

Activated carbon obtained from sapelli wood sawdust by microwave heating for *o*-cresol adsorption

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Received: 29 April 2016/Accepted: 28 July 2016/Published online: 6 August 2016 © Springer Science+Business Media Dordrecht 2016

Abstract Activated carbon (AC) was prepared from sapelli wood sawdust using a microwave heating process. The biomass was mixed with inorganic components (lime + ZnCl₂ and FeCl₃) to form a homogeneous paste. The AC samples are denoted as AC-1A (100 g sapelli wood sawdust + 20 g lime + 80 g ZnCl₂), AC-2A (150 g sapelli wood sawdust + 20 g lime + 80 g ZnCl₂), AC-1B (100 g sapelli wood sawdust + 20 g lime + 40 g ZnCl₂ + 40 g FeCl₃), and AC-2B (150 g sapelli wood sawdust + 20 g lime + 40 g ZnCl₂ + 40 g FeCl₃). The samples were placed in a microwave oven and pyrolyzed under nitrogen flow. To increase their porosity, the pyrolyzed samples were subjected to a leaching process (with 6 mol L⁻¹ HCl) under reflux to eliminate inorganic components. Several analytical techniques such as Fourier-transform infrared (FTIR) spectroscopy, scanning electron microscopy (SEM), and N₂ isotherm and vapor adsorption analyses were performed to characterize the AC materials. The samples presented high Brunauer–

Electronic supplementary material The online version of this article (doi:10.1007/s11164-016-2683-8) contains supplementary material, which is available to authorized users.

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Emmett–Teller (BET) surface areas, up to 941.08 m² g⁻¹ for AC-1A. The AC materials were tested for their *o*-cresol removal ability by determining the best fits to equilibrium and kinetic data using the Sips isotherm and fractional-order model, respectively. The maximum adsorption capacity of the AC samples as obtained from the Sips model was correlated with the surface area. The proposed adsorption mechanism suggests that hydrogen bonding, donor–acceptor complexation, and π – π interactions play key roles. The adsorbents were also tested for treatment of simulated industrial effluents, showing very good efficiency. Almost complete regeneration of the AC adsorbents was achieved using 10 % EtOH + 5 mol L⁻¹ NaOH as eluent. These results demonstrate that sapelli wood sawdust is a promising precursor for preparation of AC to remove *o*-cresol from aqueous solution.

Keywords Activated carbon · Microwave-assisted pyrolysis · *o*-Cresol · Kinetic models

Introduction

Phenols are pollutants whose presence at even low concentration can seriously impact on water use and reuse. Phenols are commonly found in wastewater generated from petrochemical, steel, and phenol-producing industries as well as coal conversion [1].

Many phenols are hazardous contaminants because of their toxicity to the environment and living beings [2, 3]. Ingestion of water contaminated with high phenol content can cause serious gastrointestinal damage, muscle tremors, difficulty in walking, and death in animals [1, 3]. Therefore, Environmental Protection Agency (EPA) regulations try to lower the phenol content in wastewater to below $1 \ \mu g \ mL^{-1}$ [4]. Because of this high toxicity, effluents containing phenols must be treated before being disposal in the environment.

Additionally, it is well known that conventional wastewater and drinking-water treatment processes are not very efficient for removal of such phenolic compound pollutants. Therefore, study of effective methods for removing phenols from water and/or wastewater has become urgent.

There are many techniques for treatment of wastewater contaminated with phenolic compounds, including biological processes [5, 6], coagulation [7], membranes [8], electro-Fenton and photoelectro-Fenton treatments [9], ozonation [10], oxidation [11, 12], and adsorption [13–16]. However, application of these methods is restricted since they are very expensive and involve high initial capital and operational costs. On the other hand, adsorption is a more attractive method because pollutants can be removed from aqueous effluent while the adsorbent can be reused several times, making the wastewater treatment process economically feasible.

The most well-known and commonly used adsorbent is activated carbon (AC), especially for removal of organic hazards [16-18], due to its good textural features such as high surface area and pore volume [16-18].

In recent years, researchers have studied various different kinds of precursor for preparation of activated carbon, including sewage sludge [17, 18], agroindustrial waste [16, 19], tire waste [20], cotton seed cake [21], biomass from wood [22], etc.

Chips and sawdust are waste biomass from wood processing. A study in 28 sawmills in the City of Ngaoundere, Cameroon revealed very high use of lignocellulosic wood species (3.120 tons per year) such as ayous and sapelli. Transformation of these species generates tons of waste per year [23]. Therefore, any attempt to reuse this waste would be useful for the country [24].

Preparation of activated carbon can be accomplished using a conventional furnace or a microwave oven [17, 25]. The main difference between these two types of pyrolysis is the way in which the heat is generated. Transfer of heat occurs by conduction in a conventional oven, unlike in microwave heating, where the energy is furnished directly to the carbon bed, being transformed into heat inside the sample by ionic conduction and dipole rotation through friction within the material [17, 25, 26]. Microwave heating is advantageous because of its shorter pyrolysis time (<10 min) [27] for activation as a result of the rapid temperature rise, as well as its remarkably lower energy consumption [25–27].

The aim of this work is to produce different AC materials using microwaveassisted pyrolysis and test these adsorbents for *o*-cresol removal from aqueous solution. The effect of different operational parameters, such as temperature and the initial pH value of the *o*-cresol solution, on the batch adsorption system was verified. Several analytical techniques were used to explore the nature of the AC samples obtained with regard to their structure and surface texture.

Experimental

Chemicals and reagents

o-Cresol (see Supplementary Fig. 1) was supplied by Vetec (São Paulo, Brazil). $ZnCl_2$ and FeCl₃ were purchased from Synth (Diadema, SP, Brazil). Analytical reagent grade of 99 % purity was employed throughout the experiments. Lime [CaCO₃ + Ca(OH)₂ + CaO] was obtained from Votoratim (Canoas, RS, Brazil). Lime is used as one of the inorganic components in preparation of activated carbon to avoid impregnation of the carbonaceous material with aqueous solution [28]. In the presence of water, lime forms a paste that holds all the solid constituents together [28].

