Tailoring Low-Cost Granular Activated Carbons Intended for CO\textsubscript{2} Adsorption

Marcos Juliano Prauchner \textsuperscript{1}\textsuperscript{*}, Silvia da Cunha Oliveira \textsuperscript{1} and Francisco Rodríguez-Reinoso \textsuperscript{2†}

\textsuperscript{1} Instituto de Química, Universidade de Brasília, Campus Darcy Ribeiro, Brasília, Brazil, \textsuperscript{2} Departamento de Química Inorgánica, Universidad de Alicante, Alicante, Spain

Physical adsorption on activated carbons has shown to be a very attractive methodology for CO\textsubscript{2} separation from flue gas streams and biogas. In this context, the goal of this work was to prepare granular activated carbons intended for CO\textsubscript{2} adsorption from an abundant and low-cost biomass residue (coconut shell) by using practical and cost-effective procedures. By the first time, parameters involved in chemical activation with dehydrating agents (H\textsubscript{3}PO\textsubscript{4} or ZnCl\textsubscript{2}) and/or physical activation with CO\textsubscript{2} were systematically screened in depth in order to obtain materials with improved performance for CO\textsubscript{2} adsorption on a volume basis. Compared with the commonly used mass basis, the data expressed on a volume basis are very important for industrial applications because they permit to estimate the efficiency of a fixed bed adsorption column. The work permitted to prepare granular activated carbons with a blend of relatively high gravimetric CO\textsubscript{2} uptake and bulk density, so that high volumetric CO\textsubscript{2} uptakes were attained. The highest values were 2.67 and 1.17 mmol/cm\textsuperscript{3} for CO\textsubscript{2} pressures of 1.0 and 0.15 bar, respectively. It is remarkable that the obtained results were similar to those reported by other authors for carbons chemically activated with KOH, the activation methodology that has been widely claimed as the one that produce ACs with the best performances for CO\textsubscript{2} adsorption, but which involves severe restrictions. Therefore, the present work can be considered a very important step in paving the way toward making CO\textsubscript{2} adsorption an each time more interesting technology to reduce the emissions of anthropogenic greenhouse gases.

Keywords: capture of CO\textsubscript{2}, CO\textsubscript{2} separation, activated carbon, adsorption, chemical activation, physical activation, flue gas, biogas

INTRODUCTION

Global warming has been considered as one of the most serious problems of the twenty-first century because it has threatened global economies and the environment. In spite of existing controversies, the problem has been widely attributed to the increase in the atmospheric concentration of greenhouse gases (GHGs), of which CO\textsubscript{2} and CH\textsubscript{4} are the most important ones. Anthropogenic CO\textsubscript{2} is extensively emitted due to the burning of fossil fuels in power plants and other combustion engines (McMillan et al., 2005). In turn, the main source of atmospheric CH\textsubscript{4} is the biogas generated by anaerobic digestion of biodegradable organic matter by microorganisms in sewage treatment plants, landfills and digestion plants to treat manure and industrial/agricultural wastes. In this
sense, the capture of CO₂ at large fixed emission sources and the recovery and utilization of biogas have been considered crucial strategies for reducing the concentration of GHGs in the atmosphere, as depicted in the next two Subsections.

**CO₂ Capture**

In spite of the efforts to replace fossil fuels by renewable fuels, the former certainly will continue to be the primary global energy source at least in the foreseeable future. Therefore, the capture of CO₂ at large fixed emission sources such as thermoelectric power plants and industrial plants (e.g., steel mills, refineries and cement kilns) has been increasingly employed as a strategy for reducing the emissions of the gas. After separated from the flue gas stream (which contains mainly CO₂ and N₂, with typical CO₂ concentrations between 10 and 15%), the CO₂ is compressed into a supercritical fluid, transported and safely deposited in the ground or an ocean-bedrock sediment layer (the so-called carbon capture and storage approach—CCS) (Huaman and Jun, 2014; Leung et al., 2014; Lee and Park, 2015; Rashidi and Yusup, 2016; Leeson et al., 2017; Bui et al., 2018). Alternatively, efforts have been directed toward CO₂ recycling and reuse in industry, agriculture, oil recovery, and energy production (the so-called carbon capture and utilization approach—CCU) (Kikuchi, 2003; Aresta and Dibenedetto, 2007; Leung et al., 2014; Bui et al., 2018; Fu et al., 2019).

**Biogas Upgrading and Use**

Biogas consists primarily of CH₄ (40–75%) and CO₂ (15–60%), besides minor quantities of other components such as H₂O, H₂S, siloxanes, hydrocarbons, NH₃, O₂, CO, and N₂. Due to its high CH₄ content, biogas is a potential fuel for the generation of electrical power and heat. In this sense, after an appropriate pre-treatment, the biogas can be used as a natural gas substitute in applications such as car fuel, production of hydrogen by steam reforming for fuel cells, and syngas production. Besides removing contaminants harmful to the natural gas grid, appliances and end-users, the referred pre-treatment (the so-called biogas upgrading or enrichment) aims to reduce the CO₂ content in the fuel in order to increase its calorific heat accordingly to the envisaged application. After that, the final product typically contains 95–97% of CH₄ and 1–3% of CO₂, being referred to as biomethane or bio-natural gas (Alonso-Vicario et al., 2010; Ryckeboch et al., 2011; Yang et al., 2014).

At this point, it is important to mention that CH₄ has a global warming potential around 28 times higher than CO₂ tackling a 100 years base (Myhre et al., 2013). Therefore, the conversion of CH₄ into CO₂ by biogas combustion is by itself very positive for the mitigation of the greenhouse effect. Furthermore, if one takes into account that the burned biogas replaced a fossil fuel, thus this positive effect becomes even higher.

**CO₂ Separation**

The separation of CO₂ is a critical step in both biogas upgrading and CCS/CCU systems, and it is still an economic burden. In the separation of CO₂ from flue gas in CCS/CCU systems, chemical absorption by amine/ammonia solutions (amine/ammonia scrubbing) is nowadays the most employed technology due mainly to the high process efficiency (up to 98%). However, the technique presents some serious shortcomings, which are most related to the high-energy consumption for solvent regeneration and pumping, corrosion of equipment and toxic emissions (Zhao et al., 2012; Rashidi and Yusup, 2016). Concerning biogas upgrading, physical absorption by water under pressure (the so-called pressurized water scrubbing, PWS) is the most used technique for CO₂/CH₄ separation. PWS is a simple and not expansive technique that presents high efficiency and low CH₄ loss, besides permitting the simultaneous removal of H₂S. However, clogging due to bacterial growth, the huge water consumption and low flexibility toward variation of input gas are serious existing drawbacks (Ryckeboch et al., 2011; Nie et al., 2013; Niesner et al., 2013; Xu et al., 2015).

Taking into account the above-mentioned shortcomings related to amine/ammonia scrubbing and PWS, the development of alternative cost-effective technologies to separate CO₂ from flue gas streams and biogas is highly desirable. Such alternatives include cryogenic separation, membrane separation, biological removal, and adsorption (Presser et al., 2011; Leung et al., 2014; Lee and Park, 2015). Amongst them, physical adsorption on solid porous sorbents has been considered the most appealing option because of the low energy required and ease of applicability over a relatively wide range of temperatures and pressures (Chaffee et al., 2007; Shafeeyan et al., 2010; Songolzadeh et al., 2012; Chen et al., 2013; Azmi and Aziz, 2019).

