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Synthetic silico-metallic mineral particles SSMMP: a new option for CO₂ capture and CO₂/N₂ separation from post-combustion technology

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Abstract

A new class of solid adsorbents based on pristine and ionic liquid (IL) functionalized synthetic silico-metallic mineral particles (SSMMP) for CO₂ capture and CO₂/N₂ separation are presented. Pristine particles were submitted to hydrothermal treatment producing synthetic talc. Samples were characterized by thermal analyses (TGA), specific surface area measurements (BET), infrared spectroscopy (FTIR), Raman spectroscopy, X-ray diffraction (XRD), NMR spectroscopy, and scanning electron microscopy (SEM). Values for specific surface areas varied from 5 m²/g for a sample with high IL content to 354 m²/g for pristine SSMMP. Samples presented distinct morphology as well as sorption capacities. IL and its increased concentration in the samples negatively influenced CO₂ sorption capacity but played an essential role in CO₂/N₂ selectivity. The best results were obtained from CO₂ sorption values for SSMMP-M1 without IL of 2.07 mmol CO₂/g at 1 bar and 4.93 mmol CO₂/g at 30 bar. In addition, a selectivity of 16.94 for the CO₂/N₂ mixture was achieved for SSMMP-5%-Im(nBu)-I sample. SSMMP particles are easy to obtain with low-cost starting materials used in the low energy and CO₂-free emission synthesis process. The sorption/desorption cycles proved the stability for both pristine and IL functionalized samples.

Keywords: synthetic silico-metallic mineral particles, synthetic talc, CO₂ sorption, CO₂ selectivity

1. Introduction

Climate change resulting from the growing CO₂ concentration is a significant concern. It is imperative to find an effective way to mitigate this gas, allowing the gradual transformation of a global economy based on fossil fuels to a new one based on low-carbon processes. The post-combustion CO₂ capture technology is considered a strong ally for this gradual transformation, given its high potential for application in retrofit processes (Fawzy et al., 2020; Younas et al., 2020). New propositions of materials for CO₂ capture combining low-cost and environmentally friendly products and processes are mandatory for the success of this technology. Aqueous amines, more specifically monoethanolamine- MEA, are the benchmark for CO₂ capture (Sexton and Rochelle, 2011; Gao et al., 2014; Xiao et al., 2018). Alkanolamines aqueous solution present some drawbacks such as high energy consumption, equipment corrosion, oxidative degradation, and high energy required to solution regeneration (Yu et al., 2012; Dinda, 2013; Espinal et al., 2013; Linneen et al., 2013). Solid sorbents appear as an alternative to aqueous amines due to lower energy requirement, ease maintenance, high tunability, and the lack of liquid waste generation (Nicot and Duncan, 2012; Linneen et al., 2013). Many kinds of solid sorbents are described as efficient for CO₂ capture. These include metal-organic frameworks (MOF) (Sabouni et al., 2014; Olajire, 2018; Younas et al., 2020), zeolitic imidasolate frameworks (ZIF) (Zhang et al., 2011; Kinik et al., 2016; Mohamedali et al., 2018), zeolites (Du et al., 2014; Wang et al., 2017; Nikolaidis et al., 2018; Kodasma et al., 2019), activated carbon (Chen et al., 2013; Houshmand et al., 2013), molecular basket sorbent (MBS) (Zhang et al., 2017), amine-modified solid sorbents (Chen et al., 2013; Houshmand et al., 2013; Linneen et al., 2013; Liu and Lin, 2013), aerogel based adsorbent (Kong et al., 2014), silica xerogels containing room temperature ionic liquids (RTIL) (dos Santos et al., 2020), mesoporous silica (He and Seaton, 2006; Kamarudin and Alias, 2013; Wang et al., 2015; Loganathan and Ghoshal, 2017; Duczinski et al., 2018; Polesso et al., 2019; Mohamedali et al., 2020), and carbon nanotubes (Razavi et al., 2011; Chen et al., 2013; Houshmand et al., 2013; Linneen et al., 2013; Liu and Lin, 2013).

Talc is a layered silicate mineral with the ideal formula of $\text{Mg}_3\text{Si}_4\text{O}_{10}(\text{OH})_2$ extensively used in industrial applications (Dumas et al., 2013b, 2013c). Structurally, talc is

composed of neutral layers stacks (forming a lamellar structure) linked together by weak Van Der Waals interactions (Claverie et al., 2018). In these layers, the atoms are linked together by covalent bonds. Its basal faces are not very reactive and constitute 90% of this material surface (Bremmell and Addai-Mensah, 2005; Claverie et al., 2018). The lateral surfaces are considered more reactive, containing SiOH and MgOH (Claverie et al., 2018). Despite its low price and high availability, natural talc presents some disadvantages depending on the intended application, such as a chemical inhomogeneity with the presence of natural cationic substitutions in the structure (Martin et al. 1999), mineralogical impurities with the presence of other crystalline phases (carbonates, sulfides, among others), and particle size distribution corresponding mainly to micrometer particles. To specifically answer to applications, it is possible to tailor synthetic talc in terms of composition, crystallinity, particle size, and layer thickness playing on synthesis parameters (Dumas et al. 2013b, c). SSMMP are formed in the first stage of synthetic talcs and can also be named amorphous or short-range nano talcs (Claverie et al., 2018). These materials undergo no hydrothermal process where the lamellar structure formation occurs by stacking octahedral and tetrahedral sheets (TOT). Therefore, the SSMMP particles present an amorphous structure, consisting of some Mg-octahedrons linked with Si-tetrahedrons distributed in the lower and upper part (Dumas, 2013; Dumas et al., 2015; Claverie et al., 2018). Unlike talc, these talc precursor materials have many reactive groups (Si-OH and Mg-OH) free on their surface (Chabrol et al., 2010; Dumas, 2013; Claverie et al., 2018). Among the applications already presented for these materials are the absorption of organic compounds (Ciesielczyk and Krysztafkiewicz, 2007; Ciesielczyk et al., 2007a, 2007b) and polymer fillers (Dietemann, 2012; Dias et al., 2015, 2016; Prado et al., 2020). The synthesis of talc precursors grafted with ionic liquids, using a fast, one-pot method, with low energy expenditure and using only an aqueous medium for dissolving the reagents, was patented (Dumas et al., 2013a). This invention presented new possibilities for applying these materials, such as catalysis and gas selective sorption processes.

