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1	Preparation of activated carbon from Xinjiang region coal by microwave
2	activation and its application in naphthalene, phenanthrene, and pyrene
3	adsorption
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23 Abstract:

Coal-based activated carbons (CAC) were prepared from coal in Xinjiang China by 24 microwave-assisted activation. The effects of preparation conditions on adsorption capabilities of 25 CAC for methylene blue (MB) and iodine were investigated. The microwave activation was carried 26 27 out with coal alkali ratio (CAR) of 1:3 for 12 min at 700 W of power, the adsorption capacities of the prepared CAC achieved the maximal values of 312 and 1048.19 mg/g for MB and iodine. The 28 increasing of CAR, activation time, and microwave power reduced the surface functional groups and 29 improved the pH_{PZC} of CAC. The BET surface area and total pore volume of CAC prepared were 30 identified as 1770.49 m^2/g and 0.99 cm³/g, respectively. Hence, the work demonstrates that the use 31 of microwave heating for preparing CAC is more rapid, cheaper and efficient. The adsorption 32 behaviors of typical polycyclic aromatic hydrocarbons (PAHs) in ethanol aqueous solution on CAC 33 34 were also explored. Equilibrium data were favorably described by the Langmuir and Freundlich isotherms, and adsorption kinetics was fitted to the pseudo-second-order model well. The 35 equilibrium adsorption amounts were 117.17 mg/g for pyrene, 111.39 mg/g for phenanthrene, and 36 78.18 mg/g for naphthalene. Thus, CAC prepared by microwave activation could have a very 37 effective application in removal of these PAHs. 38

39 Key words: Coal-based activated carbon; Microwave activation; PAHs; Adsorption

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45 **1. Introduction**

Activated carbons (AC) are porous material with extremely high surface areas; they have been 46 47 widely used in a variety of industrial applications such as separation/purification of liquids and gases, removal of toxic substances, catalysts and catalyst support, super capacitors, electrodes and gas 48 49 storage [1]. AC is usually prepared from different sources of raw materials utilizing different processing methods. Major raw materials for the preparation of AC are coal, petroleum, peat, wood 50 and agricultural wastes. Among which, coal is the most commonly used precursor for AC production 51 due to the advantage of its low cost of production and abundant resources as well as specific 52 53 beneficial properties [2]. Furthermore, coal-based activated carbon (CAC) is greater demand since it has greater density, corrosion endurance, hardness and more durable than other AC [3]. Coal is an 54 abundant resource with geological reserves exceeding 2.19 million tons. China accounts for 40% of 55 56 the total reserves. Xinjiang Province in Northwest China has become a large-scale coal production base. The high quality of coals is characterized as low ash yields, low S contents, and low mineral 57 contents [4]. Providing abundant and high quality raw materials is thus important for the 58 59 development of high value-added coal chemical products. For this reason, coal is selected as a stable alternative for the preparation of AC in Xinjiang. 60

Generally, two methods are used for the preparation of ACs via physical and chemical activation. During physical activation, the raw material is carbonized first at high temperature and then it is activated by CO_2 or steam under pressure to increase porosity and surface area of ACs. In chemical activation both carbonization and activation takes place simultaneously, in which raw material is first impregnated with activating chemical and then carbonized at desired temperature that varies according to activating chemical used [5]. Conventional heating method is usually transferred from

surface into the interior of the material by thermal conduction using the tubular furnace at relatively 67 high temperature and long time; as a result, the process of heating was slow, high energy 68 69 consumption and non-uniform with the surfaces [6]. In recent years, microwave irradiation has been widely investigated due to the tremendous thermal gradient from the interior of the char particle to 70 71 its cool surface allows the microwave-induced reaction to proceed more quickly and effectively at a lower bulk temperature [7]. Moreover, microwave radiation supply energy directly to carbon 72 skeleton at molecular level. Thus, microwave heating provides additional advantages such as energy 73 savings and shortening the processing time, possibility of selective heating, precise control of 74 75 temperature, small equipment size and reduced waste comparing with traditional heating techniques [8, 9, 10, 11, 12, 13]. 76

In microwave irradiation chemical activation, the raw material is first impregnated by activating 77 78 agents such as H₃PO₄, KOH or ZnCl₂, followed by thermal activation to create the pore structure. The microwave induced chemical activation usually has higher carbon yield and better developed 79 pore structure than the physical activation [14, 15, 16, 17]. Liu et al. [9] develop a microwave 80 induced activation process for the preparation of high surface area bamboo-based AC. The optimal 81 activation conditions were determined as: microwave power 350 W, radiation time 20 min and 82 phosphoric acid/carbon ratio 1:1, under which a surface area of 1432 m^2/g could be reached. Li *et al.* 83 [7] prepared relatively well developed porosity AC from tobacco stems by chemical activation with 84 K₂CO₃ using microwave radiation. The optimum conditions were as follow: microwave power 700 85 W, microwave radiation time 30 min, K₂CO₃/C ratio 1.5:1. Wang et al. [10] has been successfully 86 87 used to prepare AC from wood via microwave-induced ZnCl₂ activation. However, relatively few studies have focused on the preparation of CAC from anthracite by microwave activation. Revealing 88

the correlation between the microwave process conditions of preparing CAC and its adsorption
performance is also necessary.