The reagents *o*-nitrophenol, 2-naphthol, *o*-chlorophenol, *p*-nitrophenol, hydroquinone, resorcinol, *m*-cresol, bisphenol A, phenol, humic acid, sodium sulfate, sodium chloride, potassium phosphate, sodium carbonate, potassium nitrate, sodium hydroxide, and hydrochloric acid for preparation of simulated industrial effluents were purchased from Vetec (São Paulo, Brazil).

Preparation of AC adsorbents

AC adsorbents were prepared using the following procedure: 100.0 or 150.0 g of powdered sapelli wood sawdust (diameter $<250 \mu$ m) and 100.0 g of inorganic

components (20 g lime + 80 g ZnCl₂ or 20 g lime + 40 g ZnCl₂ + 40 g FeCl₃) and 45.0 mL H₂O were added and mixed thoroughly to produce a uniform mixture [28]. Before the pyrolysis step, the mixture was oven-dried at 90 °C for 120 min. Thereafter, 30.0 g of sample was placed in a quartz reactor that was inserted into a microwave oven under inert atmosphere (200 mL min⁻¹ nitrogen) [27]. The microwave was turned on for 320 s at 1200 W, then sat for 10 min for the system to cool down [27]. The material obtained after pyrolysis was named according to the composition of the activating agent and constituent ratios as follows: 1A (100 g sapelli wood sawdust + 20 g lime + 80 g ZnCl₂), 2A (150 g sapelli wood sawdust + 20 g lime + 40 g ZnCl₂ + 40 g FeCl₃), and 2B (150 g sapelli wood sawdust + 20 g lime + 40 g ZnCl₂ + 40 g FeCl₃).

To complete chemical activation, a leaching procedure was performed to eliminate remaining inorganic compounds ($ZnCl_2$ and $FeCl_3$) from the carbonized materials using 6.0 mol L⁻¹ HCl under reflux, as already described in literature [27, 28]. The carbonized materials 1A, 2A, 1B, and 2B gave rise to AC-1A, AC-2A, AC-1B, and AC-2B, respectively, after acidic leaching of inorganic contents.

Characterization of AC samples

The AC samples were ground and sieved for N₂ adsorption–desorption isotherm analysis. The resulting fine (<53 μ m) powder was used for adsorption experiments with a commercial system (TriStar II 3020; Micromeritics Instrument Corp.) at –196 °C after drying for 24 h at 150 °C under reduced pressure (<2 mbar). The surface area and pore size distribution of the AC samples were determined by the Brunauer–Emmett–Teller (BET) multipoint and Barrett–Joyner–Halenda (BJH) technique, respectively [29].

The surface of the AC samples was analyzed by scanning electron microscopy (SEM, JSM 6060; JEOL).

To determine their surface hydrophobicity/hydrophilicity, the AC samples were dried in 10-mL beakers at 70 °C for 24 h. The samples were then transferred to a desiccator and cooled to room temperature, then the accurate weight (ca. 0.3 g) of each sample was obtained. Afterwards, the beakers were arranged in capped (45/ 50 mm joint) Erlenmeyer flasks containing 60 mL solvent (water and *n*-heptane) inside a temperature-regulated shaker at 25 °C in static condition, in such a way that the glass containing the samples was not in contact with the solvent or wall of the Erlenmeyer flask. After 24 h, the sample was removed from the Erlenmeyer flask, dried carefully on the outside with laboratory tissues, and weighed again. The maximal amount of vapor adsorbed on the activated carbon was obtained as the difference between the final and initial weight, expressed in mg g⁻¹. The hydrophilic/hydrophobic properties were determined from the ratio of adsorbed *n*-heptane vapor (mg g⁻¹) to adsorbed water vapor (mg g⁻¹).

The pH_{pzc} values were obtained using a procedure described in literature [30].

The total acidity and basicity of the AC samples were determined using a modified Boehm titration method [31].

Adsorption studies

o-Cresol solution (20.00 mL) with concentration ranging from 5.00 to 500.0 mg L⁻¹ was added to 50.0-mL flat Falcon tubes containing 30 mg of each AC sample at pH values ranging from 4.0 to 10.0. The tubes were capped, then arranged horizontally inside a temperature-controlled shaker. The mixtures were shaken at speed of 150 rpm for between 1 and 120 min at 25 to 50 °C. Afterwards, the samples were centrifuged in a UniCen M centrifuge (Herolab) to separate the AC from the liquid phase, and 1–5 mL of supernatant was diluted to 10.0–50.0 mL in calibrated flasks using blank solution (aqueous solution with pH 4.0–10.0). After the batch adsorption experiment, the unadsorbed *o*-cresol was measured using a T90+ spectrophotometer (PG Instruments) at a maximum wavelength of 269 nm.

The sorption capacity for and percentage removal of o-cresol were calculated using Eqs. (1) and (2), respectively:

$$q = \frac{(C_0 - C_f)}{m} \times V, \tag{1}$$

% Removal =
$$100 \times \frac{(C_0 - C_f)}{C_0}$$
, (2)

where q is the sorption capacity of the adsorbent for o-cresol (mg g⁻¹), C_0 is the initial o-cresol concentration in contact with the AC (mg L⁻¹), C_f is the o-cresol concentration in equilibrium (mg L⁻¹), m is the weight of AC (g), and V is the volume of o-cresol solution (L).

Analytical control and statistical evaluation of nonlinear methods

To guarantee reliability, accuracy, and reproducibility of adsorption data, adsorption experiments were performed thrice [32]. The *o*-cresol solutions were placed in glass bottles which had been cleaned by submerging in 10 % HNO₃ overnight [33], washing with distilled water, drying in a oven at 70 °C, and storing [34].

Standard *o*-cresol solutions (100.0–300.0 mg L⁻¹) were used to determined analytical calibration curves. The detection limit of *o*-cresol was 0.030 mg L⁻¹ [35].