Ionic liquids (IL) are appointed as energy-efficient and environmentally benign solvents for CO₂ capture (Nonthanasin et al., 2014) due to their excellent properties.

Low vapor pressure and flammability are achieved by different possibilities of combining anions and cations (Muldoon et al., 2007; Hasib-ur-rahman et al., 2010; Nonthanasin et al., 2014). However, high cost and viscosity are the main drawback of these materials (Hasib-ur-rahman et al., 2010).

To be a viable candidate for industrial application, a solid sorbent must be easy to produce on a large scale, at low cost, using environmentally friendly reactants, and easy to regenerate as well. Synthetic talc can also be a candidate for CO₂ capture. Moreover, the possibility of functionalizing it with ionic liquids (Dumas et al. 2013a; Dumas et al. 2014a, 2014b) appears as an excellent opportunity to create highly selective and low-cost materials.

Low cost and energy demand pristine and IL functionalized SSMMP were synthesized and characterized in this work. Synthetic talcs were also obtained through hydrothermal treatment of SSMMP. Samples were evaluated as solid sorbents in CO₂ sorption tests at 25°C of temperature and pressure up to 30 bar. Yet, sample recyclability and CO₂/N₂ selectivity were also evaluated.

2. Materials and Methods

The synthetic talc (ST) and synthetic silico-metallic mineral particles SSMMP were synthesized using synthetic methods based on the literature, as shown in Table 1. In addition, for comparison purposes, commercial zeolite MCM-41-C (Sigma Aldrich), synthetic zeolites MCM-41 and MCM-48, and natural talc from Luzenac were also tested for CO₂ sorption (Table 1, entries 11-13 and 2 respectively). The natural talc sample comes from a deposit near Luzenac, in the Occitanie region, France. All chemical reagents used in this work were purchased from Sigma Aldrich and used with no further purification.

Entry	Sample	Description	Reference
1	Talc Luzenac	natural talc from Luzenac, France	
2	ST-M1	synthetic talc	(Dumas et al., 2013b)– process P1

3	ST-M2	synthetic talc	(Dumas et al., 2013b)- process P3
4	SSMMP-M1	synthetic silico-metallic mineral particles	(Dumas et al., 2013b)- P1 talc precursor
5	SSMMP-M2	synthetic silico-metallic mineral particles	(Dumas et al., 2013b)- P3 talc precursor
6	SSMMP-20%-Im(Me)-NTf ₂	SSMMP with 20% of Si bearing Im(Me)-NTf ₂	(Dumas et al., 2013a)
7	SSMMP-10%-Im(Me)-NTf ₂	SSMMP with 10% of Si bearing Im(Me)-NTf ₂	(Dumas et al., 2013a)
8	SSMMP-5%-Im(Me)-NTf ₂	SSMMP with 5% of Si bearing Im(Me)-NTf ₂	Dumas et al. 2013a
9	SSMMP-5%-Im(nBu)-I	SSMMP with 5% of Si bearing Im(nBu)-I	(Dumas et al., 2013a)
10	SSMMP-5%-Im(nBu)-NTf ₂	SSMMP with 5% of Si bearing Im(nBu)-NTf ₂	(Dumas et al., 2013a)
11	MCM-41-C	commercial MCM-41	
12	MCM-41	synthetic MCM-41	(Meléndez-Ortiz et al., 2014); Comès et al. 2020
13	MCM-48	synthetic MCM-48	(Meléndez-Ortiz et al., 2014); Comès et al., 2020)

Table 1 - Samples used in this work

2.1. ST-M1 and SSMMP-M1 synthesis

Samples were synthesized using magnesium chloride hexahydrate, sodium metasilicate pentahydrate, and hydrochloric acid (1M solution). ST-M1 (Table 1 entry 2) was synthesized according to the P1 process described in Dumas et al. 2013b, and the hydrothermal reaction was performed at 300°C for 6 hours. SSMMP-M1 (Table 1 entry 4) is a talc precursor synthesized with P1 process described in Dumas et al. 2013b, undergoing no hydrothermal treatment.