**CO₂ Adsorption on Activated Carbons**

Numerous materials have been investigated for CO₂ adsorption, which includes zeolites, silica, metal-organic frameworks (MOFs), alkali-based and metal oxide-based adsorbents, porous polymers and carbonaceous materials (Songolzadeh et al., 2012; Lee and Park, 2015; Rashidi and Yusup, 2016). Amongst them, activated carbons (ACs) are a promising option (Silvestre-Albero et al., 2011, 2014; Sevilla et al., 2019) because they: present stability in terms of resistance to thermal, mechanical and chemical strength; are safe for the environment; can be easily produced from abundant and cheap raw materials such as coal, biomass and petroleum residues, which is of overwhelming importance in large-scale applications; are hydrophobic, which is important to avoid competitive adsorption of water; can have the pore morphology easily designed by an appropriate choice of activation conditions; have fast adsorption/desorption kinetic; establish weak interactions with CO₂, which makes easy and less energy demanding the discharge process by thermal or pressure modulation (Wang et al., 2013; Parshetti et al., 2015; Li and Xiao, 2019; Singh et al., 2019).

According to the literature, the overall majority of ACs intended for CO₂ adsorption have been prepared by chemical activation with KOH. The reason is that this procedure is renowned for rendering carbons with remarkably high specific surface area and microporosity, so that the resulting adsorbents have usually shown the highest gravimetric CO₂ uptakes, as it can be verified from the data in the Supplementary Table 1. Notwithstanding, we have at least three considerations to make about this issue, as depicted in the sequence.
(i) Firstly, quite large amounts of KOH (a caustic and highly corrosive chemical) are employed in the activation process. Usual KOH:precursor ratio are in the range of 1 to 4, which can be considered unviable for a large-scale production (see, just as few examples, the works of Wahby et al., 2010; Deng et al., 2014; Parshetti et al., 2015; Haffner-Staton et al., 2016; Hong et al., 2016; Huang et al., 2019; Kutorglo et al., 2019; Li et al., 2019).

(ii) Secondly, activation with KOH invariably renders carbons in the powdered form. Therefore, before being used in fixed bed adsorption columns, the material should be extruded or conformed into monoliths. Besides implying additional costs, these procedures usually require the employment of a binder, which obstructs part of the porosity (Jordá-Beneysto et al., 2008; Munusamy et al., 2015).

(iii) Thirdly, the overwhelming majority of authors have only considered the CO$_2$ uptake on a gravimetric basis. However, considering that the adsorbent has to be confined in a given column with limited volume, so it is much more relevant, from an application point of view, to evaluate the data on a volume basis rather than on a mass basis. This issue was firstly considered by Silvestre-Albero et al. (2011); however, as far as we know, since then only Li et al. (2019), Haffner-Staton et al. (2016), and Marco-Lozar et al. (2012) have reported the CO$_2$ uptake of ACs on a volumetric basis.

**Purpose and Scope of This Paper**

Within the above-described scenario, the goal of the present work was to prepare biomass-based granular ACs with the highest volumetric CO$_2$ uptakes as possible (at atmospheric and subatmospheric pressures), but employing activation procedures that are practical and cost-effective for a large-scale production. In this sense, we used an abundant and cheap feedstock: coconut shell, a residual biomass. Furthermore, the adsorbents were prepared in the granular form because it is most practical and inexpensive. One could argue that the interparticle space of granular adsorbents reduces the volumetric adsorption uptake. Of course, it is true; however, it is needed to have in mind that, on the other hand, the existence of some free space is essential to permit adequate gas diffusion throughout a packed column.

As activating methodologies, we investigated in depth the chemical activation with dehydrating agents (H$_3$PO$_4$ or ZnCl$_2$) and the physical activation with CO$_2$. Although there are some scarce works reporting the use of these approaches in the production of ACs intended for CO$_2$ adsorption (Presser et al., 2011; Vargas et al., 2011; Bae and Su, 2013; Ello et al., 2013; Balsamo et al., 2014; Ludwinowicz and Jaroniec, 2015; Munusamy et al., 2015; Ahmed et al., 2019) and the reported gravimetric CO$_2$ uptakes are, in a general way, lower than those verified for KOH activated carbons (compare the data in Supplementary Table 1), none of the existing works concerned a systematic study in which the activation conditions are screened in order to achieve materials with improved performances. Further, the results were always considered only on a mass basis instead of a volume basis.

Finally, it is worthy to highlight that, along the present work, efforts were also directed toward the comprehension of the relationships between the procedures employed to prepare the ACs, the resulting pore morphology and the performance for CO$_2$ adsorption of the obtained materials.

**MATERIALS AND METHODS**

**Activation Procedures**

A sample of dried endocarp of coconut shell (Cocos nucifera) was used as raw material. After crushing and sieving, the fraction in the range of 2.00–2.83 mm was used in all preparations. Some characteristics of the precursor were previously published (Prauchner and Rodríguez-Reinoso, 2012). The carbonization and physical activation procedures were carried out in a horizontal tubular furnace. For carbonization, the samples were heated (2.0°C/min) under N$_2$ flow (100 mL/min) up to the desired temperature (2 h).

For chemical activation, the precursor was first impregnated with an aqueous solution of the chemical (2.0 mL per gram of precursor). The solution concentration was adjusted to provide the desired mass ratio of phosphorous or zinc to the precursor (these ratios will be termed as $X_P$ or $X_Zn$, respectively). The impregnated material was then carbonized up to 450 or 500°C (for activation with H$_3$PO$_4$ or ZnCl$_2$, respectively) and washed to remove the chemical. Some additional details on the chemical activation procedures were previously reported (Prauchner and Rodríguez-Reinoso, 2012).

Concerning the complementary carbonization, it is worthy to highlight that the chemically activated samples were firstly prepared as described above and, in a subsequent and independent step, carbonized again, this time up to 850°C.

For physical activation, the starting material (which had always been previously carbonized up to 850°C) was heated (5°C/min) under N$_2$ flow (100 mL/min). When reached the activation temperature (750°C), the gas flow was switched to CO$_2$ (100 mL/min) and the temperature was kept for the period of time necessary to achieve the desired burn-off (weight loss due to the gasification with CO$_2$).

**Activated Carbon Labels**

The ACs were labeled according to their preparing procedure and the following rules:

a) A carbonization step was indicated by the letter C followed by the maximum reached temperature.

b) For chemical activation, the letter P or Z was used to indicate the chemical (H$_3$PO$_4$ or ZnCl$_2$, respectively), followed by the $X_P$ or $X_Zn$ value multiplied by 100.

c) For physical activation, the letter B was used followed by the respective burn-off (in percentage).

Thus, for example, the Z15.C850.B20 sample concerns the material that was chemically activated with ZnCl$_2$ with a $X_Zn$ of 0.15, followed by carbonization up to 850°C and subsequent physical activation with CO$_2$ up to a burn-off of 20%.

In turn, in order to represent an entire series of samples, the letter “X” was used in the place of the parameter that varied within that series (temperature, chemical loading or burn-off). In this way, for example, the P09.C850.BXX series comprises all the activated carbons produced by chemical activation with
H₂PO₄ with a Xₚ 0.09, followed by carbonization up to 850°C and subsequent gasification with CO₂ up to different burn-offs.

**Activated Carbon Characterization**

The pore morphology was evaluated from the excess adsorption isotherms of N₂ (−196°C) and CO₂ (0°C) recorded up to 1.0 bar in a volumetric automatic system Omnisorb 610. Before any experiment, samples were degassed (10⁻⁴ Pa) at 350°C for 4 h.

The N₂ adsorption isotherms were used to determine the specific surface area (S_BET) and the micropore volume (V_mic) by applying the BET and DR equations, respectively. The volume of liquid N₂ adsorbed at p/p₀ 0.95 was termed V₀.95 and it was considered to be the sum of V_mic and the volume of mesopores (V_mes). Therefore, V_mes was calculated by subtracting V_mic from V₀.95. In turn, the CO₂ isotherms were employed in the calculation of the ultramicropore volume (V_ult) (sometimes also termed narrow micropore volume). Finally, the software Autosorb 1 was used to generate pore size distribution curves from the N₂ and CO₂ adsorption isotherms (PSD-N₂ and PSD-CO₂, respectively) by using the non-linear density functional theory (NLDFT).