Fitting of kinetic and equilibrium data was performed using nonlinear methods (simplex method and Levenberg–Marquardt algorithm) in Origin 2015 software (Microcal). The suitability of the nonlinear models was evaluated using the coefficient of determination (R^2), adjusted coefficient of determination (R^{2}_{adj}), and standard deviation of residues (SD) [36, 37], as given by Eqs. (3), (4), and (5), respectively:

$$R^{2} = \left(\frac{\sum_{i}^{n} \left(q_{i,\exp} - \overline{q}_{i,\exp}\right)^{2} - \sum_{i}^{n} \left(q_{i,\exp} - q_{i,\text{model}}\right)^{2}}{\sum_{i}^{n} \left(q_{i,\exp} - \overline{q}_{i,\exp}\right)^{2}}\right),$$
(3)

$$R_{\rm adj}^2 = 1 - \left(1 - R^2\right) \cdot \left(\frac{n-1}{n-p-1}\right),\tag{4}$$

$$SD = \sqrt{\left(\frac{1}{n-p}\right) \cdot \sum_{i}^{n} \left(q_{i,exp} - q_{i,model}\right)^{2}},$$
(5)

where $q_{i,\text{model}}$ is each individual q value predicted by the model, $q_{i,\text{exp}}$ is each individual experimental q value, $\overline{q}_{\text{exp}}$ is the average of all experimental q values, n is the number of experiments performed, and p is the number of parameters in the model [36, 37].

Adsorption kinetics

Kinetic data were fit using pseudo-first-order, pseudo-second-order, and Avrami fractional-order models, as given by the mathematical Eqs. (6), (7), and (8), respectively [36, 37]:

$$q_t = q_e \cdot [1 - \exp(-k_1 \cdot t)], \tag{6}$$

$$q_t = q_e - \frac{q_e}{[k_2(q_e) \cdot t + 1]},$$
(7)

$$q_t = q_e \cdot \{1 - \exp[-(k_{\rm AV} \cdot t)]^{n_{\rm AV}}\}.$$
(8)

Adsorption equilibrium

The experimental equilibrium data were fit using the Langmuir, Freundlich, and Sips isotherm models, as represented by Eqs. (9), (10), and (11), respectively [36, 37]:

$$q_{\rm e} = \frac{Q_{\rm max} \cdot K_{\rm L} \cdot C_{\rm e}}{1 + K_{\rm L} \cdot C_{\rm e}},\tag{9}$$

$$q_{\rm e} = K_{\rm F} \cdot C_{\rm e}^{1/n_{\rm F}},\tag{10}$$

$$q_{\rm e} = \frac{Q_{\rm max} \cdot K_{\rm s} \cdot C_{\rm e}^{1/n_{\rm s}}}{1 + K_{\rm s} \cdot C_{\rm e}^{1/n_{\rm s}}}.$$
 (11)

Synthetic effluents

Two synthetic industrial effluents, consisting of a mixture of ten phenols, humic acid, and inorganics usually present in industrial effluents, were prepared. The compositions of the effluents are presented in Table 1. The aim of using synthetic effluents is to test the sorption capacity of the AC samples for removal of a mixture

Table 1 Chemical composition of simulated industrial effluents	Phenols	Concentration (r	Concentration (mg L ⁻¹)	
		Effluent A	Effluent B	
	o-Cresol	50.0	60.0	
	Phenol	10.0	15.0	
	<i>m</i> -Cresol	5.00	10.0	
	2-Chlorophenol	5.00	10.0	
	Bisphenol A	5.00	10.0	
	2-Nitrophenol	5.00	10.0	
	4-Nitrophenol	5.00	10.0	
	2-Naphthol	5.00	10.0	
	Hydroquinone	5.00	10.0	
	Resorcinol	5.00	10.0	
	Other organic component			
	Humic acid	10.0	20.0	
	Inorganic components			
	Sodium sulfate	40.0	40.0	
	Sodium carbonate	20.0	40.0	
	Sodium chloride	20.0	40.0	
	Potassium nitrate	20.0	40.0	
	Potassium phosphate	21.0	36.4	
	рН	7.0	7.0	

of phenolic compounds from a solution containing high concentrations of organic matter and salts.

Results and discussion

Characterization of activated carbon samples

The chemical activation of the sapelli wood with inorganics and further pyrolysis assisted by microwaves generated different activated carbon materials with different adsorption characteristics. Among the main features of the adsorbents, the surface area and porosity have the greatest influence on the adsorption process. N_2 adsorption-desorption isotherms and BJH plots for samples AC-1A, AC-1B, AC-2A, and AC-2B are presented in Fig. 1.

The nitrogen isotherms for the activated carbon materials are shown in Fig. 1a. The isotherms for all samples were classified as type I according to the International Union of Pure and Applied Chemistry (IUPAC) classification, being typical of microporous materials with pore width below 2 nm [38].

The pore size distribution curves for samples AC-1A, AC-2A, AC-1B, and AC-2B are presented in Fig. 1b-e, respectively. It can be seen that all the activated



Fig. 1 a Nitrogen adsorption–desorption isotherms for activated carbon samples (filled symbols for adsorption branch, hollow symbols for desorption branch). Pore size distribution for **b** AC-1A, **c** AC-2A, **d** AC-1B, and **e** AC-2B

carbon materials presented a mixture of micropores and mesopores, with a peak maximum at around 40 Å (4 nm).

The effects of the different types of chemical agents and impregnation ratios on the BET surface area, micropore area, external surface area, total pore volume, micropore volume, mesopore volume, and yield are presented in Table 2.