2.2. ST-M2, SSMMP-M2 and SSMMP-IL synthesis

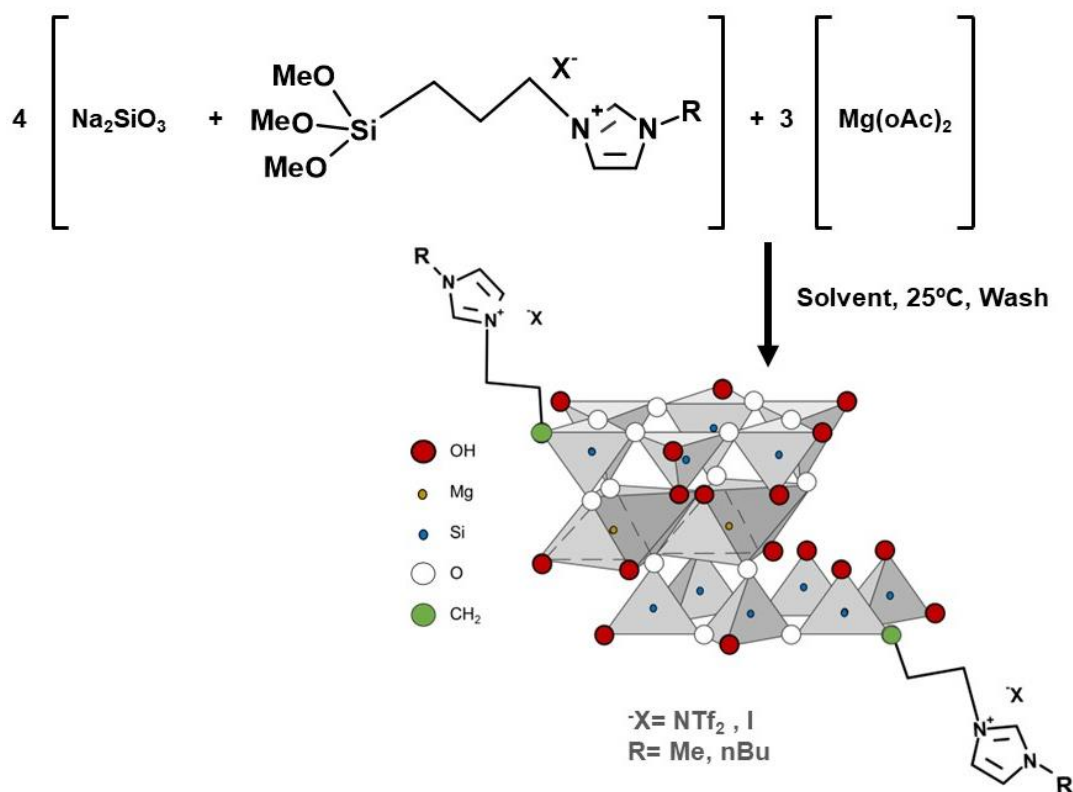


Figure 1 - Scheme of the SSMMP-IL synthesis

For the synthetic talc ST-M2 and SSMMP-M2 preparation (Table 1 entries 3 and 5), the following reagents were used: magnesium acetate tetrahydrate as Mg source, sodium metasilicate pentahydrate as Si source and acetic acid (1M solution). Dumas et al. 2013b described the P3 process used in their synthesis. The hydrothermal treatment of ST-M2 was carried out at 300°C for 6 hours. SSMMP-M2 is a talc precursor, i.e. it has undergone no hydrothermal treatment. For SSMMP-IL synthesis, corresponding to SSMMP functionalized with imidazolium molecules (Table 1 entries 6-10), the following reagents were additionally used: 1-(triethoxysilylpropyl)-3-methylimidazolium chloride [Im(Me)-Cl-silane], 1-(triethoxysilylpropyl)-3-n-butylimidazolium chloride [Im(nBu)-Cl-silane], sodium

iodide and bis(trifluoromethane)-sulfonimide lithium salt [LiNTf₂]. The synthesis method of SSMMP-IL samples was described in the patent application WO2013093339 (Dumas et al. 2013a), and the synthesis reaction shown in Figure 1. Samples functionalized with Im(nBu)-NTf₂ and Im(nBu)-I were obtained by replacing 5% of Si source by Im(nBu)-Cl-silane. After a classical exchange of Cl⁻ by NTf₂ anion (using LiNTf₂) and I⁻ (by using sodium iodide), SSMMP-5%-Im(nBu)-NTf₂ and SSMMP-5%-Im(nBu)-I were further obtained. Samples functionalized with Im(Me)-NTf₂ were synthesized by replacing 5%, 10% and 20% of Si source by Im(Me)-Cl-silane and making a classical exchange of Cl⁻ by NTf₂ anion (using LiNTf₂), resulting in SSMMP-5%-Im(Me)-NTf₂, SSMMP-10%-Im(Me)-NTf₂ and SSMMP-20%-Im(Me)-NTf₂, respectively.

2.3. MCM-41 and MCM-48 synthesis

The synthesis of MCM-41 and MCM-48 (Table 1, entries 12 and 13 respectively) were performed according to the methodology described by Meléndez-Ortiz et al. 2014, and carried out using tetraethoxysilane (TEOS), ammonium hydroxide (29% aqueous solution), and cetyltrimethylammonium bromide (CTAB). The calcination step was based on the method described by (Comès et al., 2020). MCM-41 synthesis was performed by adding 2.5 g of CTAB to 480 ml of deionized water. The solution was stirred and heated to 100°C. When the solution became translucent, keeping the temperature and stirring constant, 170 ml of ethanol was added. Subsequently, 50 ml of the ammonia solution was added to the mixture, and after another 5 minutes of stirring, 10 ml of TEOS was added as well and stirred for another 3 hours. The sample was washed with deionized water and dried at 100°C. CTAB was removed from the sample by incineration at 550°C for 8 hours. For the synthesis of MCM-48, 5.2 g of CTAB were added to 240 g of deionized water under constant agitation, then 100 ml of ethanol and 24 ml of the ammonia solution and finally, 6.8 g of TEOS. The solution remained under constant stirring for 16 hours at room temperature, then was washed and dried in the oven. CTAB was removed from the sample incineration at 550°C for 8 hours.

3. *Material characterization*

Thermogravimetric analysis (TGA) were obtained using a TA Instrument SDT-Q600. The temperature range was set at 25°C–1100°C with a heating rate of 10°C/min and under a nitrogen atmosphere. Fourier transform infrared spectroscopy (FT-IR) was performed using Perkin-Elmer FT-IR Spectrum 100 spectrometer in the range of 4000 cm⁻¹ to 600 cm⁻¹. RAMAN spectrograms were obtained by a alpha300 R access confocal Raman microscopy system from WiTec GmbH, equipped with a UHTS 300 spectrophotometer with a diffraction grating of 600g/mm BLZ=500 nm and using a He-Ne laser. The nitrogen adsorption-desorption isotherms were determined at 77 K, using a volumetric method with a Quantachrome Autosorb-1 apparatus. The isotherms were recorded in the 0.05-0.3 relative pressure range, using high purity nitrogen. Samples were outgassed for 15h at 120°C under vacuum before analysis. Specific surface areas were calculated using the Brunauer-Emmett-Teller (BET) method (Brunauer et al., 1938). X-ray diffraction (XRD) analyses were performed on disoriented powders, using a Bruker D8 Advance diffractometer operating under the reflection of the CuK α_{1+2} radiation ($\lambda = 1.5418 \text{ \AA}$), K α_2 being subtracted with Bruker Diffrac.Eva software for figures. XRD patterns were collected at room temperature over the 2-80°2 θ range, using 0.4 s counting time per 0.01°2 θ step. Solid-state ²⁹Si CP/MAS NMR spectrum was recorded on a 400 MHz Bruker Avance III spectrometer (LCC laboratory, Toulouse, France) in a static field of 9.4 T at resonance frequency of 79.49 MHz. Measurement was performed on powder in ZrO₂ rotors of 4.0 mm at a spinning frequency of 10 kHz and a 3ms CP pulse. Chemical shift is relative to tetramethylsilane (TMS). The powdered talcs skeletal density (ρ_s) was measured at 25°C using an Ultrapycnometer 1000 - Quantachrome Corporation pycnometer using ultra-high purity helium (Air Liquide / 99.999%). The particles morphology was assessed using a field emission scanning electron microscope (FESEM), Inspect F50 equipment (FEI Instruments).