The gravimetric CO₂ uptakes were determined from the CO₂ adsorption isotherms acquired as described above. In turn, the volumetric CO₂ uptakes were determined by multiplying the gravimetric uptakes by the sample bulk density (ρₕ). The latter was measured by gently tapping a weighted amount of grains in a graduate cylinder (tap density).

The CO₂ uptakes were taken at 1.0 and 0.15 bar. These pressures were chosen because they mimic typical conditions for CO₂ capture from biogas or flue gas streams. For example, if the concentration of CO₂ in the gas mixture is 15% (a typical value), thus 1.0 bar corresponds to the partial pressure of CO₂ in a pressure swing adsorption unit operating at 6.7 bar. In turn, 0.15 bar corresponds to the partial pressure of CO₂ in a vacuum swing or temperature swing adsorption unit operating at atmospheric pressure.

An automatic pycnometer Accupyc 1330 from Micromeritics was used to measure the helium density (ρₕ), which was considered equivalent to the carbon skeleton density. As previously detailed (Prauchner et al., 2016), the gravimetric waste volume (V_wg) was calculated from the following equation:

\[ V_{wg} = \frac{1}{\rho_h} - V_{0.95} - \frac{1}{\rho_h} \]

Thus, the volumetric waste volume (V_wg), determined by multiplying V_wg by ρₕ, comprises, in a bed filled by the adsorbent, the fraction of volume that corresponds to interparticle spaces plus large pores that are not filled by N₂ at p/p₀ = 0.95 and −196°C.

**RESULTS AND DISCUSSION**

For reasons of space, only some selected isotherms essential for the ongoing discussions are showed in the main text. Nevertheless, the N₂ adsorption/desorption and CO₂ adsorption isotherms of all the prepared samples are available as Supplementary Figures 4–8. The data of pore morphology and density are furnished in Table 1. A detailed discussion about porosity development during physical activation with CO₂ and chemical activation with H₃PO₄ or ZnCl₂ was reported elsewhere (Prauchner and Rodriguez-Reinoso, 2012).

**Relationships Between Pore Morphology and CO₂ Uptake**

Before presenting and discussing the results verified for each series of carbons, the data of all the prepared samples were analyzed as a whole in order to search for relationships between the pore morphology and the CO₂ uptake. These relationships will be valuable for subsequent discussions.

In this sense, the gravimetric CO₂ uptake of all prepared samples were plotted vs. the data of pore morphology. The plots showed no evident relationship between the CO₂ uptake and the parameters determined from the N₂ adsorption isotherms (S_BET, V₀.95, and V_mic; Supplementary Figure 1). On the other hand, it was possible to identify trends of increasing CO₂ uptake with increasing volume of pores determined from the CO₂ adsorption isotherms (V_ult; Supplementary Figure 2). Notwithstanding the low correlation coefficients (R²), especially in the case of adsorption at 0.15 bar, these results disclosed that, as already suggested by several authors (Presser et al., 2011; Wahby et al., 2012; Wei et al., 2012; Casco et al., 2013; Sevilla et al., 2013; Silvestre-Albero et al., 2014; Haffner-Staton et al., 2016; Serafin et al., 2017; Jang et al., 2019; Sreiscek-Nazzal and Kielbasa, 2019), CO₂ adsorption at low pressure occurs mostly in narrow micropores, which can be more accurately probed by CO₂ adsorption experiments. Namely, CO₂ isotherms are usually acquired at much higher temperature (0°C) than N₂ isotherms (−196°C), which drastically reduce the diffusional restrictions. In addition, the CO₂ saturation pressure (p₀) at 0°C is ~35 bar, so that 1.0 bar corresponds to a relative pressure (p/p₀) of only ~0.03. Therefore, it is usually accepted that, at 0°C and atmospheric pressure, CO₂ molecules are significantly adsorbed only in narrow micropores, where the proximity of neighboring walls causes an overlapping of adsorption potentials (Rodriguez-Reinoso and Linares-Solano, 1989). This is the reason why the volume of pores determined by applying the DR equation to CO₂ isotherms at 0°C is considered to correspond to the volume of ultramicropores (d < 0.7 nm, where d is the pore dimension) (Garrido et al., 1987; Rodriguez-Reinoso et al., 1989; Cazorla-Amorós et al., 1996).

The above-mentioned results triggered more in-depth studies concerning the CO₂ adsorption isotherms. In this sense, the gravimetric CO₂ uptake was plotted against the volume of pores smaller than a certain threshold size determined from the PSDs-CO₂ curves: 0.4, 0.5, 0.6, 0.7, 0.8, and 0.9 nm (V<0.4, V<0.5, V<0.6, V<0.7, V<0.8, and V<0.9, respectively). The graphs (Supplementary Figure 3) showed that, for a CO₂ pressure of 1.0 bar, the CO₂ uptake correlates better with the volume of pores smaller than 0.8 nm, with a R² of 0.9573 (Figure 1). In turn, for a CO₂ pressure of 0.15 bar, the higher R² was verified when the data were plotted against V<0.6, with a R² value of 0.8997. These findings permitted to infer that, at 0°C, pores with dimensions in the lower range of ultramicropores are efficiently filled by CO₂.
at pressures as low as 0.15 bar, whereas pores with dimensions in the higher range of ultramicropores need pressures of around 1.0 bar to be effectively filled.

**CO₂ Uptake**

Before going ahead with the discussions, it is worthy to emphasize that the volumetric uptake is determined by multiplying the gravimetric uptake by the adsorbent bulk density and, in a general way, the bulk density decreases with increasing porosity. Therefore, to reach an optimized volumetric uptake, the porosity should be increased in such a controlled way that the density of the adsorbed phase is not significantly reduced. On the otherwise, an eventual increase of the gravimetric CO₂ uptake can be surpassed by the reduction of bulk density, therefore resulting in a lower volumetric CO₂ uptake.

**Carbonization in Absence of Chemicals**

During the carbonization of lignocellulosic feedstocks in absence of chemicals, volatile matter, aliphatic carbons and hetero atoms are released, while residual carbon atoms are grouped into stacks of flat aromatic sheets cross-linked in a random manner. Thus, the irregular arrangement of carbon sheets gives rise to narrow free interstices that correspond to an incipient porosity (Cândido et al., 2020). Indeed, the sample obtained by coconut shell carbonization up to 450°C (C450 sample) presented a $V_{\text{ult}}$ of 0.14 cm$^3$/g (Table 1). The PSD-CO₂ curve (Figure 2B) showed two main peaks corresponding to pore dimensions in the range of $\sim$0.30–0.42 nm and $\sim$0.42–0.72 nm, with maxima at 0.36 and 0.49 nm, respectively. This narrow porosity resulted in gravimetric CO₂ uptakes of 1.98 and 1.03 mmol/g at 1.0 and 0.15 bar, respectively. Taking the sample bulk density (0.575 g/cm$^3$) into account, the respective volumetric uptakes were calculated as 1.14 and 0.59 mmol/cm$^3$. 