As can be seen from the results in Fig. 1a and Table 2, the adsorbed N₂ volume differed depending on the ratio of the activation agents used. The highest surface area was obtained for the AC-1A material (100 g sapelli wood sawdust + 20 g lime + 80 g ZnCl₂), followed by AC-1B (100 g sapelli wood sawdust + 20 g

Sample	BET surface area $(m^2 g^{-1})$	<i>t</i> -Plot micropore area $(m^2 g^{-1})$	<i>t</i> -Plot external surface area $(m^2 g^{-1})$	Total pore volume $(cm^3 g^{-1})$	<i>t</i> -Plot micropore volume $(cm^3 g^{-1})$	Mesopore volume $(cm^3 g^{-1})$	Yield (%)
AC-1A	914.08	518.65	395.43	0.55385	0.27025	0.28360	25.15
AC-1B	874.72	560.60	314.11	0.52327	0.25177	0.27151	25.91
AC-2A	805.06	435.17	369.89	0.44205	0.20864	0.23342	24.26
AC-2B	647.05	405.89	241.15	0.34443	0.20020	0.14424	24.88

Table 2 Textural properties of activated carbon materials

lime + 40 g ZnCl_2 + 40 g FeCl_3), then AC-2A (150 g sapelli wood sawdust + 20 g lime + 80 g ZnCl_2), and lastly AC-2B (150 g sapelli wood sawdust + 20 g lime + 40 g ZnCl_2 + 40 g FeCl_3); see Table 2. The difference in total surface area between AC-1B and AC-2A was only 8.0 % of the surface area of AC-1B. From these results, it is possible to infer that the ratio between the total amount of inorganics in the biomass and the inorganic components is very important for the surface area of, as well as pore development in, the activated carbon materials [39]. The adsorbents with higher inorganic content (AC-1A and AC-1B) presented higher surface area and higher pore volume compared with the adsorbents with lower amounts of inorganics introduced during preparation of the activated carbon (AC-2A and AC-2B) [40, 41].

Another parameter that is useful for analysis of these activated carbon materials is the ratio $S_{\text{micropore}}/S_{\text{total}}$, expressed as a percentage. This ratio was 56.74, 64.09, 54.05, and 62.73 % for AC-1A, AC-1B, AC-2A, and AC-2B, respectively. The lime + ZnCl₂ + FeCl₃ activating agent employed in the preparation of AC-1B and AC-2B resulted in a predominantly microporous adsorbent, whereas activating the carbon with lime + ZnCl₂ resulted in a material with a slightly lower amount of microporous material. However, comparing all the results, one can infer that each of the four activated carbon materials presented a mixture of micro- and mesopores.

Furthermore, analyzing these results, it is expected that the sorption capacity of AC-1B would be very close to that of AC-2A, since these materials did not exhibit remarkable differences in surface area (difference of only 8.0 %). Also the inorganic components used to form the paste with the organic precursor (20 g lime + 80 g ZnCl₂ or 20 g lime + 40 g ZnCl₂ + 40 g FeCl₃) presented practically the same performance for combinations with 1.0:1.0 (inorganic:organic) ratio. Only AC-2B presented worst surface area, being 19.6 % lower than for AC-2A, 26.0 % lower than for AC-1B, and 29.2 % lower than for AC-1A. Use of ZnCl₂ as an activating agent is well known in literature [39–41], whereas use of FeCl₃ as an activating agent for production of activated carbon is more recent in literature [27].

In addition, the ratio $V_{\text{micropore}}/V_{\text{total}}$ expressed as a percentage could be another useful parameter for analysis of these activated carbon materials. Its value was 48.79, 48.11, 47.20, and 58.13 % for AC-1A, AC-1B, AC-2A, and AC-2B, respectively. Analyzing these results, only AC-2B (150 g biomass + 20 g lime + 40 g ZnCl₂ + 40 g FeCl₃) presented a predominance of micropores,

because this ratio was slightly higher than 50 %. On the other hand, AC-1A, AC-1B, and AC-2A presented $V_{\text{micropore}}/V_{\text{total}}$ ratios lower than 50 %, indicating that these materials were predominantly mesoporous. Although this analysis differs from the analysis of the ratio $S_{\text{micropore}}/S_{\text{total}}$, for which all the activated carbon materials presented a value higher than 50 %, both analyses indicate that all the activated carbon materials possess micropores and mesopores in their structure, and these pores are responsible for the sorption capacity of the obtained adsorbents [42–45].

In preparation of activated carbon, yield is an important parameter, usually being defined as the final weight of activated carbon produced after activation, washing, and drying, divided by the initial weight of raw material, both on dry basis [27, 46]. Table 2 also presents the yield for carbon preparation achieved in this work. It is observed from Table 2 that the type and composition of activating agent was not relevant for the yield. The global yield of activated carbon varied in the range from 24.26 to 25.91 %, considering the initial mass of sawdust used.

Different results were obtained in other studies; For instance, Saucier et al. [27] used cocoa shells as precursor and achieved yield of 20 %, whereas Gomez-Serrano et al. [46] achieved 37.2–42.3 % for production of activated carbon from chestnut wood. Such differences in the yield of activated carbon can be attributed to the different pyrolysis conditions and precursor types used.

Fourier-transform infrared (FTIR) analysis was carried out to identify the functional groups on the surface of the AC materials. Such analysis enables better understanding of surface features that could promote improved uptake of *o*-cresol. The FTIR spectra for samples AC-1A, AC-1B, AC-2A, and AC-2B are presented in Supplementary Fig. 2, and their band assignments in Table 3 [27, 28, 47]. The activated carbon materials presented many functional groups on the surface, including O–H group of phenol, carboxylic acid, alcohol, aromatic rings, carboxylates, ether, and ester groups [27, 28, 47]. It seems that the main groups on the surface of the activated carbon materials were slightly hydrophilic, including oxygenated groups (carboxylates, phenol, esters, ethers, alcohols, etc.); this is in agreement with the *n*-heptane/water analysis, which provided additional evidence for the hydrophilic surface of the carbon materials (see Supplementary Fig. 2), as discussed below.