91 Polycyclic aromatic hydrocarbons (PAHs) are a class of compounds that have more than two aromatic rings in their molecules. PAHs can transport long distance in air and water, and some PAHs 92 93 are capable of interacting with DNA to promote mutagenic and carcinogenic responses [18, 19]. Because of their stability and difficult degradation by microorganisms, it is difficult to remove PAHs 94 using only biological treatment processes. Additionally, PAHs are not efficiently removed by 95 conventional physicochemical methods such as coagulation, flocculation, sedimentation, filtration, 96 97 or ozonation. AC is the most commonly used porous media for the removal of PAHs from solution [20-22]. Yuan *et al.* [23] reported that porous carbons were prepared from petroleum coke by KOH 98 chemical activation and used as adsorbents for effective adsorbing naphthalene, fluorene, 99 100 phenanthrene, pyrene and fluoranthene from aqueous solutions, at an initial concentration of 3250–6250 ng/mL, over 90% of PAHs are removed from the solution within 30 min. Kong et al. [20] 101 reported that the performance of sesame stalk-based AC by potassium hydroxide in the efficient 102 103 removal of phenanthrene from an aqueous solution, at a carbon concentration of 10 mg/32mL and a carbonization temperature of 700 °C, the removal efficiency of phenanthrene approached 100%, 104 105 which was comparable to that by the commercial carbon. Cabal et al. [21] studied the preparation of AC from bean pods waste by chemical activation was investigated as adsorbents for the adsorption 106 of naphthalene from water solutions at low concentration. Moreover, a number of studies have 107 shown that AC prepared from different materials has significant differences in the adsorption of 108 PAHs. The adsorption behavior of different PAHs on AC has some differences [24, 25, 26]. These 109 three typical PAHs (naphthalene, phenanthrene, pyrene) were selected owing to their common 110

presence in the environment and their troublesome properties. These are taken as representatives of two-, three- and four-ringed PAHs. It can be seen that these substances have low solubility in water, which tends to be even lower when the number of aromatic rings is increased. Therefore, exploring CAC prepared using microwave activation for the adsorption features of typical PAHs in 30% ethanol aqueous solution is considerably interesting and informative.

The main purpose of the present study is to use rich anthracite from Xinjiang as raw material to 116 develop a microwave-induced activation process for the preparation of CAC with high surface area 117 118 and high adsorption performance to PAHs. The effects of activator types, coal alkali ratio (CAR), 119 activation time, and microwave power on the adsorption capabilities of CAC for MB and iodine were investigated. CAC was characterized by using the Boehm method, point of zero charge, 120 nitrogen adsorption isotherm, and scanning electron microscopy (SEM). The adsorption features, 121 isotherm, and kinetics of typical PAHs such as naphthalene, phenanthrene, and pyrene were also 122 explored. The results were used to evaluate the potential application of CAC prepared by 123 microwave-induced activation for the removal of PAHs. 124

125 **2. Materials and methods**

126 **2.1. Materials**

This raw anthracite coal was purchased from TBEA, Changji, Xinjiang Uygur Autonomous Region, China. The coal was passed through a 100 mesh sieve, washed with distilled water to remove carbon and fines, and dried in an air oven at 110 °C for 4 h. The resultant coal powder was used to prepare the CAC. The selected physio-chemical properties of naphthalene, phenanthrene and pyrene are shown in Table 1. Naphthalene, phenanthrene and pyrene (>98%; Aladdin, Shanghai, China) were used to prepare the stock solutions by dissolving an appropriate amount in 30% ethanol 133 aqueous solution.

134 **Table 1 should be positioned here.**

135 **2.2. Preparation of CAC**

CAC was prepared within a microwave oven (MM823LA6-NS, Midea) at a frequency of 2.45 136 137 GHz. The oven was equipped with a power controller (100, 300, 500, 700, and 900 W) and a time controller (4, 8, 12, 16, and 20 min). The mixture of 10 g with various ratios (1:1, 1:2, 1:3, 1:4, and 138 1:5) of the dried coal powder to activator (KOH, ZnCl₂ and H₃PO₄) was placed in a quartzose tube 139 of a microwave reactor to activate under vacuum atmosphere at the given power and time. The 140 141 resultant products were washed with 10% hydrochloric acid and then with distilled water until the filtrate reached neutral. The remaining solid was dried in an oven at 110 °C for 4 h and stored in 142 143 sealed glass bottles for the adsorption experiments.

The amount of MB adsorption is often regarded as a measure of its performance for high molar-mass species and was determined according to GB/T12496.10-1999 (Detection standard of activated carbon in China) [20]. On the other hand, the adsorption performance of activated carbons for low-molar mass species is generally indicated by the iodine number, which was determined at 303 K based on the Standard Detection Method (ASTM Designation: D4607-86) [20].

149 **2.3. Characterization methods**

The textural properties of the CAC were evaluated by measuring nitrogen adsorption isotherm at 77 K with ASAP 2020C surface area and porosity analyzer (Micromeritics Instrument Corp., USA). The S_{BET} was calculated using the BET equation. The total pore volume (V_t) was calculated by converting the adsorption amount at P/P₀ = 0.95 to a volume of liquid adsorbate. The micropore area (A_m) and volume (V_m) were obtained using the t-plot method. The external volume (V_e) was the deduction of V_m from V_t and the external area (A_a) was the deduction of A_m from S_{BET}. The surface morphology of the CAC was examined using scanning electron microscopy (SEM) (JSM-6490LV, Japan Electron Optics Laboratory Ltd. Corp., Japan) at accelerating voltages of 15 kV. Before observation, the samples were coated with gold in E-1010 Ion sputter. X-ray diffraction (XRD) was carried out on a Rigaku D/MAX-YA diffractometer with Ni-filtered Cu Ka radiation as the X-ray source.