3.1. *CO₂ adsorption measurements*

CO₂ sorption capacity was measured in an equilibrium cell (Figure 2) equipped with two chambers (one being a gas chamber and the other a sorption chamber), using

the pressure decay technique reported by KOROS and PAUL (Koros and Paul, 1976).

The materials' CO₂ sorption capacity was performed according to the methodology described in previous works (Bernard et al., 2017, 2018; Rojas et al., 2017; dos Santos et al., 2020). The tests were carried out in triplicates, at a constant temperature of 25°C, and CO₂ (Air Liquide, 99.998%) pressure varying from 0 to 30 bar. Before each test, samples were dried in an oven at 100°C for five hours. The sample mass used in the tests ranged from 0.6 g to 0.9 g. Before each new analysis, the sample was placed in an oven, where it remained for 1 hour at 110°C for CO₂ desorption. Samples with the best CO₂ sorption capacity and CO₂/N₂ selectivity were submitted to sorption/desorption tests 10 times. To corroborate sample stability after 10 cycles, BET and FTIR analysis were performed.

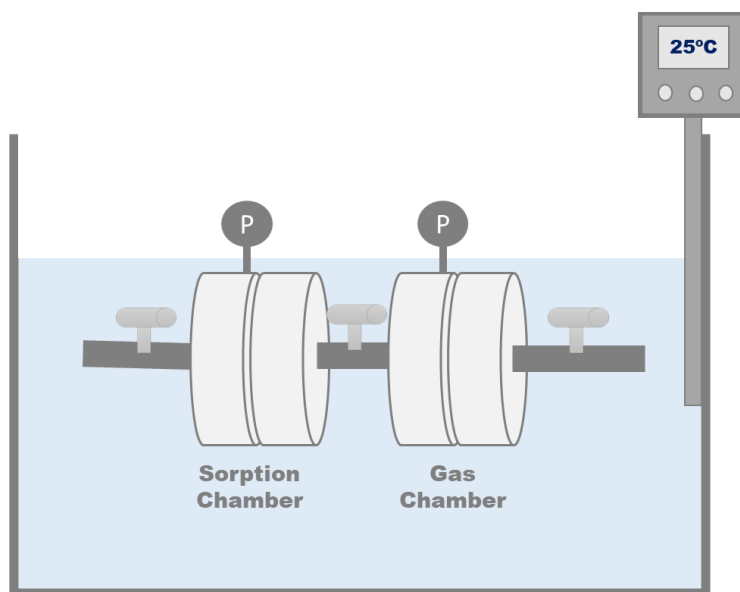


Figure 2 - System for measuring CO₂ sorption capacity.

3.2. CO₂/N₂ selectivity evaluation

Material selectivity was determined according to the procedure described in the literature by Azimi and Mirzaei 2016. The same CO₂ sorption system described above was used for the selectivity tests, replacing the pure CO₂ gas with a standard gas mixture with a composition of 15:85 (v/v) of CO₂/N₂. Selectivity tests were carried

out at an equilibrium pressure of 20 bar at 25°C. The gas composition of the mixture at the exit of the system was measured using a gas chromatograph (GC) (Shimadzu GC-14B) equipped with a thermal conductivity detector. The samples selectivity was determined by equation 1, where X_{CO_2} and X_{N_2} correspond to the molar fractions of CO_2 and N_2 sorbed by the sample, and Y_{CO_2} and Y_{N_2} are the molar fractions of CO_2 and N_2 present in the gas phase, respectively (Azimi and Mirzaei, 2016; Duczinski et al., 2018; Polesso et al., 2019).

$$S = \frac{X_{CO_2}/Y_{CO_2}}{X_{N_2}/Y_{N_2}} \quad (1)$$

4. Results

Figure 3 shows the infrared spectroscopy analysis of characteristic samples of synthetic talc, SSMMP, and SSMMP functionalized with IL (ST-M2, SSMMP-M2, and SSMMP-5%-Im(nBu)-I, respectively). The characteristic bands near 3700 cm^{-1} and 655 cm^{-1} , appearing in all samples, are attributed to $Mg_3\text{-OH}$ bond (Schroeder, 2002; Dias et al., 2015). The wide band between $3650\text{-}3000\text{ cm}^{-1}$ is related to the -OH of the water molecules corroborated by the band near 1635 cm^{-1} (Mor et al., 2017; Duczinski et al., 2018; Ouyang et al., 2018). The characteristic bands of Si-O and Si-O-Si bonds appear between $1050\text{-}900\text{ cm}^{-1}$ (Dias et al., 2015; Ouyang et al., 2018).

Figure 3 (c), samples containing IL, shows the appearance of three new characteristic bands. The band formed in the region of 1564 cm^{-1} is attributed to the $\text{C}=\text{C}$ bond of the imidazolium ring, the band around 1444 cm^{-1} related to the deformation of the CH_2 group of the IL (Duczinski et al., 2018) and the shoulder formed in the region 1142 cm^{-1} related to the Si-C bond present in the imidazolium lateral chain (Maria and Airoidi, 2001).