---

**TABLE 1 | Data of pore morphology and density.**

| Sample     | $S_{\text{BET}}$ ($m^2$/g) | $V_{\text{ult}}$ ($cm^3$/g) | $V_{\text{mic}}$ ($cm^3$/g) | $V_{\text{mes}}$ ($cm^3$/g) | $V_{0.95}$ ($cm^3$/g) | $\rho_{\text{He}}$ (g/cm$^3$) | $\rho_b$ (g/cm$^3$) |
|------------|----------------------------|------------------------------|----------------------------|-----------------------------|------------------------|-----------------------------|--------------------------|
| C450       | *                          | 0.14                         | *                          | *                           | *                      | 1.42                        | 0.575                    |
| C850       | *                          | 0.26                         | *                          | *                           | *                      | 1.89                        | 0.611                    |
| C850.B25   | 939                        | 0.41                         | 0.43                       | 0.03                        | 0.46                   | 2.01                        | 0.500                    |
| C850.B35   | 1,046                      | 0.45                         | 0.49                       | 0.05                        | 0.54                   | 2.04                        | 0.473                    |
| C850.B52   | 1,340                      | 0.46                         | 0.59                       | 0.09                        | 0.68                   | 2.08                        | 0.416                    |
| C850.B79   | 1,987                      | 0.49                         | 0.78                       | 0.23                        | 1.01                   | 2.14                        | 0.323                    |
| C850.B94   | 2,276                      | 0.48                         | 0.81                       | 0.41                        | 1.22                   | 2.19                        | 0.270                    |
| P09        | 869                        | 0.34                         | 0.37                       | 0.03                        | 0.40                   | 1.59                        | 0.536                    |
| P15        | 1,218                      | 0.35                         | 0.46                       | 0.14                        | 0.60                   | 1.66                        | 0.522                    |
| P21        | 1,294                      | 0.29                         | 0.50                       | 0.12                        | 0.62                   | 1.67                        | 0.501                    |
| P27        | 1,620                      | 0.33                         | 0.59                       | 0.21                        | 0.80                   | 1.71                        | 0.458                    |
| P30        | 1,826                      | 0.30                         | 0.62                       | 0.35                        | 0.97                   | 1.73                        | 0.422                    |
| P36        | 2,017                      | 0.37                         | 0.68                       | 0.41                        | 1.09                   | 1.76                        | 0.347                    |
| P54        | 2,212                      | 0.35                         | 0.73                       | 0.66                        | 1.39                   | 1.79                        | 0.281                    |
| Z15        | 719                        | 0.32                         | 0.32                       | 0.01                        | 0.33                   | 1.58                        | 0.544                    |
| Z25        | 1,293                      | 0.37                         | 0.52                       | 0.07                        | 0.59                   | 1.62                        | 0.519                    |
| Z32        | 1,392                      | 0.36                         | 0.55                       | 0.08                        | 0.63                   | 1.65                        | 0.507                    |
| Z40        | 1,700                      | 0.36                         | 0.66                       | 0.14                        | 0.80                   | 1.65                        | 0.445                    |
| Z50        | 1,960                      | 0.38                         | 0.71                       | 0.27                        | 0.98                   | 1.69                        | 0.405                    |
| Z65        | 1,978                      | 0.33                         | 0.73                       | 0.37                        | 1.10                   | 1.69                        | 0.330                    |
| P09.C850   | 665                        | 0.26                         | 0.27                       | 0.03                        | 0.30                   | 1.97                        | 0.680                    |
| P09.C850.B22 | 1,143                   | 0.41                         | 0.48                       | 0.03                        | 0.51                   | 2.04                        | 0.554                    |
| P09.C850.B34 | 1,470                   | 0.43                         | 0.59                       | 0.07                        | 0.66                   | 2.12                        | 0.491                    |
| P09.C850.B44 | 1,945                   | 0.54                         | 0.69                       | 0.11                        | 0.80                   | 2.15                        | 0.445                    |
| P09.C850.B54 | 2,106                   | 0.50                         | 0.76                       | 0.22                        | 0.98                   | 2.17                        | 0.393                    |
| Z15.C850   | 679                        | 0.23                         | 0.31                       | 0.00                        | 0.31                   | 1.95                        | 0.660                    |
| Z15.C850.B20 | 1,096                   | 0.40                         | 0.46                       | 0.00                        | 0.46                   | 2.05                        | 0.583                    |
| Z15.C850.B29 | 1,211                   | 0.48                         | 0.54                       | 0.03                        | 0.57                   | 2.11                        | 0.524                    |
| Z15.C850.B36 | 1,489                   | 0.53                         | 0.65                       | 0.03                        | 0.68                   | 2.14                        | 0.485                    |
| Z15.C850.B44 | 1,599                   | 0.54                         | 0.70                       | 0.07                        | 0.77                   | 2.15                        | 0.438                    |

*Not determined because the quite narrow porosity of the C450 and C850 samples hindered, at $\sim$196°C, the access of the N₂ molecules at a period of time plausible for the acquirement of the isotherms.
It is well-known that, during carbonization, pronounced condensation of aromatic rings takes place between around 600 and 800°C, thus resulting in a porosity increase (Prauchner et al., 2005). Indeed, $V_{\text{ult}}$ enhanced from 0.14 to 0.26 cm$^3$/g when the carbonization temperature increased from 450 to 850°C (Table 1). Consequently, if compared with the C450 sample, the CO$_2$ adsorption isotherm of the C850 sample presented higher CO$_2$ uptakes across the entire range of pressures (Figure 2A). At 1.0 and 0.15 bar, the gravimetric CO$_2$ uptake increased from 1.98 to 3.47 mmol/g and from 1.03 to 1.64 mmol/g, respectively. Taking the bulk densities into account (which was 0.611 g/cm$^3$ for the C850 sample), it was determined that the respective volumetric uptakes increased from 1.14 and 0.59 mmol/cm$^3$ to 2.12 and 1.00 mmol/cm$^3$.

Concerning the PSD-CO$_2$ curve (Figure 2B), C850 also presented two peaks at nearly the same positions as verified for C450, being that the area of the peak with the maximum at $\sim 0.49$ nm was pronouncedly higher for the C850 sample. Once the position of the peak did not change, it is possible to conclude that the porosity increase verified when the carbonization temperature increased from 450 to 850°C was due to the creation of new pores rather than by the growth of the existing ones.

Physical Activation

During physical activation with CO$_2$, the activating gas diffuses throughout the incipient porosity formed during the precursor carbonization and gasifies the walls accordingly to the following reaction: CO$_2$ + C $\rightarrow$ 2CO (Marsh and Rodriguez-Reinoso, 2006). Indeed, Table 1 shows that the porosity gradually increased with increasing burn-off and, consequently, the bulk density decreased.

Even though the volume of micropores plus mesopores ($V_{0.95}$) continuously increased, the volume of ultramicropores ($V_{\text{ult}}$) first increased but few changed after a burn-off of around 35% (Table 1). Therefore, taking into account that ultramicropores are more effective for CO$_2$ adsorption at atmospheric pressure, the gravimetric CO$_2$ uptake at 1.0 bar reached a maximum for the C850.B35 sample, 5.0 mmol/g, and then nearly stabilized (Figure 3A).

The isotherms of CO$_2$ adsorption and corresponding PSD curves of the C850 and C850.B35 samples (Figures 4A,B) disclose that, up to moderate burn-offs, the increase of porosity took place in both the lower and the higher range of ultramicropores. However, when the gasification was more intense, the volume of pores located in the lower range of ultramicropores diminished, as portrayed by the PSD-CO$_2$ curve of the C850.B79 sample. This supposedly occurred because the dimensions of these small pores enhanced along the gasification process, so that they partially moved toward the higher range of ultramicropores. Thanks to this behavior and the fact that CO$_2$ adsorption at low subatmospheric pressures is more efficient at small ultramicropores, the gravimetric CO$_2$ uptake at 0.15 bar reached a maximum for the B35 sample, $\sim$1.95 mmol/g, and decreased thereafter (Figure 3B).

Concerning the volumetric uptake, different behaviors were observed at 1.0 and 0.15 bar. As Table 1 discloses, the porosity development verified up to a burn-off of around 35% was basically due to increases in the volume of ultramicropores, so that almost all of the gain in porosity was effective for increasing the CO$_2$ adsorption at 1.0 bar. Therefore, within this burn-off range, the increase of gravimetric CO$_2$ uptake at 1.0 bar surpassed the reduction of bulk density and, consequently, the volumetric CO$_2$ uptake reached a maximum ($\sim$2.40 mmol/cm$^3$) for burn-offs between 25 and 35% (Figure 3A). On the other hand, only the pores located in the lower range of ultramicropores are effective for adsorbing CO$_2$ at 0.15 bar. Therefore, even for low activation degrees, the increase in the gravimetric CO$_2$ uptake at this pressure was not sufficient to compensate the reduction of bulk density. Consequently, all the physically activated samples had volumetric CO$_2$ uptake at 0.15 bar lower than that of the only carbonized sample C850 (which was 1.00 mmol/cm$^3$) (Figure 3B). Furthermore, the uptake steadily decreased with increasing burn-off.