The "Boehm titration" is a commonly used technique to determine the acidic oxygen surface 161 functional groups on carbon samples whereby bases of various strengths (NaHCO₃, Na₂CO₃, NaOH) 162 163 neutralize different acidic oxygen surface functionalities [19]. The weakest base, NaHCO₃, neutralizes only the strongest acidic carbon surface functionalities (CSFs) which are carboxylic 164 groups, while Na₂CO₃ neutralizes carboxylic and lactonic groups. The strongest base typically used, 165 166 NaOH, neutralizes carboxylic, lactonic and phenolic groups. The number of each type of CSF can be determined by difference between the uptake of each reaction base [19]. A mixture of 1.5 g CAC and 167 25.00 mL of one of the three 0.05 M reaction bases, NaHCO₃, Na₂CO₃ and NaOH was filtered and 168 wash with distilled water adequately after stirring for 24 h, then all the filtrate collected was added 169 170 0.05 M HCl to completely neutralize and end with methyl red indicator.

The point of zero charge (pH_{PZC}) is defined as the pH of the mixtures at which surface charge density on the adsorbent is zero. The pH_{PZC} values were determined by a mass titration method proposed by Franz [14]. Activated carbon (0.1 g) was added to 20 mL NaCl solutions, and then the initial pH had been adjusted with NaOH or HCl. The flasks were sealed and placed on a shaker for 24 h. The pH_{PZC} occurred when there was no change in the pH after contact with the carbon.

176 **2.4. Adsorption experiments of PAHs**

177 2.4.1. Adsorption performance analysis

Naphthalene, phenanthrene and pyrene were initially solved into ethanol aqueous solution (30 178 vol%) to prepare the 300 mg/L stock solutions, respectively. The concentrations of various adsorbate 179 solutions were measured after adsorption by a UV-vis spectrophotometer (UV-752N, Shanghai 180 181 Metash Instruments Co., Ltd, China). The concentration of naphthalene, phenanthrene and pyrene were calculated based on the following calibration equations, respectively: (1) at 218 nm, (2) at 249 182 nm, (3) at 237 nm. C = 1.7319A218 - 0.2309 (1), and R^2 was equal to 0.9995. $C = 3.4483A_{249} + 1000$ 183 0.0103 (2), and R^2 was equal to 0.9998. $C = 3.4483A_{237} - 0.0469$ (3), and R^2 was equal to 0.9998. 184 For each adsorption kinetics experiment, 40 mL of 300 mg/L adsorbate solution was placed in a 185 100 mL conical flask with a cover. The solution was shaken at 25 °C and 60 rpm. After preheating 186 for 5 min, 100 mg of CAC prepared was added to the solution. Then, the samples were taken 187 188 periodically and filtrated immediately by suction. The supernatant solution was analyzed for the remaining concentration of adsorbate by a UV-vis spectrophotometer. 189 For each adsorption isotherm experiment, approximately 40 mL of different adsorbate solution of 190

various concentrations was placed in a 100 mL conical flask with cover. To establish the thermal equilibrium before adsorption, the solution was shaken at 25 °C and a stirring rate of 60 rpm. After preheating for 5 min, 150 mg of CAC prepared was added to the solution. Then, the samples were taken after 50 min and filtrated immediately by suction. The supernatant solution was analyzed for the remaining concentration of adsorbate by a UV–vis spectrophotometer. The amount of adsorbate (Q_e) was calculated as follow:

197
$$Q_e = \frac{(C_0 - C_e)V}{m} \tag{1}$$

198 where C_0 (mg/L) and C_e (mg/L) are liquid-phase concentrations of adsorbate at initial and

equilibrium, respectively. V (mL) represents the volume of the solution and m (g) stands for the massof CAC used.

201 2.4.2. Adsorption isotherm

The Langmuir adsorption Eq. (2), predicts the existence of monolayer coverage of the adsorbate at the outer surface of the adsorbent [9].

204
$$\frac{C_{\rm e}}{Q_{\rm e}} = \frac{1}{q_{\rm m}K_{\rm L}} + \frac{C_{\rm e}}{q_{\rm m}}$$
(2)

where $C_{\rm e}$ and $Q_{\rm e}$ are as defined in Eq. (1), $q_{\rm m}$ is adsorption maximum (mg/g); $K_{\rm L}$ is sorption equilibrium constant (L/mg).

The Freundlich isotherm [9] is an empirical equation employed to describe heterogeneous systems, and is expressed by the following Eq. (3):

209
$$1 \circ \mathcal{Q}_{e} = \frac{1}{n} 1 \circ \mathcal{C}_{e} + 1 \circ \mathcal{K}_{F} \quad (3)$$

where $K_{\rm F}$ is a constant which represents a measure of the adsorption capacity of the adsorbent for specific solute, and *n* is a measure of intensity of adsorption.

212 2.4.3. Adsorption kinetics

217

Kinetic models are used to examine the rate of the adsorption process and potential rate controlling step, i.e., particle diffusion or chemical reaction. The capability of pseudo-first-order, pseudo-second-order and intraparticle diffusion, were examined in this study.

The pseudo-first-order kinetic rate equation is expressed as [15]:

$$\frac{1}{Q_{\rm t}} = \frac{1}{Q_{\rm e}} + \frac{k_{\rm 1}}{Q_{\rm e}\,{\rm t}}$$
(4)

where k_1 is the rate constant of pseudo-first-order sorption (min), which can be determined from the slope of the linearized pseudo-first-order rate equation. 220 The pseudo-second-order kinetic rate equation is expressed as [15]:

221
$$\frac{t}{Q_{\rm t}} = \frac{1}{k_2 Q_{\rm e}^2} + \frac{t}{Q_{\rm e}}$$
(5)

where k_2 is the second-order rate constant (g/mg/min) can be determined from the intercept of the linearized pseudo-second-order rate equation.

In the intraparticle diffusion model, the relationship between the adsorption capacity at time *t*, Q_t and $t^{1/2}$ could be written as [27]:

226
$$Q_t = K_p t^{1/2} + C$$
 (6)

where K_p is the intraparticle diffusion constant (mg/g/min^{1/2}) and *C* is the intercept of the line, which

is proportional to the boundary layer thickness.

