Activated carbon from flash pyrolysis of eucalyptus residue

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Abstract

Forestry waste (eucalyptus sp) was converted into activated carbon by initial flash pyrolysis followed carbonization and CO\textsubscript{2} activation. These residues were obtained from a pilot plant in Spain that produces biofuel, the biochar represented 10–15\% in weight. It was observed that the highest activation was achieved at a temperature of 800 °C, the specific surface increased with time but, on the contrary, high loss of matter was observed. At 600 °C, although there was an important increase of the specific surface and the volume of micropores, at this temperature it was observed that the activation time was not an influential parameter. Finally, at 400 °C it was observed that the activation process was not very significant. Assessing the average pore diameter it was found that the lowest value corresponded to the activation temperature of 600 °C, which indicated the development of microporosity. When the activation temperature increases up to 800 °C the pore diameter increased developing mesoporosity.

Keywords: Engineering, Agriculture

1. Introduction

Activated carbon is a form of carbon that has been processed to create pores, resulting in an increased surface area. The surface area of activated carbon makes the material suitable for many processes: metal extraction from industrial waste
water [1, 2, 3 and 4], water purification [5, 6], sewage treatment [7, 8], air conditioning filters [9], gas masks manufacturing [10, 11], etc.

Activated carbon can be produced from carbonaceous source materials such as eucalyptus wood [12] and acacia [13, 14], waste wood from old wooden houses and tips in saw-mills [15], lignin (which is a waste emitted from paper mill and that normally is used like fuel) [16], cork oak [17], apricot stone [18], peanut shell [19], rice husks [20], ears of corn [21] or cellulose-based paper impregnated with a petroleum pitch [22].

The process to obtain an activated carbon begins with the pyrolyzation, dehydration and volatilization of the specimen. Therefore, biochar with high carbon content is obtained after elements that are non-carbonaceous such as hydrogen and oxygen are eliminated. The pyrolysis process is performed in the absence of oxygen in order to avoid combustion. The specimen starts to develop an internal porosity, low compared to that developed afterwards during the activation process. Tar substances and carbonaceous residues detached during heating are eliminated during the activation process, opening current pores and developing new ones. The activation can be achieved following a physical [23] or a chemical [24, 25 and 26] procedure.

Physical activation involves carbonization followed by activation of the resulting char in the presence of activating agents. As to the possible physical treatments, the one standing out uses water vapour [27, 28, 29 and 30] or carbon dioxide as activating gas [31, 32], or a mixture of both [33].

Chemical activation is known as a single step method for the preparation of activated carbon in the presence of chemical agents. Chemical activation usually requires lower temperatures and shorter times for activation of the material. It can improve the pore development in the carbon structure because the effect of chemicals. The carbon yields obtained by chemical activation are higher than by physical activation. However, it requires a process after the thermal treatment that recovers the chemical agents, which might restrict its application due to environmental issues [34].

The main characteristic of the activated carbon is that possess a high adsorption capacity, as a result of the high degree of microporosity. The micropores have molecular dimensions, the effective radii being less than 2 nm. The adsorption in these pores occurs through volume filling, and there is no capillary condensation taking place. They usually constitute the 95% of the total surface area of the activated carbon (1000–1500 m²/g), being micropore volumes from 0.15 to 0.7 cm³/g.

The activated carbons have surface area and pore volume that range between 250–2410 m²/g and 0.022–91.4 cm³/g, respectively [35]. Although the major part
of the adsorption take place through micro pores, the mesopores and macropores are also important due to enable sorbent molecules passing rapidly to smaller pores situated more internally.

Adsorption isotherms are usually experimentally determined in physical adsorption. The shape of an adsorption isotherm can provide preliminary qualitative information on the adsorption mechanism as well as on the porous structure of the carbons [36, 37 and 38]. The amount of gas adsorbed at a given temperature for different relative gas pressures is known as adsorption isotherms (IUPAC recognize 6 types of isotherms of physical adsorption [39]).