Chemical Activation

During the carbonization of lignocellulosic precursors impregnated with H$_3$PO$_4$ or ZnCl$_2$, the carbon matrix develops around the chemicals, which leaves the structure in an expanded state. Thus, the subsequent removal of the chemical by leaching makes a pore structure available (Prauchner and Rodriguez-Reinoso, 2012). Indeed, Table 1 shows that chemical activation with H$_3$PO$_4$ or ZnCl$_2$ rendered carbons with higher porosity and lower bulk density than the sample carbonized at similar temperature in absence of chemicals (C450 sample). Further, the higher the chemical loading, the higher the $V_{0.95}$ and the lower the bulk density.

In comparison with the C450 sample, the higher porosity of the samples chemically activated with the lowest loadings (P09 and Z15 samples) was primarily due to their higher volume of ultramicropores (compare the data in Table 1). In turn, for higher loadings, $V_{\text{ult}}$ oscillated without a defined trend, while the volume
FIGURE 2 | (A) CO$_2$ adsorption isotherms and (B) corresponding PSDs-CO$_2$ curves of the samples carbonized up to 450 and 850°C in absence of chemicals.

FIGURE 3 | Gravimetric and volumetric CO$_2$ uptakes at (A) 1.0 and (B) 0.15 bar as a function of the burn-off during physical activation with CO$_2$.

FIGURE 4 | (A) CO$_2$ adsorption isotherms and (B) corresponding PSDs-CO$_2$ curves of the carbon obtained by carbonization up to 850°C and some selected samples physically activated with CO$_2$. 
of larger pores increased. Therefore, since CO$_2$ adsorption at atmospheric pressure takes place mostly at ultramicropores, the gravimetric CO$_2$ uptake at 1.0 bar was abruptly higher for the P09 and Z15 samples (3.2 and 3.3 mmol/g, respectively) if compared with the C450 sample (1.98 mmol/g), and slightly decreased for higher chemical loadings (Figures 5A,C).

Conversely to what was observed for a CO$_2$ pressure of 1.0 bar, the chemical activation with low chemical loadings did not promote considerable increases in the gravimetric CO$_2$ uptake at 0.15 bar (Figures 5B,D). Instead of that, the uptake for the Z15 sample was only marginally higher than the value verified for the C450 sample, while the uptake of the P09 sample was even lower. In addition, the uptake pronouncedly decreased with increasing chemical loading. This profile of behavior can be explained based on the isotherms of CO$_2$ adsorption of the chemically activated samples (Figures 6A,C) and respective PSD curves (Figures 6B,D). The curves show that, if compared with carbonization in absence of chemicals, chemical activation provoked a reduction in the volume of pores below ~0.42 nm; the higher the chemical loading, the more intense was this reduction. Thus, since pores located in the lower range of ultramicropores are responsible for most of the CO$_2$ adsorption at low subatmospheric pressures, the reduction in the volume of pores below 0.42 nm diminished the CO$_2$ uptake at 0.15 bar.

The above-mentioned reduction in the volume of pores below ~0.42 nm can be explained as it follows. As already discussed in the Subsection “Carbonization in absence of chemicals”, the irregular arrangement of carbon sheets during the carbonization of non-impregnated biomass generates free interstices that correspond to pores in the lower range of ultramicropores. In turn, if the carbonization is performed in presence of dehydrating agents such as H$_3$PO$_4$ or ZnCl$_2$, the chemical decomposes the lignocellulosic chains and the resulting fragments reorganize and give rise to a more ordered carbon structure. Therefore, at the same time that the chemical acts as a physical template for the development of larger pores, the material reorganization supposedly reduces the occurrence of free interstices that correspond to small ultramicropores.

**Comparing the Activations With CO$_2$, H$_3$PO$_4$, and ZnCl$_2$**

The comparison of the results verified for the samples prepared by the different methodologies shows that the carbons physically activated with CO$_2$ reached higher CO$_2$ uptakes than the chemically activated samples, whether at 1.0 or 0.15 bar, whether
on a mass or a volume basis. This behavior is disclosed by Figure 7, in which the highest CO$_2$ uptake reached for each series of samples is plotted in a bar graph. Furthermore, in the case of the gravimetric uptake, the Supplementary Figure 1 makes clear that the physically activated carbons (C850.BXX series) exhibited always higher performances than the chemically activated carbons (PXX and ZXX series). These results can be mainly attributed to the higher volume of ultramicropores reached by the physically activated carbons in both the lower and higher range of ultramicropores, as portrayed by the PSD-CO$_2$ curves in Figure 8. At this point, it is valid to emphasize that the chemical procedures did not permit to increase considerably the volume of ultramicropores because the use of higher chemical loadings leads only to increases in the volumes of supermicropores or even mesopores.

In the specific case of the volumetric uptake, another factor positively affects the performance of the physically activated carbons: their higher skeleton density. Figure 9 portrays that all the carbons that underwent carbonization at 850°C (as it was the case of the physically activated samples) presented higher helium density than the samples that were treated up to only 450 or 500°C (as it was the case of the chemically activated samples). This occurs because temperatures in the range of ~600–800°C promote extensive aromatization, which increases the skeleton density (Prauchner et al., 2005).

Complementary Carbonization and Physical Activation of Chemically Activated Carbons

As mentioned in the previous Subsection, the carbons prepared by chemical activation with H$_3$PO$_4$ or ZnCl$_2$ had lower CO$_2$ uptakes than those prepared by physical activation with CO$_2$. On the other hand, we have shown elsewhere (Prauchner and Rodriguez-Reinoso, 2012; Prauchner et al., 2016) that chemical activation with H$_3$PO$_4$ or ZnCl$_2$ permits to suppress the formation of large macropores that are present in physically activated carbons produced from lignocellulosic feedstocks. These macropores, which correspond to empty spaces that originate from conductor vessels present in the botanical structure of the precursor, few contribute for adsorption and reduce the material bulk density, thus reducing the
volumetric adsorption capacity. However, the acid attack promoted by H₃PO₄ and ZnCl₂ degrades the lignocellulosic structure during early carbonization stages, so that the resulting fragments have sufficient mobility to promote a material reorganization and redistribution. As mentioned in the previous Subsection, the reorganization suppresses the formation of small ultramicropores that correspond to free interstices generated during the carbonization process. In turn, the redistribution promotes the filling of existing empty spaces that, otherwise, would result in the formation of macropores.

In this context, we decided to research deeper the feasibility of using chemical activation to prepare ACs with improved volumetric CO₂ adsorption capacities. We first investigated the possibility of raising the performance of chemically activated carbons by subjecting them to complementary carbonization up to 850°C. For this purpose, we choose the samples activated with the lower chemical loadings (P09 and Z15) because, as already discussed, amongst the chemically activated samples, they presented the more appropriate porosity for CO₂ adsorption. Afterward, we investigated the possibility of optimizing the porosity of the materials obtained from the complementary carbonization by means of subsequent physical activation with CO₂.