3. Results and discussion

230 **3.1 Preparation of CAC**

231 3.1.1. Effects of activator

MB molecule has been estimated that the minimum pore size it can enter is 1.3 nm [28] and can be used to indicate the capacity of a carbon for sorption of medium-sized molecules. The iodine molecule provides information about the surface area that is contributed by pores larger than 1 nm [29], and the iodine number provides a good indication of the capacity of a carbon for sorption of small molecules [30]. The average pore width of KOH-activated CAC was approximately 2.8 nm in Table 2, which was a sufficiently large pore width to allow MB and iodine molecules to pass through for a large amount of adsorption.

According to the preliminary experiment, the ratio for $ZnCl_2$ or H_3PO_4 was also 1:3, whose is optimum conditions are the same as KOH. The influences of activator types on the adsorption amounts of MB and iodine on CAC prepared at a coal activator ratio of 1:3, activation time of 12 min, and microwave power of 700 W are shown in Fig. 1. In the chemical activation, the adsorption property of CAC prepared with various activating agents showed a significant difference. The adsorption amounts of MB and iodine on CAC prepared with KOH reached the maximum. Alkaline hydroxide activation involves the redox reduction and carbon oxidation to generate porosity. During the reactions, the evolution of CO, CO₂ and H₂ constituents, and additional reactions [11] between the active intermediates with the carbon surface are possible (Eqs. (1) - (3)).

- $6KOH + C \rightarrow 2K + 3H_2 + 2K_2CO_3 \tag{1}$
- 249

$$K_2CO_3 + C \rightarrow K_2O + 2CO_2 \tag{2}$$

(3)

 $250 \qquad \qquad 2K + CO_2 \rightarrow K_2O + 2CO$

The produced alkaline is intercalated to the carbon matrix responsible for both stabilization and 251 widening of the spaces between the carbon atomic layers. Thus, KOH activated samples exhibited 252 253 the best adsorption capacity. Metallic potassium formed during the redox reaction can be intercalated into the carbon walls independently of the structural order responsible for separation and degradation 254 of graphitic layers thus develop the microporosity and mesoporosity. The resulting CAC by the 255 activation agents of ZnCl₂ and H₃PO₄ had very low adsorption amounts of MB and iodine 256 (particularly the adsorption amounts of MB), which may be due to the activation of ZnCl₂ and 257 H₃PO₄ producing a lower specific surface area and smaller pore size on CAC, which is showed in 258 Table 2. The activation agents of ZnCl₂ and H₃PO₄ could be widely used to prepare AC from 259 lignocellulosic products [31]. So KOH was selected as appropriate activator in this investigation. 260

- 261 **Table 2 should be positioned here.**
- 262 Fig. 1 should be positioned here.
- 263 3.1.2. Effects of coal to KOH ratio (CAR)

Table 3 presents the influences of the CAR on the adsorption amounts of MB and iodine on CAC prepared at microwave power of 700 W for 8 min. The activation reaction was strengthened and a large number of pores were formed with the rising ratio, enhancing the CAC adsorption capacity. When CAR reached 1:3, the carbons on the active sites could be reacted completely, and the adsorption capacity reached the maximum [7]. When CAR was larger than 1:3, the pores were widened. Besides, excessive KOH molecules might decompose following the reactions:

$$270 \qquad 2KOH \rightarrow K_2O + H_2O \qquad (4)$$

$$H_2O + C \rightarrow CO + H_2 \tag{5}$$

Therefore, the catalytic oxidation entailed the widening of mesoporous structures and carbon burn off, so the adsorption capacity of AC was decreased.

Surface functional groups such as phenolic group (-OH), lactonic group (-COOCO-), and 274 275 carboxylic group (-COOH) were determined by Boehm titration. The results of these surface functional groups for CAC with various CARs are shown in Table 3. The CAR apparently had a 276 destructive influence on the functional groups of CAC. With increased CAR, the amount of lactonic 277 278 groups, phenolic and carboxylic groups decreased, and phenolic and carboxylic groups are fairly low. A possible explanation for this could be that the alkali activator affected the number of surface acidic 279 functional groups. The results of pH_{PZC} titrations for CAC under various CARs can also be seen in 280 Table 3. pH_{PZC} increased with increased CAR. Previous research has indicated that the existence of 281 surface functional groups, such as carboxyl, carbonyl, and phenol, constitutes the source of surface 282 acidity [9, 12, 32] and the opposite is observed for alkali treatment. Considering the results of the 283 adsorption capacity of CAC, CAR was set at 1:3 for this study. 284

285 3.1.3. Effects of activation time

Table 3 demonstrates the effects of activation time on the adsorption amounts of MB and iodine 286 on CAC prepared with CAR of 1:3 at microwave power of 700 W. The MB adsorption amounts of 287 288 CAC increased from approximately 234 to 312 mg/g with increased activation time from 4 to 12 min, and then decreased to 238.5 mg/g for 20 min activation. Similarly, the iodine number of the CAC 289 290 increased gradually from approximately 962.12 to 1048.19 mg/g with increased time from 4 to 12 min, and then slightly decreased to 1045.87 mg/g for 20 min activation. However, when the 291 activation time reached a certain value, the carbon pores were burnt off by microwave heating, 292 thereby lowering the CAC adsorption capacity. A similar tendency was observed in the preparation 293 294 of AC from rice husk [17].

The results of the surface functional content and pH_{PZC} titrations for CAC under various activation times can be seen in Table 3. With increased activation time from 4 to 20 min, the number of surface functional groups decreased. pH_{PZC} generally showed the opposite trend. The phenomenon implied that increasing activation time promoted an acceleration of energy, leading to a destruct of surface functional groups and developed the rudimentary of the pore structure [11]. Considering the results of the CAC adsorption capacity, the activation time was set at 12 min in this study.