Numerous research works can be found in the literature concerning the estimation of properties related to the adsorption capacity of the activated carbons, i.e. specific surface, total volume of pores, pore size distribution, initial concentration of sorbent, temperature, degree of carbonization [40], contact time [24] and pH on the adsorption process [41, 42]. The effect of the temperature on the development of the porosity of a coal depends on the source material, heating rate and the residence time in the oven. The temperatures, where generally are reached good results in the pyrolysis and later on in the activation with CO₂ of lignocellulosic materials, are in the range between 350 °C and 850 °C [13, 23 and 43]. Out of these limits usually is not observed a significant improvement of the porosity.

A challenge in the production of activated carbon is to obtain very specific carbons with a given pore size distribution, from low cost materials [44] at low temperature. In this work, the preparation of activated carbon from a carbonaceous residue generated in a flash pyrolysis process of forestry waste (eucalyptus sp.) has been investigated.

The thermal degradation of a lignocellulosic material through a process of pyrolysis flash, for a short time and a high rate of heating, is commonly used for obtaining bio-oil [45, 46 and 47]. In contrast with other research works that have used eucalyptus wood as precursor material [12, 13], in this research is used a residue from a pyrolysis flash in order to valorize it. This residue was subjected to carbonization followed by carbon dioxide activation, in order to study the evolution of the specific surface and the porosity in relation to the activation temperature and the time of treatment.

2. Experimental

The main aspects that influence the activation of carbons are: the nature of the specimen, particle size, temperature, heating rate and residence time, in addition to the type of furnace and the process followed [48].
2.1. Raw material

The residue that has been investigated in this research is the product resulting from the flash pyrolysis of eucalyptus (barks, leaves and branches) at 650 °C in a fluidized bed reactor, vertical up-flow, at low pressure, and in the presence of a non-oxidizing atmosphere, sand bed and with recirculation of gases. Table 1 shows the main chemical features of this product. In the selection of the type of residue for the preparation of activated carbon, factors such as high carbon content, low inorganic compounds (i.e. low ash) content, or sufficient volatile matter content were taken into account [49].

By means of laser diffraction of the residue and the ashes were identified a significant presence of quartz (SiO₂). Because of the high intensity signal emitted by SiO₂, the presence of other minerals could be hidden. Therefore, it was carried out a demineralization with hydrochloric acid (HCl) to eliminated oxides, and later it was attacked with hydrofluoric acid (HF) to eliminate silicates. The resultant diffraction of the attacked product did not showed any peak. This could mean that the forestry waste was composed basically of organic matter, being the unique mineral substance presented the SiO₂, probably obtained when the lignocellulosic residues from pruning were collected.

2.2. Test set-up

Carbonization and activation tests of the residue have been carried out in a horizontal tubular furnace shown in Fig. 1.

With the introduction of N₂ into the circuit an inert atmosphere is achieved. N₂ from a cylinder is introduced into the circuit at a rate of 15 l/h, regulating the amount of gas by a flow controller.

The specimen is placed into a quartz tube of 75 mm diameter closed at both ends in order to confine the activation gases.

Table 1. Chemical characterization raw material & carbonaceous residue.

|                     | Proximate analysis (% dry basis) | Moisture (as received) | Total sulfur (% dry basis) |
|---------------------|---------------------------------|------------------------|---------------------------|
|                     | Ashes   | Volatile matter | Fixed carbon |                      |                     |
| Forest residue      | 4.11    | 73.75          | 22.14        | 4.50                  | 0.08                |
|                     | 3.25    | 24.90          | 71.85        | 4.70                  | 0.06                |
| Carbonaceous residue| C       | H              | N            | S                     | O*                  |
|                     | 74.25   | 3.67           | 0.18         | 0.05                  | 21.85               |

*By difference.
The programmable control of the oven allows, by means of a thermopar “Type K” situated in the centre of the tube, to establish the final heating temperature (electrically heated). It has been proved that the temperature gradient along the half-metre of the tube that is inside the oven is negligible. Therefore, in this part of the tube the temperature can be considered homogeneous.

The initial sample is placed in three ceramic vessels, and located in the central zone of the horizontal tube. By means of an oxygen detector is controlled the absence of this gas in the interior of the circuit.