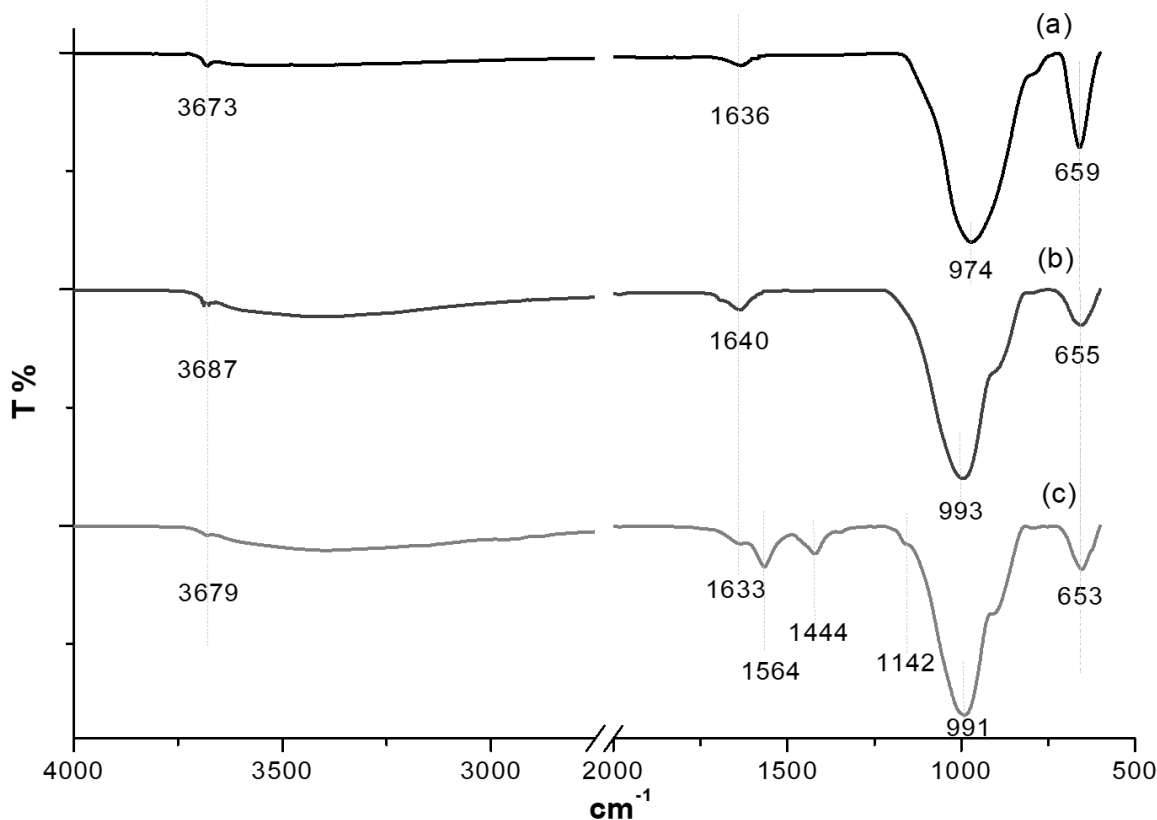


Figure 3 - FTIR spectra of (a) ST-M2, (b) SSMMP-M2 and (c) SSMMP-5%-Im(nBu)-I

Raman spectroscopy analysis was performed for SSMMP-M2, SSMMP-5%-Im(nBu)-I and SSMMP-5%-Im(nBu)-NTf₂ samples. A band in the region of 680 cm⁻¹, appearing in the Raman spectra of all samples, is attributed to the Si-O-Si symmetrical elongation mode, present in the SSMMP structure (see Figure S1) (Kloprogge, 2017). In the spectra of samples containing IL, the appearance of weak bands near 1400 cm⁻¹ is attributed to the in plane asymmetric stretch of the imidazole ring (H-C-H, C-C and C-N bonds) (Talaty et al., 2004; Chen et al., 2017; Paschoal et al., 2017). The bands in the region from 2839 to 3066 cm⁻¹ are attributed to the cation lateral chains and imidazolium ring C-H bonds stretching (Grondin et al., 2011; Chen et al., 2017).

Synthesized samples were characterized by thermogravimetric analysis (see Table 2). Results show that all samples, except the natural talc sample (Table 2, entry 1),

have a first mass loss starting between 20°C-46°C. This first mass loss is attributed to the loss of physisorbed water (Dumas et al., 2013c; Duczinski et al., 2018). For MCM-41-C, MCM-41, and MCM-48 (Table 2, entries 11-13 respectively), only this first mass loss at low temperatures was observed, indicating that all the surfactant used in the MCM-41 and MCM-48 synthesis was removed in the calcination step. The second mass loss refers to the loss of silanols (Si-OH) and magnesium hydroxide (Mg-OH) present on the synthetic talc sheet edges (Table 2, entries 2 and 3) (Dumas et al 2013c) and in the SSMMP surface (Table 2, entries 4-10). For samples grafted with IL, this second mass loss is also related to the loss of the imidazolium cation (Dumas et al., 2013c; Seyed and Nazemzadeh, 2017; Duczinski et al., 2018). Thus, particle size allied to the presence of the imidazolium cation are responsible for the significant variation on the second mass loss. Natural talc (Table 2, entry 1) showed higher thermal stability, evidencing no mass loss at low temperatures due to its hydrophobicity. A single-step mass loss was observed at high temperature (774°C to 1084°C), corresponding to the talc sheets dehydroxylation, occurring simultaneously with the enstatite crystallization and the silica formation (Chabrol et al., 2010). For ST-M1 and ST-M2 synthetic talcs, and their precursors SSMMP-M1 and SSMMP-M2 (Table 2, entries 2, 3, 4 and 5, respectively), dehydroxylation, enstatite crystallization and silica formation steps occurred in different ways. For ST-M1 and SSMMP-M1 samples, this step occurs continuously (corresponding to the second mass loss), and for ST-M2 and SSMMP-M2 in two distinct steps corresponding to the second and third mass loss, here attributed to the dehydroxylation of the sheet edges and of the bulk, respectively (Chabrol et al., 2010; Dumas et al., 2013c). Typical thermograms are presented in the supplementary file (Figures S2-S4).