302 3.1.4. Effects of microwave power

The effects of microwave power on the adsorption amounts of MB and iodine of CAC prepared by KOH activator with CAR of 1:3 for 12 min, as shown in Table 3. The adsorption amounts of MB rose from 54 to 312 mg/g with increased power from 100 to 700 W and then decreased. The adsorption amounts of iodine increased from 769.02 to 1048.19 mg/g with increased power from 100 to 700 W and then decreased. More carbon molecules were activated along with the higher 308 microwave power participate in the activation reaction, and lead to the development of pore structure 309 adequately. However, the exceeded of a certain power may lead to excessive destruction of pore 310 structure [11].

The results of the surface functional content and pH_{PZC} titrations for CAC under various microwave powers can be seen in Table 3. With increased microwave power from 100 to 900 W, the number of lactonic groups decreased. The surface functional content and pH_{PZC} had the opposite trend. Menéndez *et al.* [12] pointed out that microwave treatment reduced the functional group, especially those of acidic in nature, and increased the pH_{PZC} of the carbon. In this study, CAC produced directly by microwave heating agreed with the reported findings [12]. The optimum microwave power in this experiment was thus 700 W.

318 **Table 3 should be positioned here.**

319 **3.2. Characterization of CAC**

320 3.2.1. Porous structure analysis

The N_2 adsorption isotherms of the CAC at 77 K are presented in Fig. 2. The adsorption isotherms demonstrated a sharp rise at low P/P₀ range, and a gradual increase was seen at relatively high P/P₀ range. According to the classification of IUPAC [33], all adsorption isotherms were of type I, indicating that the CACs were mainly microporous. The N_2 adsorption isotherms on activated carbons from Zonguldak region coals using chemical activation also conformed to type I isotherm by Kopac and Toprak [3]. Moreover, the saturated adsorption amounts of N_2 increased after microwave activation, implying an increase in the pore volume.

The textural properties obtained from the N₂ adsorption isotherms are summarized in Table 2. CAC-4 had a well-developed porosity with a surface area of 1770.49 m^2/g , which was primarily

microporous under optimal conditions. Liu et al. [9] reported that bamboo-based AC using 330 microwave radiation has a maximum surface area of 1409 m²/g. Yagmur *et al.* [13] reported that the 331 maximum BET surface area was 1157 m²/g for the AC from waste tea treated with chemical 332 activation with microwave energy. Compared with previous works, CAC obtained a higher specific 333 334 surface area. This observation suggested that coal could also form a considerable porous structure through microwave-assisted chemical activation. The specific surface area obtained by the BET 335 equation (S_{BET}), micropore area (A_m), external surface area (A_e), micropore volume (V_m), and total 336 pore volume (Vt) of CAC-2 were larger than those of CAC-1, whereas the average pore width of 337 338 CAC-2 was smaller than CAC-1. This result indicated that the increasing potassium hydroxide amount enhanced the pore development and created new pores of coal, resulting in further increased 339 specific surface area and decreased average pore width. CAC-2, CAC-3, and CAC-4 were obtained 340 341 using different activation times. With increased activation time from 4 min to 12 min, the SBET, Ae, and Vt gradually increased. The Am and Vm of CAC-2 and CAC-4 had no significant differences, and 342 they were larger than those of CAC-3 but not significantly. The phenomenon implied that the 343 344 increase in activation time promotes an acceleration of energy, which in turn increases the reaction rates, thus developed the rudimentary of the pore structure [11]. The six surface physical parameter 345 values of CAC-5 prepared at 100 W of microwave power were small, indicating no strong reaction 346 occurred between the coal and activation agent at low microwave power. 347

348 Fig. 2 should be positioned here.

Enhancing microwave power from 100 to 700 W drastically improved textural properties, possibly ascribed to the combined effect of internal and volumetric heating responsible for the expansion of carbon structure [11]. Also, considerable changes in the surface properties were

achieved within a short time, which should be attributed to the distinct mechanism of microwave 352 heating. As listed in Table 4, the samples obtained by microwave heating have higher BET surface 353 354 area. It can be seen advantage of microwave heating is that the treatment time as well as the consumption of gases used in the treatment can be considerably reduced, which resulted in a 355 356 reduction in the energy consumption, compared with conventional surface heating. In addition, microwave heating technique has overall shown better effect in terms of porous structure, relatively 357 greater surface area than conventional heating method [3]. Hence, the work reported here clearly 358 demonstrates that the use of microwave heating for preparation CAC is more rapid, cheaper and 359 360 efficient than the use of a conventional furnace.

361 **Table 4 should be positioned here.**

362 3.2.2. Surface morphology analysis

SEM images of the coal and CAC prepared under optimum conditions (CAC-4) are shown in Fig. 3. The surface of raw material (Fig. 3(A)) was fairly smooth without any pores except for some occasional cracks. After activation, it shows the obtained CAC has an irregular surface with pores and craters. From Table 2, it was also known that the pores are micro- and mesopores. As shown in Fig. 3(B), the external surface of the CAC-4 demonstrated a well pronounced pore structure, indicating constitution of new matrix during the activation stage.

- 369 Fig. 3 should be positioned here.
- 370 3.2.3. Crystal structure

X-ray diffraction technique is a powerful tool to analyze crystalline nature of materials. Fig. 4
 shows the XRD profiles of coal and CAC-4. The coal shows an intense diffraction peaks at 26°. The
 CAC-4 prepared from coal via microwave-induced KOH activation shows an intense diffraction

- peaks at around 29° and several not obvious diffraction peaks at around 44° , corresponding to the (002) and (100) diffraction of disordered stacking of micrographites, respectively [14].
- 376 Fig. 4 should be positioned here.