Activation and carbonization processes are influenced by the heating rate and the decomposition temperature of biomass. Based on studies of other authors, where it has been observed that the heating rate vary between 5–20 K/min [50, 51, 52, 53 and 54], in this research it has been chosen a slow heating ramp of 7 K/min, where the development of secondary reactions are ensured (oil cracking and repolymerisation) [55] and it is allowed to obtain higher quantities of biochar.

The carbonization tests with CO₂ were carried out at different temperatures 400 °C, 600 °C and 800 °C, following research carried out from other authors [56, 57 and 58] using the same procedure. Each of the carbonized products obtained is activated to these same temperatures (i.e. the carbonized product at 400 °C was activated at 400 °C). The temperature of 400 °C was selected because is when the devolatilization of the three constituent polymers of lignocellulosic materials (hemicellulose, cellulose and lignin) is almost complete. In this research work an intermediate temperature of 600 °C was also selected, and then a maximum

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**Fig. 1.** Sketch of the experimental set and description of elements.

01: Gas cylinder (N₂/CO₂)  06: Oven thermocouple
02: Mass flow controller  07: Ceramic vessels
03: Silicone caps  08: Oximeter
04: Cylindrical quartz tube  05: Horizontal tubular furnace

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temperature of 800 °C where the efficiency to obtain active carbon significantly decrease [59].

2.3. Carbonization

About 7.5 g of the sample (particle size <0.125 mm) was used in these experiments, which was carbonized under ultra-high purity nitrogen flow (250 ml. min\(^{-1}\) STP). The preheating rate was 7 K/min to reach the carbonization temperatures (400 °C, 600 °C and 800 °C) maintaining these temperatures during 2 h, 4 h and 8 h.

2.4. Activation

In the following stage the physical activation was carried out with CO\(_2\) at flow rate of 200 ml.min\(^{-1}\)STP. The activation temperatures were 400 °C, 600 °C and 800 °C with a heating rate of 7 K/min, and an activation time of 2 h, 4 h and 8 h. At the end of the activation process N\(_2\) flow was again used during 2 h to reach the ambient temperature. The specimens of activated carbon were classified and preserved in a dryer to avoid the absorption of moisture from the ambient. Later on, the degree of porosity developed was analysed.

2.5. Characterization

Gas adsorption experiments were performed by using a Micromeritics ASAP 2000 analyzer, in order to determine the pore structure of both the chars and the activated carbons by N\(_2\) adsorption-desorption at 77 K.

The surface area was determined by using the Brunauer, Emmett and Teller (BET) method, the micropore area and volume by using the Boer method and the total pore volume and pore size distribution by using the Barret-Joyner-Halenda (BJH) method.

Scanning electron micrographs (SEM) of selected samples were also obtained. The apparatus used was a Hitachi S-570 Scanning Electron Microscope to observe the presence of porosities and microporosities of the samples. The resolution was 3.5 nm, 20x-200.000x with an accelerating voltage between 0.5 and 30 kV. As the samples were not electrically conductive, they were gold plated (<10 nm) using a Polaron-Bio-Raid apparatus.

3. Results and discussion

3.1. Surface area and pore volume

Table 2 shows the results obtained during carbonization at three temperatures, needed for the characterization of the porous structure of the chars obtained. It is
Table 2. Characterization of the porous structure of biochar produced at different carbonization temperatures.

| Temp. (°C) | Time (h) | Ash (%) | Loss of weight (%) | Surface area BET (m²/g) | Micropore area (m²/g) (%) | Total vol. BJH (cm³/g) | Micropore volume (cm³/g) (%) | Average pore diameter (nm) |
|------------|----------|---------|-------------------|-------------------------|--------------------------|------------------------|-----------------------------|---------------------------|
| 400        | 2        | 3.9     | 8.90              | 52.13                   | 4.82                     | 0.0358                 | 0.0017                      | 4.71                      | 2.749                     |
|            | 4        | 4.2     | 8.98              | 37.52                   | 4.08                     | 0.0348                 | 0.0013                      | 3.80                      | 3.711                     |
| 600        | 2        | 4.4     | 23.11             | 370.49                  | 296.39                   | 0.1948                 | 0.1381                      | 70.14                     | 1.594                     |
|            | 4        | 4.0     | 23.97             | 367.25                  | 298.97                   | 0.1928                 | 0.1389                      | 72.03                     | 1.576                     |
| 800        | 2        | 4.8     | 30.53             | 435.09                  | 366.69                   | 0.2314                 | 0.1728                      | 74.68                     | 1.582                     |
|            | 4        | 5.4     | 29.92             | 436.51                  | 371.10                   | 0.2285                 | 0.1736                      | 75.95                     | 1.571                     |
shown that the temperature has a remarkable influence in the increase of the surface area BET and microporosity.

