) and activation times (e and f).

due to the more complete activation reaction between the mixture and steam, exhibiting a high adsorption performance of methylene blue. The MBV adsorption performances are similar to the nitrogen adsorption and desorption results, but excessive activation temperature decreases the yield (5.9% of 8-YLHPC-950-3) of HPCs.³⁶ Therefore, the optimum activation temperature was selected to be 900 °C by considering the heating energy consumption and yield of HPCs.

3.2.3 Effect of activation time. In this part, the HPCs were prepared under different activation times from 2 to 4 h at the activation temperature of 900 °C. It can be concluded that the

proportion of mesopores increased significantly with the rise of activation time according to the larger hysteresis loop (8-YLHPC-900-4) of the N₂ adsorption-desorption isotherm curves in Fig. 8(e) and the pore size distribution patterns in Fig. 8(f). Moreover, the mesopore volumes and the average pore sizes increase from 0.564 cm³ g⁻¹ and 2.44 nm in 8-YLHPC-900-2 to 0.676 cm³ g⁻¹ and 2.95 nm in 8-YLHPC-900-4, respectively, also indicating that the percentage of mesopores in the HPCs increases with the increase of activation time. It shows that the pore enlargement reaction (micropores etched into mesopores or macropores) is dominant at longer activation times.



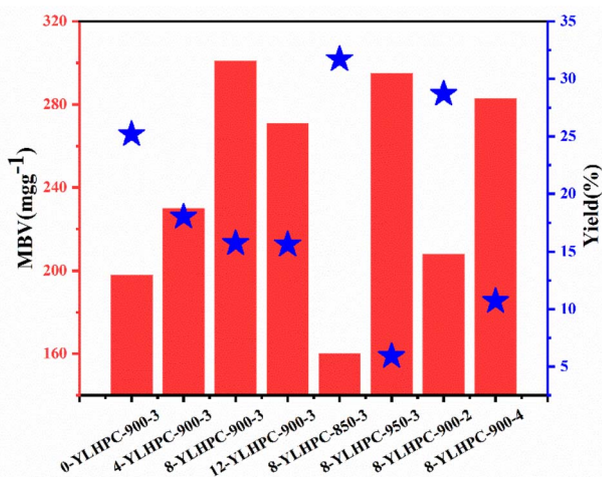


Fig. 9 Methylene blue adsorption values and yields of x -YLHPC- w - y with different WPC doping ratios, activation temperatures and activation times.

The methylene blue adsorption behaviors of HPCs show a good positive correlation with SSA and total pore volume; that is, the MBVs (Fig. 9) increase with the rise of SSA and total pore volume, which are closely related to the activation time. However, an overlong activation time also leads to ablation and collapse of the hole wall, resulting in a poorer MBV (the MBV of 8-YLHPC-900-4 is less than that of 8-YLHPC-900-3). Normally, the adsorption performance of HPCs mainly depends on micropores and small size mesopores, due to their large proportion of pore volume and specific surface area. However, the channel effect of macropores and transitional role of mesopores allow the adsorbate molecules pass into the micropores, which can promote the adsorption performance of micropores and small mesopores. Sometimes mesopores can also absorb a certain amount of macromolecular organic substances in the liquid phase.³⁷ Therefore, porous carbon with a wider distribution of micropores, mesopores and macropores prepared at a longer activation time shows better adsorption of methylene blue.

4. Conclusion

Three-dimensional structural HPCs with excellent methylene blue adsorption performance in aqueous solution were successfully prepared through a synergistic pyrolysis carbonization and steam activation reaction of bitumite mixed with waste powder char. The SSA and pore size distribution of HPCs can be efficiently controlled by different activation temperatures and times. The obtained HPCs present a better hierarchical pore structure and more excellent adsorption performance than materials prepared without WPC doping. 8-YLHPC-950-3 shows an excellent SSA and total pore volume up to 1299.75 m² g⁻¹ and 0.853 cm³ g⁻¹, respectively. The obtained HPCs exhibit good adsorption performance and the highest MBV of 8-YLHPC-900-3 reaches 301 mg g⁻¹. Meanwhile, the prepared HPCs possess abundant oxygen-containing functional

groups and the highest oxygen content reached 15.92 wt%, according to the XPS analysis. Hence, this mild, sustainable and industrially available strategy not only prepares promising coal-based HPCs for adsorption materials with a better performance but also achieves the high value-added utilization of the coal-based solid waste. Future work will focus on the evolution mechanism of the oxygen-containing functional groups of HPCs during the preparation process by *in situ* characterization techniques.

Author contributions

Yufeng Yin provided the idea of this work and designed relevant experiments, analyzed experiment results and composed manuscript. Yiting Zhao prepared and characterized the samples, and analyzed experiment results. Jing Wang offered great help during the process of experiment. All authors reviewed the manuscript.

Conflicts of interest

There are no conflicts to declare.

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