377 3.3. Adsorption isotherm of selected PAHs on CAC-4

378 Table 5 lists the parameters of the Langmuir and Freundlich adsorption isotherm models of naphthalene, phenanthrene, and pyrene in ethanol aqueous solution onto CAC-4 at 25 °C, along with 379 their regression coefficients (R^2), the adsorption capacity and sorption equilibrium constants (q_m and 380 $K_{\rm L}$), and intensity parameters ($K_{\rm F}$ and n). R^2 reflected the goodness of fit of each of the models to the 381 data. For the Langmuir isotherm, the R^2 values were >0.9934. The regression coefficients R^2 of the 382 Freundlich isotherm were >0.9809. Both Langmuir and Freundlich isotherm models efficiently 383 predicted the adsorption of the three PAHs on CAC-4 with high correlation coefficients. This 384 385 observation indicated that the sorption of naphthalene, phenanthrene, and pyrene onto CAC-4 may be confined to a monolayer adsorption and heterogeneous sorption. Similarly, adsorption isotherm of 386 PAHs from vegetable oil on AC also fitted well by both isotherm models [26]. While Yuan et al. [22] 387 reported that the equilibrium adsorption isotherms of PAHs from water on petroleum coke-derived 388 porous carbon fitted the Freundlich equation well, and Kong et al. [20] reported that the 389 phenanthrene removal from aqueous solution on sesame stalk-based carbon fit the Freundlich 390 equation well. 391

With carbon, the sorption process results from molecular interactions between the carbon surface and the adsorbate, such as van der Waals attractions, electrostatic forces, and weak intermolecular association [34]. The properties of the adsorbate greatly influence the sorption process. Since PAHs are nonpolar compounds, sorption is governed mainly by hydrophobic interactions [24]. The values of $K_{\rm L}$, $K_{\rm F}$, and $q_{\rm m}$ followed the trend of pyrene > phenanthrene > naphthalene, which confirmed that hydrophobic interactions follow the same trend. Kong *et al.* [20] reported that the hydrophobic interactions were the key factor for phenanthrene removal from aqueous solution by the sesame stalk-based carbon. Moreover, the values of *n* were all greater than 1 and did not significantly change, showing that the prepared CAC favorably adsorbed the three PAHs [22].

401 **Table 5 should be positioned here.**

402 **3.4. Adsorption kinetics of selected PAHs on CAC-4**

Fig. 5 shows the plots of the adsorption amounts of naphthalene, phenanthrene, and pyrene at 403 404 different time intervals. The adsorption process consisted of two distinguished periods. In the first stage, the three PAHs were rapidly adsorbed onto easily accessible hydrophobic sites in the CAC-4 405 406 matrix within 5 min. In a well-stirred, solid-liquid system, the first stage of diffusive transport of 407 PAHs molecules from the bulk solution to the exterior surface of porous CAC particles, as well as the intraparticle diffusion of the PAHs through the interior pores of CAC, were assumed to be rapid. 408 409 In the second stage, adsorption was probably limited by the slow migration of adsorbates to less 410 accessible sites associated with micropores within the CAC-4 matrix, which could take a long time. Valderrama et al. [25] reported similarly that the adsorption process of PAHs onto granular AC also 411 412 was the two-stage. The adsorption amount increased with increased contact time. The maximum adsorption was observed at 20 min, beyond which almost no further increase in the adsorption 413 amount was observed. 414

The experimental data for the naphthalene, phenanthrene, and pyrene adsorption onto CAC-4 were fitted by three kinetic models, the values of k_1 , k_2 , k_p , and Q_e , and the correlation coefficients are listed in Table 6. The experimental kinetic data followed the pseudo-second-order kinetic curves

for the three PAHs, which can be demonstrated by higher R^2 (>0.9999) and closer Q_e values to the 418 experimental ones, q_{exp} . The adsorption progress of the PAHs on the CAC surface were presumably 419 420 caused by the chemical interaction [14]. Cabal *et al.* [21] found the experimental data of adsorption kinetics of aqueous naphthalene on the AC were fitted to the pseudo-second kinetic model with a 421 422 high correlation coefficient. While a literature showed that sorption systems of naphthalene and pyrene on AC followed a pseudo-first-order reaction model, although the pseudo-second-order 423 reaction model provides an acceptable description of the sorption process [25]. The equilibrium 424 adsorption amounts at 20 min were 117.17 mg/g for pyrene, 111.39 mg/g for phenanthrene, and 425 426 78.18 mg/g for naphthalene, respectively, decreased as follows: pyrene > phenanthrene > naphthalene. Compared with the carbonaceous materials such as the phenanthrene adsorption 427 amount at 24 h from aqueous solution of sesame stalk- based AC was 73.74 mg/g [20], the 428 429 naphthalene adsorption amount at 60 min from aqueous solution of reed straw- based AC was 107.00 mg/g [35], and the pyrene adsorption amount at 24 h from aqueous solution of rice husk- based AC 430 was 104.50 mg/g [36], the PAHs adsorption performance of prepared carbon in this work was higher 431 432 and faster. In addition, coal as a precursor is relatively cheaper. Thus, these results indicate that CAC 433 prepared with microwave activation had excellent capability to remove PAHs from aqueous solution. The agreement between the equilibrium adsorption values determined (q_{exp}) and the adsorption 434 maximum (q_m) obtained from the Langmuir isotherm was clear and sufficient. The observed trend in 435 436 adsorption amounts was generally consistent with the trend in the amount of aromatic rings and molecular weight shown in Table 2. The adsorption amounts were higher for larger molecular PAHs 437 438 such as pyrene, which suggested that the PAHs were able to form a π - π complex between the π -electrons of benzene rings and active sites on the AC surface [18]. Therefore, the π - π dispersive 439

interactions between the aromatic ring in PAHs and the graphene layers on CAC were the key 440 parameters that determined the adsorptive affinity of those molecules with a polycyclic aromatic 441 442 skeleton structure [19]. This phenomenon can explain why molecules with more aromatic rings showed a significantly higher adsorptive affinity than aromatics with fewer rings. The electron 443 444 donor-acceptor mechanism could involve lactonic groups that predominated on the basic carbon surface. Lactonic groups acted as electron donors. The aromatic rings in the adsorbate accepted 445 electrons [21], consistent with the number of lactonic groups and pH_{PZC} CAC-4 prepared at 0.35 446 mmol/g and 8.35, respectively. 447