The functional groups present on the AC materials were also confirmed using a modified potentiometric Boehm titration method [31]. The total acidity was composed of carboxylic acid, phenols, and lactone groups. In this work, the fractions of these functional groups with NaCO₃ and NaHCO₃ were not determined, because it is very difficult to eliminate CO₂ from the aqueous solution, as observed in our laboratory by potentiometric titrations and as also discussed in literature [31]. The total acidity of samples AC-1A, AC-2A, AC-1B, and AC-2B was 0.559, 0.594, 0.601, and 0.606 mmol g⁻¹, respectively. The total basicity of all the activated carbon materials was 0.000 mmol g⁻¹, with the exception of AC-1B, for which it was 0.009 mmol g⁻¹. Based on these results, it is seen that lime + ZnCl₂ + FeCl₃ promoted a slightly higher amount of acidic groups on the activated carbon adsorbent compared with lime + ZnCl₂. It should be noted that such an increase of the functional groups on the surface of the activated carbon does not mean an increase in the sorption capacity, since the sorption depends on the interaction of the

FTIR band (cm ⁻¹)	
AC-1A	Assignment
3411	O-H stretching
2921	C-H asymmetric stretching
2853	C-H symmetric stretching
1610	Aromatic ring mode
1563	Asymmetric carboxylate stretching and aromatic ring mode
1452	Symmetric carboxylate stretching and aromatic ring mode
1421	Aromatic ring mode
1099	Asymmetric C-O-C ether and O-C-C of aromatic ester, C-O phenol
803	CH out-of-plane bends of aromatic rings
AC-2A	
3404	O-H stretching
2920	C-H asymmetric stretching
2852	C-H symmetric stretching
1572	Asymmetric carboxylate stretching and aromatic ring mode
1383	C-H bending
1136	Asymmetric C-O-C ether and O-C-C of aromatic ester, C-O phenol
885, 811	CH out-of-plane bends of aromatic rings
AC-1B	
3384	O-H stretching
2923	C-H asymmetric stretching
2852	C-H symmetric stretching
1583	Asymmetric carboxylate stretching and aromatic ring mode
1379	C-H bending
1167	Asymmetric C-O-C ether and O-C-C of aromatic ester, C-O phenol
885, 812, 752	CH out-of-plane bends of aromatic rings
AC-2B	
3417	O-H stretching
2918	C-H asymmetric stretching
2853	C-H symmetric stretching
1617, 1565	Asymmetric carboxylate stretching and aromatic ring mode
1460	Aromatic ring mode
1383	C-H bending
1104	Asymmetric C–O–C ether and O–C–C of aromatic ester, C–O phenol
803	CH out-of-plane bends of aromatic rings

Table 3 FTIR vibrational bands of AC-1A, AC-2A, AC-1B, and AC-2B

Assignments based on literature [25, 26, 40]

adsorbate with the adsorbent, while textural characteristics of the adsorbent such as the surface area and total pore volume also play a decisive role in the overall adsorption.

Surface features of the materials could influence the interaction between adsorbent and adsorbate in the adsorption process. The adsorption ratio of vapors of solvents with different polarity (water = hydrophilic, *n*-heptane = hydrophobic) [48-50] was used to characterize the surface of the AC materials; the results are shown in Fig. 2. As seen from this figure, the adsorption ratios between water and *n*heptane were below 1, implying that the AC materials have a more hydrophilic surface. Comparing the activating agents, samples AC-1B and AC-2B with FeCl₃ $(20 \text{ g lime} + 40 \text{ g ZnCl}_2 + 40 \text{ g FeCl}_3)$ were less hydrophilic compared with the samples prepared using $ZnCl_2$ (20 g lime + 80 g $ZnCl_2$), i.e., AC-1A and AC-2A (Fig. 2). For hydrophobic materials, the *n*-heptane/water ratio reaches values of 10 or more, e.g., for hydrophobic polysiloxanes [48]. However, composite materials containing silicate structure mixed with carbonized materials present *n*-heptane/ water adsorption ratios ranging from 2 to 4 [48]. Based on the adsorption results for the solvent vapors and the FTIR data, it can be concluded that the process of producing activated carbon using microwaves generated an oxygen-rich surface with hydrophilic characteristics.

The results shown in Supplementary Fig. 2 (FTIR analysis) and Fig. 2 (water/*n*-heptane adsorption) suggest that varying the composition or type of chemical activating agent did not have remarkable effects on the surface of the four activated carbon materials, being insufficient to explain the difference in their sorption capacities. Therefore, textural properties such as the specific surface area, total pore volume, and micropore and mesopore volumes will be the main characteristics causing the different sorption capacities of the activated carbon materials.

In the field of environmental science, pH_{pzc} determines how easily a substrate can adsorb potentially harmful charged components [30, 51]. When the pH of the solution exceeds pH_{pzc} , the adsorbent surface is negative and can adsorb positively charged species, whereas when $pH < pH_{pzc}$, the adsorbent surface presents positive charge, being suitable for adsorption of negatively charged species [30].

Figure 3 shows that the pH_{pzc} value for AC-1A and AC-2A (activated 20 g lime + 80 g ZnCl₂) was 4.96 and 5.28, respectively. However, for samples AC-1B



Fig. 2 Solvent vapor adsorption analysis



Fig. 3 Point of zero charge curves for AC samples

and AC-2B (activated with 20 g lime + 40 g ZnCl_2 + 40 g FeCl_3), the pH_{pzc} values were slightly different: 4.30 and 3.97, respectively. This means that the activated carbon materials obtained using FeCl₃ (AC-1B and AC-2B) as activating agent had surfaces with more acidic behavior compared with those prepared using ZnCl₂ (AC-1A and AC-2A). This behavior is in agreement with the total acidity of the activated carbon materials as described above, where the activated carbon materials obtained using FeCl₃ (20 g lime + 40 g ZnCl₂ + 40 g FeCl₃) presented more total acidic groups than those obtained using only ZnCl₂ (20 g lime + 80 g ZnCl₂) as activating agent.

Experiments using *o*-cresol solutions with initial pH in the range of 4.0-10.0 revealed that the initial pH of the adsorbate solution had no significant effect when its value was varied from 4.0 to 9.0. However, at pH 10, the percentage removal of *o*-cresol was decreased by at least 25 %. Considering that the pH of treated effluents should be close to neutrality for release into the environment, the initial pH of the *o*-cresol solution was fixed at 7.0 for all subsequent experiments.

Adsorption kinetics

Adsorption kinetics is known to be important in many heterogeneous systems and time-dependent processes. Kinetic analysis allows for determination of the rate of adsorption, which determines the time of contact between the adsorbent and adsorbate required for the system to reach equilibrium. Therefore, for evaluation and regeneration of an adsorbent, it is important to understand the rate of the adsorption process [18, 27, 36, 52].