**Complementary Carbonization of Chemically Activated Carbons**

Since during chemical activation the materials were treated only up to moderate temperatures (450 or 500°C), complementary carbonization of the P09 and Z15 samples up to 850°C led to additional weight loss of around 8%. This weight loss shrank the particles, thus causing porosity contraction. Therefore, there was considerable reduction in the volume of pores that were accessible to N₂ at −196°C. As disclosed by the PSD-N₂ curves in Figure 10, these pores were mostly situated in the range of supermicropores (0.7 nm < d < 2.0 nm). Furthermore, there was also some reduction in the volume of pores located in the higher range of ultramicropores, as portrayed by the PSD-CO₂ curves in Figure 11B. On the other hand, this same Figure evidences that the contraction of larger pores ended by increasing the volume of pores located in the lower range of ultramicropores. This effect was remarkably pronounced in the case of the Z15.C850 sample, whose PSD-CO₂ curve exhibited a somewhat sharp peak centered at around 0.47 nm.
The comparison of the CO\textsubscript{2} adsorption isotherms of the Z15.C850 and P09.C850 samples with those of the Z15 and P09 samples (Figure 11A) reveals that complementary carbonization increased the gravimetric CO\textsubscript{2} uptake. In line with the changes in porosity reported in the previous paragraph, the increases were more pronounced at lower pressures: while at 1.0 bar the uptake increased only from 3.18 and 3.28 mmol/g for the P09 and Z15 samples to 3.31 and 3.48 mmol/g for the P09.C850 and Z15.C850 samples (increases of around 4 and 8%, respectively), at 0.15 bar the increases were much higher, from 0.95 and 1.11 mmol/g to 1.40 and 1.78 mmol/g (increases of around 47 and 60%, respectively).

Besides increasing the gravimetric CO\textsubscript{2} uptake, complementary carbonization considerably increased the bulk density: the increases were from 0.536 and 0.544 g/cm\textsuperscript{3} for the P09 and Z15 samples to 0.680 and 0.660 g/cm\textsuperscript{3} for the P09.C850 and Z15.C850 samples (Table 1). This increase can be attributed to two effects: (i) the porosity contraction; (ii) the increase of skeleton density (see Figure 9 and related discussion).

Since both the gravimetric CO\textsubscript{2} uptake and bulk density increased with complementary carbonization, the volumetric uptake also increased: at 1.0 bar, it increased from 1.70 and 1.78 mmol/cm\textsuperscript{3} for the P09 and Z15 samples to 2.25 and 2.30 mmol/cm\textsuperscript{3} for the P09.C850 and Z15.C850 samples; at 0.15 bar, the respective increases were from 0.51 and 0.60 mmol/cm\textsuperscript{3} to 0.95 and 1.17 mmol/cm\textsuperscript{3}.

**Physical Activation of Chemically Activated Carbons**

As disclosed by the $V_{0.95}$ values in Table 1, physical activation with CO\textsubscript{2} permitted to develop the porosity of the P09.C850 and Z15.C850 samples. Regarding the ultramicropores, Figure 12 shows that the volume of pores in the higher range of ultramicropores (higher than nearly 0.5 nm) pronouncedly increased up to a burn-off of around 35% and, from that point on, it stopped to increase and even slightly diminished. Therefore, the gravimetric CO\textsubscript{2} uptake at 1.0 bar passed through a maximum at a burn-off of nearly 35% and decreased thereafter. The maxima were $\sim 4.80$ and $4.99$ mmol/g for the P09.C850.BXX and Z15.C850.BXX series, respectively (Figures 13A,C).

Maximum values were also observed for the volumetric CO\textsubscript{2} uptake at 1.0 bar (Figures 13A,C). However, thanks to the effect of the decreasing bulk density, the burn-off required to achieve the maximum volumetric CO\textsubscript{2} uptake at 1.0 bar ($\sim 20\%$) was lower than that needed to attain the highest gravimetric value ($\sim 35\%$). This phenomenon shows that, even though the volume of ultramicropores and the gravimetric CO\textsubscript{2} uptake keep increasing throughout the burn-off range of around 20–35%, the density of the adsorbed phase diminishes, so that the volumetric uptake decreases. In the case of the P09.C850.BXX series, the maximum (2.40 mmol/cm\textsuperscript{3} for the P09.C850.B22 sample) was only slightly higher than that of the correspondent non-gasified sample P09.C850 (2.25 mmol/cm\textsuperscript{3}). However, in the case of the Z15.C850.BXX series, significant gain was achieved: the volumetric uptake increased from 2.30 mmol/cm\textsuperscript{3} for the non-gasified carbon Z15.C850 to 2.67 mmol/cm\textsuperscript{3} for the Z15.C850.B20 sample. This result can be attributed to the already mentioned narrow size distribution of ultramicropores presented by the Z15.C850 sample, which could be carefully tailored by gasification in order to render an AC with optimized pore size distribution for CO\textsubscript{2} adsorption.

The volumetric uptake at 1.0 bar verified for the Z15.C850.B20 sample (2.67 mmol/cm\textsuperscript{3}) was considerably higher than that observed for the C850.B25 sample (2.34 mmol/g), the one that presented the maximum value in the series prepared by physical activation of the carbons that were not submitted to a previous chemical activation. However, it is interesting to note that the Z15.C850.B20 sample presented an even somewhat lower gravimetric uptake at 1.0 bar (4.58 mmol/g) than the C850.B25 sample (4.68 mmol/g), so that the higher volumetric uptake of the former can be attributed to its higher bulk density. In turn, since the samples Z15.C850.B20 and C850.B25 presented similar $V_{0.95}$ (0.464 and 0.455 cm\textsuperscript{3}/g, respectively) and helium density (2.06 and 2.04 g/cm\textsuperscript{3}), thus it is possible to infer that the higher bulk density of the Z15.C850.B20 sample is due to its lower waste volume. Indeed, Figure 14 shows that the samples that underwent chemical activation with ZnCl\textsubscript{2} or H\textsubscript{3}PO\textsubscript{4} (those pertaining to the PXX, ZXX, P09.C850.BXX and Z15.C850.BXX series, besides the samples P09.C850 and Z15.C850) had lower waste volume than the samples that were heat treated only in absence of chemicals (samples pertaining to the C.850.BXX series, besides the samples C450 and C850). As already discussed along the text, such lower waste volume can be attributed to the action of the chemical during the early stages of carbonization, which prevents the occurrence of macropores.

If by one hand the gasification with CO\textsubscript{2} increased the volume of pores in the higher range of ultramicropores, on the other hand it decrease the volume of pores in the lower range (Figure 12). In the case of the samples that were previously chemically activated with H\textsubscript{3}PO\textsubscript{4} (P09.C850.BXX series), this decrease was restricted to the pores smaller than $\sim 0.4$ nm, so that it did not considerably affect the capacity of adsorbing CO\textsubscript{2}.
FIGURE 11 | (A) CO$_2$ adsorption isotherms and corresponding (B) PSDs-CO$_2$ curves of the chemically activated samples P09 and Z15 and the samples resulting from their complementary carbonization up to 850$^{\circ}$C.

Therefore, the gravimetric CO$_2$ uptake at 0.15 bar somewhat increased with low burn-offs, being that a maximum value was verified for the P09.C850.B22 sample (0.88 mmol/cm$^3$; Figure 13B). In turn, in the case of the samples that were previously chemically activated with ZnCl$_2$, the volume of pores pronouncedly decreased in the range of nearly 0.42–0.51 nm. Thus, all ACs pertaining to the Z15.C850.BXX series had gravimetric CO$_2$ uptake lower than the corresponding non-gasified Z15.C850 sample (Figure 13D).

Thanks to the results verified for the gravimetric CO$_2$ uptake at 0.15 bar and the fact that the bulk density decreased with increasing burn-off, all the gasified samples pertaining to the P09.C850.BXX and Z15.C850.BXX series had lower volumetric CO$_2$ uptake than the corresponding non-gasified P09.C850 and Z15.C850 samples. Furthermore, the volumetric uptake steadily decreased with increasing burn-off.

FIGURE 12 | PSDs-CO$_2$ curves of carbons produced through complementary carbonization of the chemically activated samples (A) P09 and (B) Z15, besides some selected samples obtained by subsequent physical activation with CO$_2$.