- 448 Fig. 5 should be positioned here.
- 449 **Table 6 should be positioned here.**
- 450 **4. Conclusions**
- The optimum conditions for CAC preparation were identified to be KOH activator, CAR of 1:3,
 activation time of 12 min, and microwave power of 700 W. Under the optimum conditions
 adsorption amounts of MB and iodine on CAC was 312 and 1048.19 mg/g, respectively.
- 454 2. The CAR, activation time, and microwave power destructed surface functional groups and 455 raised the pH_{PZC} of CAC. The BET surface area and pore volume of the optimal CAC were 456 evaluated as 1770.49 m²/g and 0.99 cm³/g, which was primarily microporous. The SEM image 457 of the optimal CAC showed that the external surface of CAC had a well-pronounced and almost 458 uniform pore structure. Hence, the work clearly demonstrates that the use of microwave heating 459 for preparing CAC is more rapid, cheaper and efficient.
- 3. The adsorption equilibrium of naphthalene, phenanthrene, and pyrene on CAC-4 wassatisfactorily represented by both the Langmuir and Freundlich isotherm models with high

correlation coefficients. The prepared CAC favorably adsorbed naphthalene, phenanthrene, and 462 pyrene. The experimental kinetic data followed the pseudo-second-order kinetic curves for 463 naphthalene, phenanthrene, and pyrene, presuming that the adsorption progress is affected by 464 chemical interactions. The equilibrium adsorption amounts were 117.17 mg/g for pyrene, 111.39 465 mg/g for phenanthrene, and 78.18 mg/g for naphthalene. Overall, CAC from Xinjiang region 466 coal prepared by microwave activation could be effectively applied in liquid-phase adsorption 467 involving PAHs. 468 Acknowledgements 469 470 This work was supported financially by funding from the National Natural Science Foundation

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473 **References**

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575 Figure captions:

- 576 Fig. 1 Adsorption amounts of MB and iodine by CAC with various activator types (preparation
- 577 conditions: activation time = 12 min; microwave power = 700 W; CAR= 1:3).
- 578 Fig. 2 Nitrogen adsorption isotherm for the prepared CACs (CAC-1: activation time = 8 min;
- 579 microwave power = 700 W; CAR= 1:1, CAC-2: activation time = 8 min; microwave power = 700 W;
- 580 CAR= 1:3, CAC-3: activation time = 4 min; microwave power = 700 W; CAR= 1:3, CAC-4:
- activation time = 12 min; microwave power = 700 W; CAR= 1:3, CAC-5: activation time = 12 min;
- 582 microwave power = 100 W; CAR= 1:3).
- 583 Fig. 3 SEM micrographs of the selected samples at 5000× magnification: (A) coal, (B) CAC-4
- 584 (preparation conditions: activation time = 12 min; microwave power = 700 W; CAR= 1:3).
- 585 Fig. 4 XRD profiles of coal and CAC-4 (preparation conditions: activation time = 12 min;
- 586 microwave power = 700 W; CAR= 1:3).
- 587 Fig. 5 Adsorption amounts of naphthalene, phenanthrene and pyrene in ethanol aqueous solution on
- 588 CAC-4 (preparation conditions: activation time = 12 min; microwave power = 700 W; CAR= 1:3) at 589 various times and at 25 °C.
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Fig. 1 Adsorption amounts of MB and iodine by CAC with various activator types (preparation conditions: activation time = 12 min; microwave power = 700 W; CAR= 1:3).





⁶⁰⁵ Fig. 2 Nitrogen adsorption isotherm for the prepared CACs (CAC-1: activation time = 8 min;

606 microwave power = 700 W; CAR= 1:1, CAC-2: activation time = 8 min; microwave power = 700 W;

607 CAR= 1:3, CAC-3: activation time = 4 min; microwave power = 700 W; CAR= 1:3, CAC-4:

activation time = 12 min; microwave power = 700 W; CAR= 1:3, CAC-5: activation time = 12 min;

609 microwave power = 100 W; CAR= 1:3).

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Fig. 3 SEM micrographs of the selected samples at 5000× magnification: (A) coal, (B) CAC-4
(preparation conditions: activation time = 12 min; microwave power = 700 W; CAR= 1:3).



Fig. 4 XRD profiles of coal and CAC-4 (preparation conditions: activation time = 12 min;

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617 microwave power = 700 \text{ W}; CAR= 1:3).
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Fig. 5 Adsorption amounts of naphthalene, phenanthrene and pyrene in ethanol aqueous solution on CAC-4 (preparation conditions: activation time = 12 min; microwave power = 700 W; CAR= 1:3) at various times and at 25 °C.