To investigate the adsorption kinetics of *o*-cresol on the four AC materials, pseudo-first-order, pseudo-second-order, and fractional-order kinetic models were used. The fitting parameters for the kinetic models for all the AC samples for two initial concentrations of *o*-cresol are presented in Table 4. To verify the suitability of the models, the standard deviation of residues (SD) was taken into account, with a

	AC-1A		AC-2A		AC-1B		AC-2B	
	500.0 mg L^{-1}	700.0 mg L^{-1}	500.0 mg L^{-1}	700.0 mg L^{-1}	500.0 mg L^{-1}	700.0 mg L^{-1}	500.0 mg L^{-1}	700.0 mg L^{-1}
Pseudo-first-order								
$k_{\rm f}~({\rm min}^{-1})$	2.947	3.035	2.981	2.963	4.128	3.858	2.691	2.381
$q_{\rm e} \; ({ m mg} \; { m g}^{-1})$	215.3	238.9	161.1	179.7	166.1	247.5	152.3	160.6
R^2 adjusted	0.9986	0.9966	0.9971	0.9971	0.9992	0.9993	0.9983	0.9993
SD (mg g^{-1})	2.066	3.606	2.252	2.498	1.161	1.667	1.615	3.203
Pseudo-second-order								
$k_{\rm s}$ (g mg ⁻¹ min ⁻¹)	0.07238	0.05939	0.09036	0.07861	0.2520	0.1359	0.08331	0.07882
$q_{\rm e}~({ m mg~g}^{-1})$	216.9	241.1	162.5	181.3	166.6	248.4	153.6	162.9
R^2 adjusted	0.9999	0.9992	0.9992	0.9994	0.9996	0.9998	0.9998	0.9993
SD (mg g^{-1})	0.4411	1.728	1.1739	1.065	0.8411	0.8881	0.4324	1.037
Avrami fractional-ord	er							
$k_{\rm AV}~({\rm min}^{-1})$	0.3628	0.3648	0.3140	0.3360	0.3113	0.3251	0.2608	0.3608
$q_{\rm e}~({ m mg~g}^{-1})$	216.8	244.3	165.9	182.8	167.1	252.1	153.4	163.7
$n_{\rm AV}$	0.2742	0.1383	0.1179	0.1598	1.267	0.07278	0.3008	0.2301
R^2 adjusted	0.9999	0.9999	0.9998	0.9998	0.9998	0.9999	0.9998	0.9999
SD (mg g^{-1})	0.3422	0.6170	0.6194	0.5896	0.4764	0.2409	0.5435	0.3674
All values expressed t	to four significant e	digits. Conditions: i	initial pH of adsor	bate 7.0, adsorbent	dosage 1.5 g L^{-1}			

Table 4 Kinetic fitting parameters for adsorption of o-cresol onto adsorbents AC-1A, AC-2A, AC-1B, and AC-2B

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lower value indicating less difference between theoretical and experimental q_t values [27, 51].

The fractional-order model presented the lowest SD values (varying from 0.2409 to 0.6194) for all adsorbents and initial concentrations, meaning that the q_t values predicted by the fractional-order model were closer to those measured experimentally. The pseudo-first-order model presented SD values varying from 1.161 to 3.606, while the SD values for the pseudo-second-order model varied from 0.4324 to 1.728.

Such a fractional-order model indicates that the adsorption process is complex or has multiple pathways, with the possibility that the adsorption mechanism changes during the adsorption process [37, 51]. Instead of the adsorption mechanism exhibiting only an integer kinetic order, it could follow multiple kinetic orders that change during contact of the *o*-cresol with the adsorption process, usually being a fractional value.

To ensure that adsorption equilibrium was reached, in subsequent experiments, the contact time between the AC materials and *o*-cresol was kept at 120 min.

Adsorption equilibrium

The isotherm provides information about the relationship between the *o*-cresol adsorbed on the solid phase and its concentration in the liquid phase when the adsorption process reaches equilibrium [53, 54]. The fit curves (at 45 °C) and the parameters of the Langmuir, Freundlich, and Sips isotherm models are presented in Fig. 4 and Table 5, respectively.

The experimental procedure to obtain the isotherm data was carried out at various temperatures between 25 and 45 °C with contact time of 120 min, initial pH of *o*-cresol solution of 7.0, and adsorbent dosage of 1.5 g L⁻¹. All isotherms presented similar trends for all samples and temperature ranges. Table 5 presents the fitting parameters and adsorption isotherms for *o*-cresol onto AC-1A, AC-1B, AC-2A, and AC-2B at various temperatures. Based on the SD values, the Sips model was the most suitable to describe the equilibrium data for adsorption of *o*-cresol onto the four AC materials (AC-1A, AC-1B, AC-2A, and AC-2B) at all temperatures (25, 35, and 45 °C) (Table 5). The Sips model showed the lowest SD values, meaning that its theoretical q_e values were closer to those found experimentally (Table 5). Strengthening the argument for the suitability of the Sips model, as for the SD values, the R_{adj}^2 values also confirm that the Sips isotherm is best for modeling the adsorption of *o*-cresol onto our AC materials.

The effect of temperature on the percentage removal of *o*-cresol by the AC materials was also evaluated in this work; the results are presented in Table 5. For all the AC materials, it seems that temperature did not have a remarkable effect on the Q_{max} value, albeit increasing from 221.4 to 239.4 mg g⁻¹ for sample AC-1A, from 220.5 to 234.6 mg g⁻¹ for sample AC-2A, from 219.5 to 234.9 mg g⁻¹ for sample AC-1B, and from 183.4 to 198.8 mg g⁻¹ for sample AC-2B.