Final Considerations
The highest volumetric CO$_2$ uptakes attained in the present work were 2.67 and 1.17 mmol/cm$^3$ for CO$_2$ pressures of 1.0 and 0.15 bar. These values were verified for the Z15.C850.B20 and Z15.C850 samples, respectively. It is noteworthy that the preparation of both these samples involved a preliminary chemical activation with a low ZnCl$_2$ loading, followed by complementary carbonization up to 850$^{\circ}$C. In the case of the sample Z15.C850.B20, a soft gasification with CO$_2$ was also employed. These results and the discussions along the text permitted to make important insights about the way the processing methodology influences the characteristics of the resulting adsorbents and their performance for CO$_2$ adsorption. Firstly, a chemical activation step (with H$_3$PO$_4$ or ZnCl$_2$) permits to suppress the occurrence of macropores, what is beneficial for the bulk density. Nevertheless, it is needed to...
mention that a low chemical loading must be used because, otherwise, the pores widen and, therefore, become less efficient for CO$_2$ adsorption at low pressures. Secondly, it is important to submit the material to a subsequent thermal treatment up to relatively high temperatures (like 850°C) in order to increase the material skeleton density and reduce the average size of the pores created by chemical activation. Finally, if compared with H$_3$PO$_4$, ZnCl$_2$ is a more appropriate chemical agent because it renders a more homogeneous distribution of ultramicropores. After carbonization at 850°C, these pores become appropriate for CO$_2$ adsorption at low subatmospheric pressures (e.g., 0.15 bar). In turn, for adsorption at atmospheric pressure, a subsequent soft gasification with CO$_2$ makes it possible to enhance the pores in such a controlled way that improved volumetric CO$_2$ uptakes can be attained.

If taken on a gravimetric basis, the CO$_2$ uptakes obtained in the present work were lower than those usually reported in the literature for ACs prepared by chemical activation with KOH. For example, the highest values we attained at 1.0 and 0.15 bar were \(\sim5.00\) and \(2.00\) mmol/g, respectively, whereas the corresponding values reported by Li et al. (2019) for petroleum coke-based ACs were \(6.47\) and \(2.52\) mmol/g. Notwithstanding, if a volumetric basis is taken into account, the differences disappear. Namely, the highest volumetric CO$_2$ uptakes verified in the present work, \(2.67\) and \(1.17\) mmol/cm$^3$, were quite similar to those verified by the team of Li, \(2.64\) and \(1.18\) mmol/cm$^3$ (the authors considered the bulk density as the tap density measured for the powdered AC with particle size 50–74 µm).
At this point, it is worthy to highlight that the adsorbents prepared by Li et al. (2019) were chemically activated with a high proportion of a caustic and corrosive chemical agent (the employed KOH:precursor ratio was 1.5), whereas the samples that displayed the highest volumetric uptakes in our work were activated with a softer chemical, ZnCl₂, with a Zn:precursor ratio of only 0.15. Further, the ACs prepared by the team of Li were in the form of a fine powder, which is unpractical for fixed bed adsorption columns, whereas our ACs were prepared as grains with high mechanical resistance, which makes them very appropriate adsorbents for this kind of application.

In addition, it is also valid to mention that the interparticle space of granular carbons can be reduced by providing and adequate mix of samples with different granulometries, in which smaller grains partially occupy the free space among the larger grains (Prauchner and Rodríguez-Reinoso, 2008). Therefore, we foresee that it is possible to improve by around 8–10% the maximum volumetric adsorption capacities we reported in the present work.

As already mentioned in the Introduction Section, there are in the literature three other papers that reported the CO₂ adsorption capacity of ACs on a volumetric basis, the papers of Silvestre-Albero et al. (2011), Marco-Lozar et al. (2012), and Haffner-Staton et al. (2016). However, the comparison of the results attained in our work with those reported by these authors is hampered because they: (i) measured the CO₂ uptake at a different temperature, 25°C; (ii) used different approaches to determine the bulk volume. For example, the teams of Silvestre-Albero and Haffner-Staton measured the bulk volume by compacting the powered ACs at high pressures, what does not correspond to a realistic situation at an operating fixed bed adsorption column. In turn, Marco-Lozar et al. considered the bulk volume as the geometric volume of the monoliths. However, the resulting values cannot also be considered realistic because it is unfeasible to have a large scale adsorption column completely filled by monoliths, because the gas diffusion would be precluded. Therefore, it is possible to state that in both works the authors overestimated the bulk densities and, therefore, the volumetric adsorption capacities.

CONCLUSIONS

Granular ACs intended for CO₂ capture at atmospheric and subatmospheric pressures were prepared from an abundant and low-cost biomass residue by using practical and cost-effective procedures, which is essential for large scale applications such as CO₂ capture and biogas upgrading. By the first time, parameters involved in the chemical activation with dehydrating agents (H₃PO₄ or ZnCl₂) and/or physical activation with CO₂ were systematically screened in depth in order to obtain materials with improved performance for CO₂ capture. The prepared ACs exhibited a blend of relatively high gravimetric CO₂ uptake and bulk density, so that high volumetric adsorption capacities were attained. The highest values were 2.67 and 1.17 mmol/cm² for CO₂ pressures of 1.0 and 0.15 bar, respectively. They were verified for the samples that were first chemically activated with ZnCl₂, followed by carbonization up to 850°C; in the case of the pressure of 1.0 bar, a subsequent soft gasification with CO₂ was also employed to optimize the pore size distribution.

Remarkably, these results were similar to those reported by other authors for carbons chemically activated with KOH, the activation methodology that has been widely claimed as the one that produce ACs with the best performances for CO₂ adsorption, but which involves severe restrictions such as the need of using very large proportions of a caustic and corrosive chemical and the fact that the ACs are obtained in the powdered form. Furthermore, it is possible to foresee that the volumetric results reported in the present work can be additionally improved by using a mix of samples with different granulometries, which would permit to reduce the interparticle space. Therefore, the present work can be considered a very important step in paving the way toward making CO₂ adsorption at each time more interesting technology to reduce the emissions of anthropogenic greenhouse gases.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Materials, further inquiries can be directed to the corresponding author/s.

AUTHOR CONTRIBUTIONS

MP and FR-R conceived the study. MP developed most of the experimental work and wrote the first draft of the manuscript. SO assisted MP in data treatment and manuscript writing. All authors read and approved the final manuscript.

A TRIBUTE TO FRANCISCO RODRÍGUEZ-REINOSO (1941-2020)

Some time after the submission of this paper, we were shocked about the death of Professor Francisco Rodríguez-Reinoso (on August 25th). I am not able to describe the profound sadness that the news provoked in me. He was a brilliant scientist in the area of activated carbons, with a world-wide reputation. However, much beyond a great scientist, professor Rodríguez-Reinoso was also a great person, a precious friend and a great family man. His teachings will certainly remain forever, influencing all those who, like me, had the privilege of having him as professor, supervisor and friend. Dear professor Rodríguez-Reinoso, thank you very much for all you have provided to me and other hundreds of scientists all over the world.

ACKNOWLEDGMENTS

The authors thank the Fundação de Amparo à Pesquisa do Distrito Federal (FAPDF, Brazil; grant numbers 0193.000719/2016 and 0193.001613/2017) and the Coordenação...
The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fchem.2020.581133/full#supplementary-material
Myhre, G., Shindell, D., Bréon, F.-M., Collins, W., Fuglestvedt, J., Huang, J., et al. (2013). “Anthropogenic and natural radiative forcing”, in: Climate Change 2013: The Physical Science Basis, eds T. F. Stocker, D. D. Qin, G.-K. Plattner, M. Tignor, S. K. Allen, J. Boschung, A. Nauels, Y. Xia, V. Bex, and P. M. Midgley (Cambridge: Cambridge University Press), 659—740.