Entry	Sample	1 st mass loss		2 nd mass loss		3 rd mass loss	
		<i>T</i> (°C)	w/w %	<i>T</i> (°C)	w/w %	<i>T</i> (°C)	w/w %
1	Talc Luzenac	-	-	-	-	774-1084	4.6
2	ST-M1	20-150	10.8	150-830	5.5	-	-
3	ST-M2	26-152	0.3	152-598	4.4	598-865	3.5

4	SSMMP-M1	24.3-281.8	20.3	281.8-791.7	6.9	-	-
5	SSMMP-M2	25.8-159.9	10.5	159.9-417.2	6.9	417.2-850.9	4.5
6	SSMMP-20%-Im(Me)-NTf ₂	29-234	11.8	234-708	28.4	-	-
7	SSMMP-10%-Im(Me)-NTf ₂	29-232	19.9	232-782	20.9	-	-
8	SSMMP-5%-Im(Me)-NTf ₂	38-170	18.8	170-584	18.4	-	-
9	SSMMP-5%-Im(nBu)-I	26-250	18.8	250-643	12.6	-	-
10	SSMMP-5%-Im(nBu)-NTf ₂	22-286	18.7	286-712	11.7	-	-
11	MCM-41-C	44.6-795	2.6	-	-	-	-
12	MCM-41	46-791.3	19.6	-	-	-	-
13	MCM-48	22.8-791.7	26.4	-	-	-	-

Table 2 - Thermogravimetric analyses results

BET measurement was also used to characterize the samples. As seen in Table 3, the synthetic talcs ST-M1 and ST-M2 (329 m²/g and 125 m²/g, respectively) present higher specific surface area values when compared to natural talc from Luzenac (20 m²/g). For SSMMP-IL, functionalized with imidazolium molecules, increasing imidazolium content decreases the specific surface area values (compare SSMMP-5%-Im(Me)-NTf₂, 79 m²/g; SSMMP-10%-Im(Me)-NTf₂, 55 m²/g; and SSMMP-20%-Im(Me)-NTf₂, 5 m²/g). Silica gel immobilized with tetraethylenepentamine (Nonthanasin et al., 2014) shows the same behavior as SSMMP-IL samples. In addition, the decrease in the samples' specific surface area with increasing ionic liquid concentration indicate that the functionalization of SSMMP-M1 with imidazolium molecules was successful (Younas et al., 2020). The synthetic method directly influences the specific surface area values (compare samples ST-M1, 329 m²/g and ST-M2, 125 m²/g). Table 3 also shows the density values of the samples under study in the range of 1.89-2.70 g/cm³, attributed to the similarity of the structure of synthetic samples with the one of natural talc, and the presence of functional groups attached to their structure for the functionalized samples.

Entry	Sample	Specific surface area (m ² /g)	Density (g/cm ³)
1	Talc Luzenac	20	2.70
2	ST-M1	329	2.53
3	ST-M2	125	2.71
4	SSMMP-M1	354	2.55
5	SSMMP-M2	325	2.31
6	SSMMP-20%-Im(Me)-NTf ₂	5	2.21
7	SSMMP-10%-Im(Me)-NTf ₂	55	1.90
8	SSMMP-5%-Im(Me)-NTf ₂	79	1.96
9	SSMMP-5%-Im(nBu)-I	156	2.05
10	SSMMP-5%-Im(nBu)-NTf ₂	151	2.23
11	MCM-41-C	841	2.33
12	MCM-41	1188	2.42
13	MCM-48	1361	1.89

Table 3 - Sample physical parameters

XRD analysis (Figure S5) shows for ST-M1 and ST-M2 the presence of intense diffraction peaks, very similar to those found for Luzenac talc corroborating the crystallinity of these materials (Dias et al., 2015; Prado et al., 2020). SSMMP-5%-Im(Bu)-I present an X-ray diffraction pattern with broad and low-intensity diffraction peaks, confirming the lower stacking order and growth in the ab plane, and the amorphous structure (Claverie et al., 2018).

NMR analysis confirms that organic groups are well present in the precursor structure. ²⁹Si NMR spectra (Figure S6) shows two signals regions. The first one between -75 and -95 ppm corresponds to Q-type Si atoms (with Si-O-Si, Si-O-Mg and Si-OH bonds), probably Q¹ and Q² for the most part because of the small TOT entities of the precursor (based on literature, please see discussion in the next

paragraph). The second region between -40 and -65 ppm corresponds to T-type Si atoms with Si-C bond, more probably T¹ around -49 ppm and T² around -53 ppm (basing upon reported studies, for example Fuji and Hayashi 2005, or Gallégo et al. 2007). These results indicate that the organic group is covalently bonded with the phyllosilicate-like structure of SSMMP (see structure in Figure 1).

Table 4 and Figure 4 present the CO₂ sorption capacity test results for each of the synthesized samples, at equilibrium pressures of 1-30 bar of CO₂ at a temperature of 25°C. When comparing the results of CO₂ sorption capacity of ST-M1 and ST-M2 (Table 4, entries 2 and 3, respectively) with their respective precursors (precipitated material without hydrothermal process) SSMMP-M1 and SSMMP-M2 (Table 4, entries 4 and 5), we observed that for both precursor materials, CO₂ sorption capacity was higher than for ST at all CO₂ equilibrium pressures. At equilibrium pressure of 1 bar, SSMMP-M1 evidenced a CO₂ sorption capacity of 2.07 mmol CO₂/g, an increase of 1.99 mmol CO₂/g, compared to ST-M1 (0.08 mmol CO₂/g, 1 bar CO₂). This higher sorption capacity of SSMMP is directly related to the distinct structures presented by the precipitated materials (SSMMP-M1 and SSMMP-M2) compared to synthetic talcs obtained after hydrothermal treatment process. Literature (Chabrol et al., 2010; Dumas et al., 2015; Claverie et al., 2018) points out that unlike ST, presenting a lamellar shape with several stacked layers, the precipitated SSMMP, before going through hydrothermal treatment process, consists of structures composed of small TOT entities, formed by Mg-octahedrons [MgO₄(OH)₂] and Si-tetrahedrons (SiO₄) distributed in the lower and upper part of its Mg-octahedral "sheet" resulting in a structure with a number of Si-OH and Mg-OH reactive groups on their surfaces. When the precipitated material (SSMMP) is subjected to hydrothermal treatment, both the Mg-octahedral sheet and the two Si-tetrahedron sheets grow in the (ab) crystallographic plane and along the c axis, with a large amount of Si-tetrahedrons connected to three other Si-tetrahedrons and only few ones connected to only two or one Si-tetrahedrons (located on the edge of the sheets). The same happens for the Mg-octahedral sheet (Chabrol et al., 2010; Dumas et al., 2013c, 2015; Claverie et al., 2018). According to literature (Bremmell and Addai-Mensah, 2005), only 10% of talc surfaces have Si-OH and Mg-OH