Fig. 4 Adsorption isotherms of o-cresol on activated carbon materials at 45 °C

Plots of Q_{max} as obtained from the Sips isotherm model at 45 °C versus S_{BET} and total pore volume are shown in Fig. 5a and b, respectively. It is observed that higher S_{BET} and V_{total} values correspond to activated carbon materials with higher sorption capacity. In practice, samples AC-2A and AC-1B had very similar Q_{max} values (234.6 and 234.9 mg g^{-1} , respectively), while the difference in their surface area was 8.0 % (805.06 versus 874.72 m²g⁻¹). The adsorbent material AC-1A presented the highest Q_{max} , compatible with its higher S_{BET} value of 914.08 m² g⁻¹. The value of Q_{max} for AC-1A was only 1.8 % higher than for AC-1B, since the S_{BET} value was only 4.5 % higher for AC-1A than AC-1B. From these plots, it is also possible to conclude that the activated carbon materials with inorganic:sapelli sawdust ratio of 1:1 presented higher surface area and consequently higher Q_{max} . In addition, FeCl₃ is not a bad activating agent when mixed with ZnCl₂. Analysis of total pore volume is provided in Fig. 5b. As in the earlier analysis, higher pore volume of the adsorbent led to higher sorption capacity. Based on this analysis, the order of the samples in terms of decreasing Q_{max} versus V_{total} was as follows: AC- $1A > AC-1B \approx AC-2A > AC-2B$. In general, the sorption capacity of an adsorbate (e.g. o-cresol) from aqueous solution by AC depends, in addition to S_{BET} , on various other factors including pore size, pore volume, micropore and mesopore volumes, etc. In this work, it was found that S_{BET} was not the only parameter influencing the o-cresol adsorption: basically, samples with higher total pore (micropore and mesopore) volume had higher o-cresol adsorption capacity (Fig. 5a, b). Sample AC-1A presented the highest total pore volume, both micropore and

Table 5 Langmuir, Freundlic	h, and Sips	isotherm pa	trameters for	r adsorption	of o-cresol	onto the ac	ctivated carl	oon materia	s			
	AC-1A			AC-2A			AC-1B			AC-2B		
	25 °C	35 °C	45 °C	25 °C	35 °C	45 °C	25 °C	35 °C	45 °C	25 °C	35 °C	45 °C
Langmuir												
$Q_{ m max}~(m mg~g^{-1})$	201.2	211.5	217.4	204.2	212.5	218.1	206.1	213.6	216.2	172.6	176.2	180.7
$K_{\rm L} ({\rm L mg}^{-1})$	0.08552	0.07202	0.08092	0.05684	0.05984	0.06727	0.03877	0.03670	0.03011	0.04417	0.04088	0.03047
$R^2_{ m adj}$	0.9055	0.9095	0.9112	0.8922	0.9279	0.8842	0.9393	0.9298	0.9415	0.9439	0.9352	0.9413
SD (mg g^{-1})	19.51	19.96	20.13	16.29	17.37	23.34	15.01	16.78	15.14	11.88	13.03	12.48
Freundlich												
$K_{\mathrm{F}} \Big(\mathrm{mg}\mathrm{g}^{-1} \left(\mathrm{mg}\mathrm{L}^{-1} ight)^{-1/n_{\mathrm{F}}} \Big)$	57.20	55.67	54.88	50.24	48.72	56.95	41.84	40.69	37.26	41.38	40.08	34.37
$n_{ m F}$	4.409	4.173	4.354	4.070	3.875	4.155	3.662	3.515	3.330	4.130	3.993	3.621
$R^2_{ m adj}$	0.9976	0.9993	0.9981	0.9932	0.9988	0.9981	0.9996	0.9985	0.9975	0.9987	0.9983	0.9958
SD (mg g^{-1})	3.063	1.729	2.715	4.071	2.204	2.956	1.081	2.449	3.303	1.789	2.132	3.372
Sips												
$Q_{\max} \ (\mathrm{mg} \ \mathrm{g}^{-1})$	222.4	228.4	239.2	220.5	229.7	234.6	219.5	227.7	234.9	183.4	190.2	198.8
$K_{\rm s}$ (L mg ⁻¹)	0.8651	0.9165	0.9235	0.8547	0.8854	0.9325	0.8555	0.8566	0.8635	0.8432	0.8755	0.8821
$N_{ m s}$	0.2363	4.063	4.172	3.196	3.530	3.424	3.432	3.515	3.222	3.263	3.827	3.613
$R^2_{ m adj}$	0.9975	0.9996	0.9991	0.9941	0.9989	0666.0	0.9997	0.9988	0.9977	0.9993	0.9984	0.9975
SD (mg g^{-1})	3.204	1.100	2.453	3.821	2.178	2.986	1.013	2.329	3.213	1.371	2.088	3.312
All values expressed to four 1.5 g L^{-1}	significant d	igits. Condi	tions: initia	bH fixed	at 7.0, conti	act time bet	ween adsor	bate and ac	lsorbent fixe	ed at 120 m	nin, adsorbe	nt dosage

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Fig. 5 Relations between Q_{max} of Liu isotherm at 45 °C and **a** surface area and **b** total pore volume



mesopore, as well as the highest *o*-cresol adsorption capacity, followed by samples AC-1B, AC-2A, and AC-2B, respectively.

Comparison of adsorption capacity of different adsorbents

Various adsorbents have been identified for adsorption of *o*-cresol as well as *m*-cresol and *p*-cresol; therefore, it is important to verify the performance of the adsorbents used in this work versus other adsorbents reported in literature. The Q_{max} value of *o*-cresol onto AC and other adsorbents is listed in Table 6 [55–60]. The Q_{max} values reported in Table 6 were obtained using the best experimental conditions of each study. As shown in this table, samples AC-1A, AC-2A, AC-1B, and AC-2B presented higher Q_{max} values compared with all other adsorbents presented in Table 6 [55–60]. This result is very important from the scientific point of view, because the adsorbents proposed in this work present the best performance when compared with the other adsorbents highlighted in Table 6, giving an indication that these adsorbents are very efficient.