A change is observed in the transition from 400 °C to 600 °C, and also it continues increasing, to a lesser degree, at 800 °C; but on account of a high loss of matter (about 30%). However, there is not an influence of the time of treatment in the carbonization stage. Ash production remains practically constant in this period as it is shown in Table 2.

Table 3 reports the characterization of the porous structure of activated carbons prepared by using CO2 flow at the same temperatures than the carbonization stage.

The maximum activation is achieved at the temperature of 800 °C, where there is a great influence of the activation time but, on the contrary, a high loss of matter (about 90%) is produced. Because of that, it was not considered to carry out the testing over 4 h.

At 600 °C there is a remarkable increase of the area and volume of micropores remaining practically invariant along this temperature, showing that the influence of time is not so significant. The figures of the BET surface area are influenced by the use of a precursor for the activated carbon. However, the activation process is less significant at 400 °C.

A difference among the activated samples is observed at 800 °C, where the time parameter increases the specific surface in a more pronounced manner than at 400 °C and 600 °C. Furthermore, in this temperature the greater increase of the specific area and volume of micro pores is produced. However, in comparison with the activated samples at 600 °C the micropores decrease at 800 °C, indicating that the material is mainly microporous at that temperature, and when is activated at higher temperature a development of mesoporosity is produced. This result is confirmed in the assessment of the figures obtained from average diameters, where at 600 °C the activated carbon presents the smallest pore diameter with the presence of plentiful micropores. When the activation temperature increases up to 800 °C the pore diameter grows in favour of the mesoporosity.

Gases removed during the thermal degradation of the lignocellulosic material are essentially CO2, CO, H2, CH4, C2H6 and C2H4. These are significantly removed within the range of temperatures 305–375 °C [19]. So, it could be an explanation to the low sensitivity at higher temperatures (600 °C and 800 °C in this study) when activation occurs. This can be also observed in other studies [13], once it has been developed microporosity (as in the case of the eucalyptus in this study at 600 °C) the activation time also seems to have less influence.
Table 3. Characterization of the porous structure of activated carbon by CO₂ flow at different temperatures.

| Temp. (°C) | Time (h) | Ash (%) | Loss of weight (%) | Surface area BET (m²/g) | Micropore area (m²/g) (%) | Total vol. BJH (cm³/g) | Micropore volume (cm³/g) (%) | Average pore diameter (nm) |
|------------|----------|---------|-------------------|-------------------------|--------------------------|------------------------|----------------------------|---------------------------|
| 400        | 2        | 3.6     | 9.10              | 72.97                   | 8.15                     | 0.0391                 | 0.0016                    | 4.17                      | 2.144                     |
|            | 4        | 3.5     | 10.11             | 134.56                  | 17.76                    | 0.0689                 | 0.0038                    | 5.59                      | 2.048                     |
|            | 8        | 3.9     | 11.11             | 181.78                  | 34.10                    | 0.0933                 | 0.0257                    | 27.60                     | 2.054                     |
| 600        | 2        | 4.7     | 13.33             | 376.70                  | 304.74                   | 0.1948                 | 0.1425                    | 73.17                     | 1.546                     |
|            | 4        | 5.0     | 25.03             | 389.96                  | 336.69                   | 0.1987                 | 0.1563                    | 78.66                     | 1.539                     |
|            | 8        | 5.2     | 27.25             | 410.06                  | 326.02                   | 0.2186                 | 0.1570                    | 71.81                     | 1.547                     |
| 800        | 2        | 10.6    | 66.77             | 905.02                  | 544.36                   | 0.5488                 | 0.2538                    | 46.24                     | 1.794                     |
|            | 4        | 25.4    | 89.88             | 1034.07                 | 570.09                   | 0.6129                 | 0.2596                    | 42.35                     | 1.779                     |
3.2. Pore size distribution