			Molecular	Molecular	Malar	Water	logKow
PAHs	Structure	Formula	weight	dimension	volume	solubility	
			(g/mol)	(nm×nm×nm)	(cm ³ /mol)	(ng/g)	
Naphthalene		$C_{10}H_8$	128.17	0.91×0.73×0.38	148	30800	3.30
Phenanthrene	, A	$C_{14}H_{10}$	178.22	1.17×0.80×0.34	199	1283	4.57
Pyrene	()	C ₁₆ H ₁₀	202.25	0.92	214	135	5.18
			202.20			100	
Table 2 Activati	on conditior	is and sur	face chara	cteristics of the r	orepared CA	ACs.	
ample Activ	ation condit	ions	SBET	$A_m = A_e$	Vm	V _t V _m /	V _t Average

Table 1 Selected properties of PAHs used [22]

										width
	Microwave	e Microwave	Coal	m²/g	M^2/g	M^2/g	cm ³ /g	cm ³ /g	%	nm
	time	power	activator							
	min	W	ratio							
CAC-1	8	700	1:1	623.84	417.31	206.53	0.23	0.32	71.88	3.25
CAC-2	8	700	1:3	1651.50	543.09	1108.40	0.33	0.91	36.26	2.86
CAC-3	4	700	1:3	1007.27	379.35	627.92	0.22	0.54	40.74	2.86
CAC-4	12	700	1:3	1770.49	508.87	1261.62	0.61	0.99	61.62	2.82
CAC-5	12	100	1:3	0.21	0.00	0.00	0.00	0.00	0.00	0.00
CAC-6	12	700	1:3	934.54	425.64	534.22	0.37	0.49	75.51	1.95
CAC-7	12	700	1:3	843.56	384.34	423.55	0.31	0.42	73.81	1.88

 S_{BET} : specific surface area obtained by BET equation, A_m , A_e : micropore area, external surface area, V_m , V_t : micropore volume, total pore volume.

636 Activator: The CAC-1, CAC-2, CAC-3, CAC-4, CAC-4 and CAC-5 with KOH; the CAC-6 with 637 $ZnCl_2$; the CAC-7 with H₃PO₄.

Table 3 Adsorption performance and surface properties of CAC prepared with various CAR,
activation time and microwave power.

Treatments		MB	Iodine	Lactonic	Carboxylic	Phenolic	pH _{PZC}
		adsorption	adsorption	group	groups	groups	
		amount	amount	(mmol/g)	(mmol/g)	(mmol/g)	
		(mg/g)	(mg/g)				
	1:1	77.7	999.38	0.5667	0.0333	0.0812	8.10
	1:2	268.5	1034.24	0.4933	0.0167	0.0530	8.20
CAR	1:3	291.0	1038.90	0.4700	0.0063	0.0233	8.30
	1:4	270.0	1034.23	0.3500	0.0056	0.0100	8.35
	1:5	256.5	1017.95	0.2333	0.0028	0.0067	8.30
	4	234.0	962.12	0.5267	0.0900	0.0267	8.10
Activation	8	282.0	1033.54	0.4700	0.0733	0.0190	8.30
time	12	312.0	1048.19	0.3500	0.0700	0.0058	8.35
(min)	16	238.5	1045.87	0.2267	0.0333	0.0047	8.32
	20	241.5	1045.87	0.1833	0.0083	0.0033	8.31
	100	54.0	769.02	0.5733	0.0933	0.0562	7.50
Microwave	300	180.0	1003.99	0.4567	0.0267	0.0333	7.60
power	500	210.0	1031.91	0.4133	0.0133	0.0193	7.90
(W)	700	312.0	1048.19	0.3500	0.0093	0.0097	8.35
	900	219.0	1043.54	0.3333	0.0045	0.0033	8.30

Sample	Carbon	Heating	$\mathbf{S}_{\mathrm{BET}}$	Vtotal (cm ³ /g)	Data sources
	source	methods	(m ² /g)		
Commercial AC	Pitch	Convention	1403.00	0.583	[19]
Commercial AC	Wood	Convention	1100.00	1.061	[19]
Lotus stalk based - AC	Lotus stalk	Convention	1220.00	1.191	[14]
Jatropha hull based - AC	Jatropha hull	Convention	748.00	0.580	[2]
CAC	Coal	Convention	830.5m ² /g	0.250	[3]
Rice husk based - AC	Rice husk	Microwave	752.00	0.640	[37]
Cotton stalk based - AC	Cotton stalk	Microwave	729.33.00	0.380	[38]
CAC	Coal	Microwave	1770.49	0.990	This work
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Table 4 Comparison of different kinds of ACs.

PAHs		Langmuir		Freundlich		
	$K_{\rm L}({\rm L/mg})$	$q_{\rm m}({\rm mg/g})$	R^2	$K_{\rm F}({\rm L/mg})$	n	R^2
Naphthalene	0.03	89.29	0.9934	5.50	1.71	0.9921
Phenanthrene	0.28	116.28	116.28 0.9990		1.60	0.9848
Pyrene	0.90	117.65	0.9991	52.16	1.61	0.9809

Table 5 Isotherm constants for the adsorption of naphthalene, phenanthrene, and pyrene in ethanol aqueous solution onto CAC-4 at 25 °C.

Table 6 Kinetics constants for the adsorption of naphthalene, phenanthrene, and pyrene in ethanolaqueous solution onto CAC-4.

	$q_{ m exp}$	Pseudo-	first-orde	r kinetic	Pseudo-sec	ond-order	Intraparticle diffusion		
			model		:	model		model	
PAHs	(mg/g)	k_1	$Q_{ m e}$	R^2	k_2	$Q_{ m e}$	R^2	k_p	R^2
		(min ⁻¹)	(mg/g)		(g/mg min)	(mg/g)		(mg/g/ min ^{1/2})	
Naphthalene	78.18	0.61	80.65	0.9615	0.02	81.30	0.9999	1.84	0.9125
Phenanthrene	111.39	0.25	112.36	0.9970	0.04	112.36	1.0000	0.76	0.9124
Pyrene	117.17	0.13	117.65	0.9644	0.06	117.65	1.0000	0.61	0.9049