reactive groups, present only in small amounts on the talc side surface (Claverie et al., 2018). These groups (-SiOH and -MgOH) present high affinity to CO₂, so the greater amount of these groups on the SSMMP surface is responsible for the higher CO₂ sorption capacity of these samples when compared to their respective synthetic talc (Rimola et al., 2013; Polesso et al., 2019). For SSMMP-IL, as seen in Table 4, the increase in the IL content caused a decrease in CO₂ sorption capacity, lower for all SSMMP-IL than for SSMMP-M1 and SSMMP-M2 regardless of the analyzed equilibrium pressures. SSMMP-M2 exhibited CO₂ sorption capacity of 95,8% (1 bar), 123,7% (10 bar) and 212,9% (30 bar) higher than SSMMP-5%-Im(nBu)-NTf₂. The CO₂ adsorption process on the IL and the SSMMP surface is physical; there is no chemical reaction between the material and the CO₂ (Ramdin et al., 2012; Houshmand et al., 2013). The incorporation of imidazolium molecules in the sample results in a decrease in the specific surface area of SSMMP-M2, decreasing its CO₂ sorption capacity due to limited interactions of SSMMP/CO₂ (Ramdin et al., 2012). These results evidenced the relevance of the SSMMP/CO₂ interaction compared to the IL/CO₂. As previously discussed, SSMMP-M1 and M2 have an amorphous structure consisting of small TOT entities (Dumas et al., 2015), containing several reactive groups (-SiOH and -MgOH) on their surface which can interact with CO₂. Increasing in the IL load results in a decrease in both the SSMMP specific surface area and surface reactive groups, consequently reducing SSMMP/CO₂ interaction (Ramdin et al., 2012; Rimola et al., 2013; Kinik et al., 2016; Polesso et al., 2019). Malherbe et al. (2010) analyzed the CO₂ molecule interaction with different silica-based materials surface. They stated that the interaction between CO₂/silica surface occurs by physical adsorption through dispersive and electrostatic interactions and by CO₂ weak interaction with the OH group present in the silica surface ($\text{H}^{\delta+} \dots \delta^- \text{O}=\text{C}=\text{O}^{\delta-}$). Our group demonstrated that for IL grafted in silica-based solids, the anion size and charge density play an important role in CO₂ sorption capacity being the solid binding sites the primary driver for CO₂ capture (Aquino et al., 2015). It was also evidenced that CO₂ establish weaker interactions with the cation showing no competition with the anion for the same locations. The preferred location for the CO₂ molecule in the IL is near methyl groups explained by the existence of a

local conformational equilibrium and, consequently, a higher cavity formation probability (Corvo et al., 2013). These finds, combined with the results presented by Simon et al. (2018), evidencing that the increase in the cation carbon chain increases the CO₂ sorption capacity in the IL, corroborate the results of this work evidencing the higher sorption capacity achieved by the SSMMP grafted with the IL containing the Im(nBu)⁺ cation (entries 9 and 10, Table 4) compared to Im(Me)⁺ (entries 6 to 8, Table 4).

These results indicate that an ideal balance between the IL and OH content allied to specific surface area value would be necessary to achieve an adequate CO₂ absorption capacity.

Comparing SSMMP-M1 and SSMMP-M2, one can see that the sample specific surface area only influenced CO₂ sorption capacity at low pressures (up to 5 bar). SSMMP-M1 exhibited a surface area 8.9% superior to SSMMP-M2 and sorption capacity 9.5% higher than SSMMP-M2 at 1 bar of CO₂ equilibrium pressure. For all pressures above 10 bar, the SSMMP-M1 presented the best sorption capacity performance.

Entry	Sample	mmol CO ₂ /g sorbent at 1 bar	mmol CO ₂ /g sorbent at 10 bar	mmol CO ₂ /g sorbent at 30 bar
1	Talc Luzenac	0.50	0.68	0.79
2	ST-M1	0.08	0.15	0.35
3	ST-M2	0.19	0.38	0.74
4	SSMMP-M1	2.07	3.19	4.93
5	SSMMP-M2	1.86	3.78	6.32
6	SSMMP-20%-Im(Me)-NTf ₂	0.37	0.63	0.56
7	SSMMP-10%-Im(Me)-NTf ₂	0.50	1.05	1.00
8	SSMMP-5%-Im(Me)-NTf ₂	0.58	1.47	1.68
9	SSMMP-5%-Im(nBu)-I	0.89	1.54	2.16
10	SSMMP-5%-Im(nBu)-NTf ₂	0.95	1.69	2.02

11	MCM-41-C	0.40	2.04	5.25
12	MCM-41	1.58	4.12	6.21
13	MCM-48	1.71	4.28	7.75

Table 4 - CO₂ sorption values for all samples at 1, 10 and 30 bar

Mesoporous silica MCM-41-C, MCM-41, and MCM-48 were also evaluated for CO₂ sorption as well-known solid sorbents and support materials for capturing CO₂ (He and Seaton, 2006; Kamarudin and Alias, 2013; Wang et al., 2015; Loganathan and Ghoshal, 2017; Mohamedali et al., 2020). Comparing these three materials with SSMMP-M1 and SSMMP-M2, it is observed (Figure 4) that, up to 2.5 bar, these last showed higher sorption capacity than the mesoporous silica and up to 20 bar compared to MCM-41-C. At higher pressures from 4 bar, MCM-41 and MCM-48 have greater sorption capacity compared to SSMMP-M1 and SSMMP-M2, except for MCM-41 similar but just below SSMMP-M2 above 20 bar. However, as the primary purpose of these materials is to capture CO₂ in post-combustion processes, where the CO₂ partial pressure is 1 bar, SSMMP-M1 and, to a lesser extent, SSMMP-M2 are still considered the best materials. Furthermore, they are synthesized in just one step (one-pot), quantitative precipitation showing excellent atom economy process, with low-cost reagents and no energy expenditure (no hydrothermal treatment or calcination step).