Fig. 2 shows the 77 K adsorption-desorption isotherms of activated carbons obtained at different temperatures by using CO\textsubscript{2}. It is observed that at 400 °C and 600 °C the adsorption isotherms are typical of the microporous activated carbons, showing the phenomenon so-called low pressure hysteresis, due to the deformation of the micropores caused by the sorbent during the adsorption process. Assessing the volume of the micropores, it is checked that the microporosity is inside the rank of the narrow micropores, i.e. those micro pores filled at very low relative pressures as consequence of the high adsorption potential between their walls. In the series of 800 °C the isotherms of the activated samples showed hysteresis, characteristic of capillary condensation in the mesopores. This suggests that the activation at high temperature develops the mesoporosity, as it could be seen by the increase of the slope in the adsorption branch when the activation time increases. These types of hysteresis cycles seem to be associated to the narrow pores with the shape of “horseshoe”, frequent in active carbons.

In order to provide further information, the sample was analysed performing SEM analysis. The sample was firstly carbonized and then activated with CO\textsubscript{2}. Therefore, the internal porous structure of the material obtained could be observed.

3.3. Particle morphology characterization

In Fig. 3 are shown Scanning Electron Microscopy (SEM) micrographs of a carbonized sample 120x-1.200x at 800 °C during 4 h. In this figure it is observed the morphological structure, where disorganized narrow pores are found.
As a consequence of the carbonization, changes were produced in the original structure of the sample of eucalyptus. In this way, it was obtained a solid and homogeneous material with presence of abundant grooves. A grooved structure of carbon is observed in the surface that folds up because of the temperature, as well as the presence of loose material that must be eliminated through sweep activation gas in the later stage.

In Fig. 4 are shown SEM micrographs of an activated sample, at 800 °C during 4 h. In these figures narrow pores appeared with a characteristic “horseshoe” shape being consistent the results shown in Fig. 2.

![Fig. 3. SEM of a sample prepared by carbonization at 800 °C during 4 h.](image)

![Fig. 4. SEM of a sample activated at 800 °C during 4 h.](image)
In the activation stage the superficial morphology of the material change to cylindrical with the presence of abundant pores connected that allows the access to the micro porous internal zones. The carbonaceous structure that is obtained is much weaker as a consequence of the material loss.

4. Conclusions

The carbonization stage of the carbonaceous residue generated in a flash pyrolysis process of eucalyptus devolatilises the char and creates a microporous structure which falls mainly within the rank of narrow micropores.

The maximum physical activation with CO₂ flow has been produced at 800 °C with a well-developed porosity compatible with BET apparent surface area values, similar to those of commercial activated carbons used as adsorbents in gas and liquid-phase separation processes, purification of products and gas and water cleaning operations. At this temperature there is great influence of the time parameter of activation, but on the contrary a high loss of matter is produced (about 90%).

Although the activation at 600 °C is higher than at 400 °C, in both of them the influence of the time is not significant, remaining practically constant.

The activated carbon obtained shows the lowest pore diameter at 600 °C, and when the activation temperature increases up to 800 °C the pore diameter grows in favour of the mesoporosity.

The N₂ adsorption-desorption isotherms forms of the active carbon obtained by CO₂ activation indicate the form and size of the pores. Those verify at 800 °C the presence of mesoporosity, while at 400 °C and 600 °C the isotherms are typical of microporosity carbons, with low pressure hysteresis phenomenon.

The temperature reached during the pyrolization and activation processes is responsible of the internal folding of the carbonaceous structure, obtaining morphology of the residue organized in cylindrical layers.

5. Declarations

Author contribution statement

Carlos Grima-Olmedo: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Álvaro Ramírez-Gómez: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Wrote the paper.
Dulce Gómez-Limón: Performed the experiments; Analyzed and interpreted the data.

Carmen Clemente-Jul: Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data.

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The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.

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