Nie, H., Jiang, H., Chong, D., Wu, Q., Xu, C., and Zhou, H. (2013). Comparison of water scrubbing and propylene carbonate adsorption for biogas upgrading process. Energy Fuels 27, 3329—3345. doi: 10.1021/ef400233w

Niesner, J., Jecha, D., and Stehlík, P. (2013). Biogas upgrading technologies: state of art review in European region. Chem. Eng. Trans. 35, 517–522. doi: 10.3303/CET1335086

Parshetti, G. K., Chowdhury, S., and Balasubramanian, R. (2015). Biomass derived low-cost microporous adsorbents for efficient CO$_2$ capture. Fuel 148, 245—254. doi: 10.1016/j.fuel.2015.01.032

Prachner, M. J., Pasa, V. M. D., Molhallem, N. D. S., Otani, C., Otani, S., and Pardini, L. C. (2005). Structural evolution of Eucalyptus tar pitch-based carbons during carbonization. Biomass Bioenergy, 28, 53–61. doi: 10.1016/j.biombioe.2004.05.004

Prachner, M. J., and Rodríguez-Reinoso, F. (2008). Preparation of granular low-cost microporous adsorbents for efficient CO$_2$ capture. Fuel, 87, 146–154. doi: 10.1016/j.fuel.2007.06.017

Prachner, M. J., and Rodríguez-Reinoso, F. (2012). Chemical versus physical transformation of biogas to biomethane. Energy Fuels, 26, 4323–4328. doi: 10.1021/ef200861a

Presser, V., Mc Donough, J., Yeon, S.-H., and Gogotsi, Y. (2011). Effect of pore size on carbon dioxide sorption by carbide derived carbons. Energy Environ. Sci. 4, 3059—3066. doi: 10.1039/c1ee00117f

Rashidi, N. A., and Yusup, S. (2016). An overview of activated carbons utilization for the post-combustion carbon dioxide capture. J. CO$_2$ Util. 13, 1—16. doi: 10.1016/j.jcou.2015.10.002

Rodríguez-Reinoso, F., Garrido, J., Martín-Martínez, J. M., Molina-Sabio, M., and Torregrosa, R. (1989). The combined use of different approaches in the characterization of microporous carbons. Carbon 27, 23—32. doi: 10.1016/0008-6223(89)90153-X

Rodríguez-Reinoso, F., and Linares-Solano, A. (1989). “Microporous structure of activated carbons as revealed by adsorption methods.” in Chemistry and Physics of Carbon, ed P. A. Thrower (New York, NY: Marcel Dekker), 1–146.

Ryckebosch, E., Drouillon, M., and Vervaeren, H. (2011). Techniques for transformation of biogas to biomethane. Biomass Bioenergy, 35, 1633—1645. doi: 10.1016/j.biombioe.2011.02.033

Serafín, J., Narkiewicz, U., Morawski, A. W., Wrobel, R. J., and Michalkiewicz, B. (2017). Highly microporous activated carbons from biomass for CO$_2$ capture and effective micropores at different conditions. J. CO$_2$ Util. 18, 73—79. doi: 10.1016/j.jcou.2017.01.006

Sevilla, M., Ferrero, G. A., and Fuertes, A. B. (2019). “CO$_2$ storage on nanoporous”, in Nanoporous Materials for Gas Storage, Green Energy and Technology, eds K. Kaneko, F. Rodríguez-Reinoso (Singapore: Springer), 287—330.

Sevilla, M., Parrá, J. B., and Fuertes, A. B. (2013). Assessment of the role of micropore size and N-doping in CO$_2$ capture by porous carbons. Appl. Mater. Interfaces 5, 6360—6368. doi: 10.1021/am401423b

Shafeeyan, M. S., Daud, W. M. A. W., Houshmand, A., and Shamiri, A. (2010). A review on surface modification of activated carbon for carbon dioxide adsorption. J. Anal. Appl. Pyrol. 89, 143—151. doi: 10.1016/j.jaap.2010.07.006

Silvestre-Albero, A., Silvestre-Albero, J., Martínez-Escandell, M., and Rodríguez-Reinoso, F. (2014). Micro/mesoporous activated carbons derived from polyaniline: promising candidates for CO$_2$ adsorption. Ind. Eng. Chem. Res. 53, 15398—15405. doi: 10.1021/ie5013129

Silvestre-Albero, J., Wahby, A., Sepúlveda-Escribano, A., Martínez-Escandell, M., Kaneko, K., and Rodríguez-Reinoso, F. (2011). Ultrahigh CO$_2$ adsorption capacity on carbon molecular sieves at room temperature. Chem. Comm. 47, 6840–6842. doi: 10.1039/c1cc16118e

Singh, J., Basu, S., and Bhunia, H. (2019). CO$_2$ capture by modified porous carbon adsorbents: effect of various activating agents. J. Taiwan Inst. Chem. Eng. 102, 438—447. doi: 10.1016/j.jtice.2019.06.011

Songolzadeh, M., Ravanchi, M. T., and Soleimani, M. (2012). Carbon dioxide capture and storage: a general review on adsorbents. Int. J. Chem. Mol. Eng. 6, 900—907.

Srećscek-Nazhal, I., and Kielbasa, K. (2019). Advances in modification of commercial activated carbon for enhancement of CO$_2$ capture. Appl. Surf. Sci. 494, 137—151. doi: 10.1016/j.apsusc.2019.07.108

Vargas, D. P., Giraldo, L., Silvestre-Albero, J., and Moreno-Piraján, J. C. (2011). CO$_2$ adsorption on binderless activated carbon monoliths. Adsorption 17, 497—504. doi: 10.1007/s10450-010-9309-z

Wahby, A., Ramos-Fernández, J. M., Martínez-Escandell, M., Sepúlveda-Escribano, A., Silvestre-Albero, J., and Rodríguez-Reinoso, F. (2010). High-surface-area carbon molecular sieves for selective CO$_2$ adsorption. ChemSusChem 3, 974—981. doi: 10.1002/cssc.20100083

Wahby, A., Silvestre-Albero, J., Sepúlveda-Escribano, A., and Rodríguez-Reinoso, F. (2012). CO$_2$ adsorption on carbon molecular sieves. Micropor. Mesopor. Mat. 164, 280—287. doi: 10.1016/j.micromeso.2012.06.034

Wang, J., Senkovska, I., Otschalt, M., Lohe, M. R., Borchardt, L., Heerwig, A., et al. (2013). Highly porous nitrogen-doped polyimine-based carbons with adjustable microstructures for CO$_2$ capture. J. Mater. Chem. A 1, 10951—10961. doi: 10.1039/c3ta11995e

Wei, H., Deng, S., Hu, B., Chen, Z., Wang, B., Huang, J., et al. (2012). Granular bamboo-derived activated carbon for high CO$_2$ adsorption: the dominant role of narrow micropores. ChemSusChem 5, 2354—2360. doi: 10.1002/cssc.201200570

Xu, T., Huang, Y., Wu, B., Zhang, X., and Zhang, S. (2015). Biogas upgrading technologies: energetic analysis and environmental impact assessment. Chin. J. Chem. Eng. 23, 247—254. doi: 10.1016/j.cjche.2014.09.048

Yang, L., Ge, X., Wan, C., Yu, F., and Li, Y. (2014). Progress and perspectives in converting biogas to transportation fuels. Renew. Sustain. Energy Rev. 40, 1133—1152. doi: 10.1016/j.rser.2014.08.008

Zhao, B., Su, Y., Tao, W., Li, L., and Peng, Y. (2012). Post-combustion CO$_2$ capture by aqueous ammonia: a state-of-the-art review. Int. J. Greenhouse Gas Control 9, 355—371. doi: 10.1016/j.jgcgc.2012.05.006

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2020 Prauchner, Oliveira and Rodríguez-Reinoso. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.