Desorption studies

Adsorbent regeneration can be important to reduce the cost of the adsorption process in practical wastewater treatment systems [52, 58, 59]. Regeneration of the

Adsorbent	Phenolic adsorbate	$Q_{\rm max}~({\rm mg}~{\rm g}^{-1})$	Reference
Activated carbon from water hyacinth	o-Cresol	123.6	[55]
Activated carbon from water hyacinth	m-Cresol	123.1	[55]
Polymer-supported iron nanoparticles	o-Cresol	1.02	[56]
Activated carbon from coconut shell	p-Cresol	30.23	[56]
Commercial activated carbon	Cresol	4.74	[57]
Fly ash	Cresol	6.7	[58]
Clay and membrane	o-Cresol	8.8	[58]
Montmorillonite	o-Cresol	10.1	[58]
Carbon composites	p-Cresol	86	[59]
Parthenium-based activated carbon	p-Cresol	62.91	[<mark>60</mark>]
AC-1A microwave-activated carbon	o-Cresol	239.2	This work
AC-1B microwave-activated carbon	o-Cresol	234.6	This work
AC-2A microwave-activated carbon	o-Cresol	234.9	This work
AC-2B microwave-activated carbon	o-Cresol	198.8	This work

 Table 6 Maximum sorption capacity of different adsorbents used for removal of various phenolic compounds

activated carbon materials was examined through desorption experiments. Therefore, desorption experiments were carried out to regenerate all the adsorbents. NaOH (1.0–5.0 M) and 10 % EtOH + NaOH M and 50 % EtOH + NaOH M were used as eluents in the *o*-cresol desorption process (Fig. 6).

Pure NaOH solutions, up to 5 M, presented good recovery of adsorbents (between 74.84 and 81.56 %). However, the mixture of EtOH (10 and 50 %) + NaOH (5 M) presented excellent desorption efficiency, desorbing more than 99.9 % of *o*-cresol from all the activated carbon materials. These results show that addition of EtOH caused obvious enhancement of the amount of *o*-cresol desorbed.

Fig. 6 Desorption of *o*-cresol from samples AC-1A and AC-2A



Proposed adsorption mechanism

Based on the characterization analysis, pH studies, kinetic and adsorption processes, and desorption experiments, the adsorption mechanism of o-cresol onto the activated carbon materials can be summarized as shown in Fig. 7. The adsorption mechanism involved in the adsorption process between the carbon surface and ocresol might be governed by physical interactions such as $\pi - \pi$ interactions, hydrophobic interactions, and/or electron donor-acceptor complexation [57-60]. The electron donor-acceptor mechanism relies on aromatic rings of the o-cresol to act as electron acceptors while basic zones on the AC surface serve as electron donors [57]. The π - π interactions are physical interactions between π electrons of aromatic rings of the o-cresol and with π electrons of the rings of AC [57–60], as shown in Fig. 8. Moreover, there is also a strong possibility that o-cresol sorption occurs through formation of hydrogen bonding with the -OH group present on the AC surface (Fig. 8) [57–60]. The AC materials present strong adsorption capacity for o-cresol, which could be related to formation of such hydrogen bonds [57-60]between o-cresol and oxygenated groups of the activated carbon. In the desorption experiments, the organic solvent EtOH helped to break up the interactions of ocresol with the activated carbon surface, leading to desorption of o-cresol.

Adsorption of simulated phenolic effluents

Two simulated effluents containing several phenolic compounds, humic acid, and some salts with different compositions and concentrations were prepared (Table 1) to investigate the effectiveness and efficiency of treatment of simulated synthetic effluents using the activated carbon materials (Fig. 8).

The spectra of the synthetic phenolic effluents before and after treatment with the AC materials were recorded between 190 and 800 nm in the ultraviolet-visible (UV-Vis) region (Fig. 8). The areas under the absorption bands from 190 and 800 nm were used to measure the amount of phenolic compounds removed from the synthetic effluents. All of the activated carbon materials presented very good performance in treatment of the simulated effluents. The removal percentages for effluent A were 98.87, 97.42, 98.13, and 96.61 % for AC-1A, AC-1B, AC-1B, and AC-2B, respectively. For effluent B, which was slightly more concentrated than effluent A (Table 1), the removal of phenolic compounds was 96.81, 95.95, 96.31,



Fig. 7 Proposed adsorption mechanism between AC and o-cresol



and 93.82 % for AC-1A, AC-1B, AC-1B, and AC-2B, respectively. These results are in accordance with the porosity and adsorption data, which showed that the sorption capacity followed the order: AC-1A > AC-1B > AC-2A > AC-2B, which is the same order as for Q_{max} versus S_{BET} in Fig. 4.

Based on the removal efficiency for these synthetic effluents, it can be inferred that these AC materials might have very good applicability for removal of phenolic compounds from real industrial wastewater contaminated with phenols.

Conclusions

This study has proven that activated carbon materials with high S_{BET} values can be prepared from sapelli wood sawdust via a microwave heating process. The AC materials were microporous with S_{BET} values up to 914.08 m² g⁻¹. Variation of the

ratios of the activation agent resulted in noticeable changes in the textural properties of the prepared activated carbon materials, as well as their surface chemistry.

Kinetic and equilibrium studies confirmed that the Avrami fractional-order model and Sips isotherm provided the best fit, indicating multilayer adsorption. The maximum amount of *o*-cresol adsorbed at 45 °C was 239.2, 234.6, 234.9, and 198.8 mg g⁻¹ for AC-1A, AC-2A, AC-1B, and AC-2B, respectively. The AC materials showed excellent results in treatment of simulated industrial effluents, effectively removing at least 93.82 % of mixtures containing high concentrations of phenols, organic matter, and saline concentrations. An adsorption mechanism is proposed, suggesting that donor–acceptor complexation, hydrogen bonding, and π – π interactions play key roles in the adsorption process.

Based on the presented data, it is possible to conclude that microwave heating enables production of efficient activated carbon with high surface area from sapelli wood sawdust.

Acknowledgments We would like to thank the National Council for Scientific and Technological Development (CNPq, Brazil) and The Academy of Sciences for Developing World (TWAS, Italy) for financial support and sponsorship. We also thank The Centre of Electron Microscopy (CME-UFRGS) for use of the SEM. We are once again grateful to Chemaxon for providing us with an academic research license for Marvin Sketch software version 16.6.6.0 (http://www.chemaxon.com) 2016, used for *o*-cresol physical–chemical properties.

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