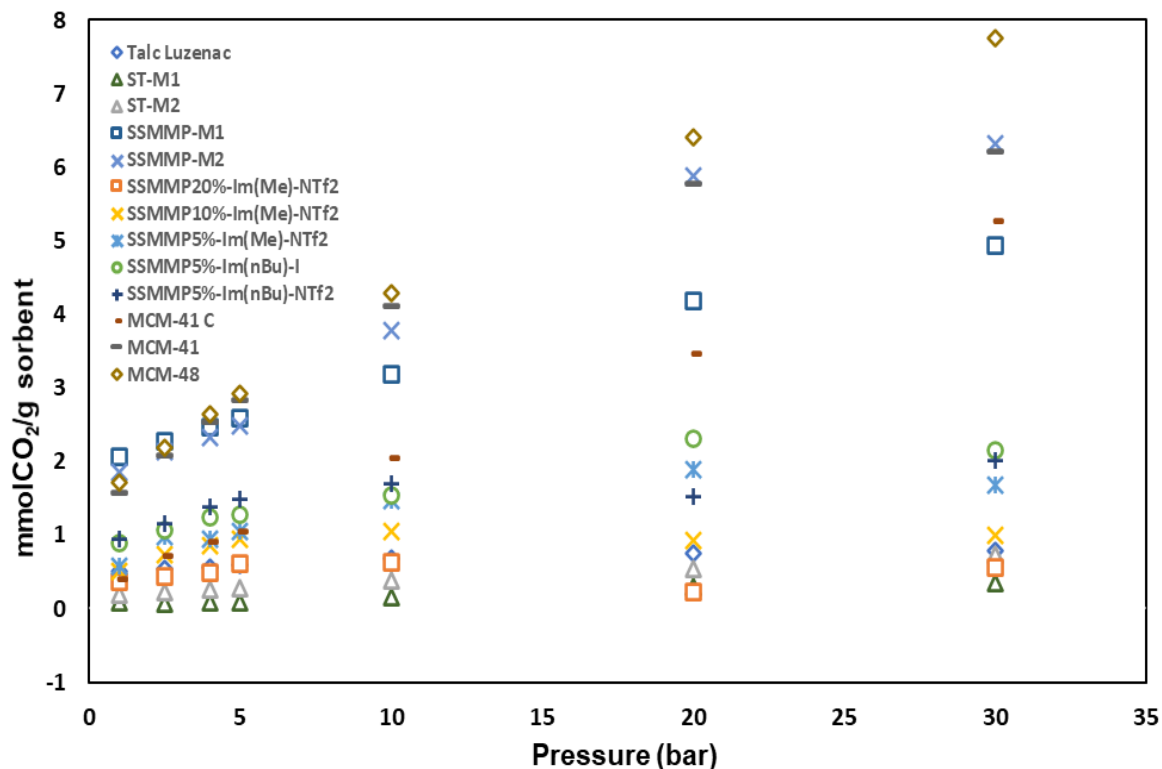


Figure 4 - Behavior of the samples as sorbents for CO₂ capture at 25°C and a pressure range of 1-30 bar.

Table 5 presents sorption capacity (at 1 bar) of other inorganic silicate materials described in literature. It is clear from this table that SSMMP-M1 presents greater CO₂ sorption capacity at low pressure. Moreover, SSMMP-M1 is cheaper when compared with other solid sorbents considered suitable for capturing CO₂ at low pressure, such as MOFs (Mg-MOF-78 in (Yu and Balbuena, 2013) and ZIFs (IL@ZIF-8 in (Mohamedali et al., 2018)). Furthermore, MOFs are considered expensive materials to produce, unstable in basic and acidic solutions and humid environments, turning their commercial application highly limited (Yu and Balbuena, 2013; Younas et al., 2020). On the other hand, ZIFs, (a subfamily of MOFs) present a greater or comparable sorption capacity with SSMMP-M1 only when impregnated with ILs (Mohamedali et al., 2018).

Sample	Sample type	mmol CO ₂ /g sorbent	Analysis conditions	Reference
SX	Silica xerogel	1.13	1 bar; 25°C	(dos Santos et al., 2020)
MCM-41-30	Mesoporous silica	<0.5	1 bar; 30°C	(Loganathan and Ghoshal, 2017)
SBA-15	Mesoporous silica	0.45	1 bar; 45°C	(Sanz et al., 2010)
MCSU-H	Mesoporous silica	<0.2	1 bar; 40°C	(Fujiki et al., 2015)
NaMnSi ₁₀ O _x	Metal silicate	<1.8	1 bar; 25°C	(Li et al., 2020)
NaNiSi ₁₀ O _x	Metal silicate	<0.9	1 bar; 25°C	(Li et al., 2020)
NaCuSi ₁₀ O _x	Metal silicate	<1.6	1 bar; 25°C	(Li et al., 2020)
MMT	Porous Montmorillonite	<0.5	1 bar; 25°C	(Atilhan et al., 2016)
E-VER	Exfoliated Vermiculite	<1.0	1 bar; 25°C	(Zhang et al., 2020)
SSMMP-M1	Talc precursor	2.07	1 bar; 25°C	This work

Table 5 - CO₂ sorption values for different inorganic silicate materials found in the literature

Figure 5 highlights that morphology appears to play a role in CO₂ capture. It is possible to observe that samples ST-M1 and ST-M2 (entries (a) and (b)) have a compact and ordered surface compared to all other studied samples (SSMMP-M1, SSMMP-M2, all SSMMP-IL samples, MCM-41 and MCM-48). The two latter have relatively smaller particles and rougher surface. ST-M1, Talc Luzenac, ST-M2, and SSMMP-20%-Im(Me)-NTf₂ have low CO₂ sorption capacity, less than 1 mmol CO₂/g of sorbent even at 30 bar of CO₂ pressure. All other samples have sorption capacity equal to or greater than 1 mmol CO₂/g- of sorbent at 30 bar.

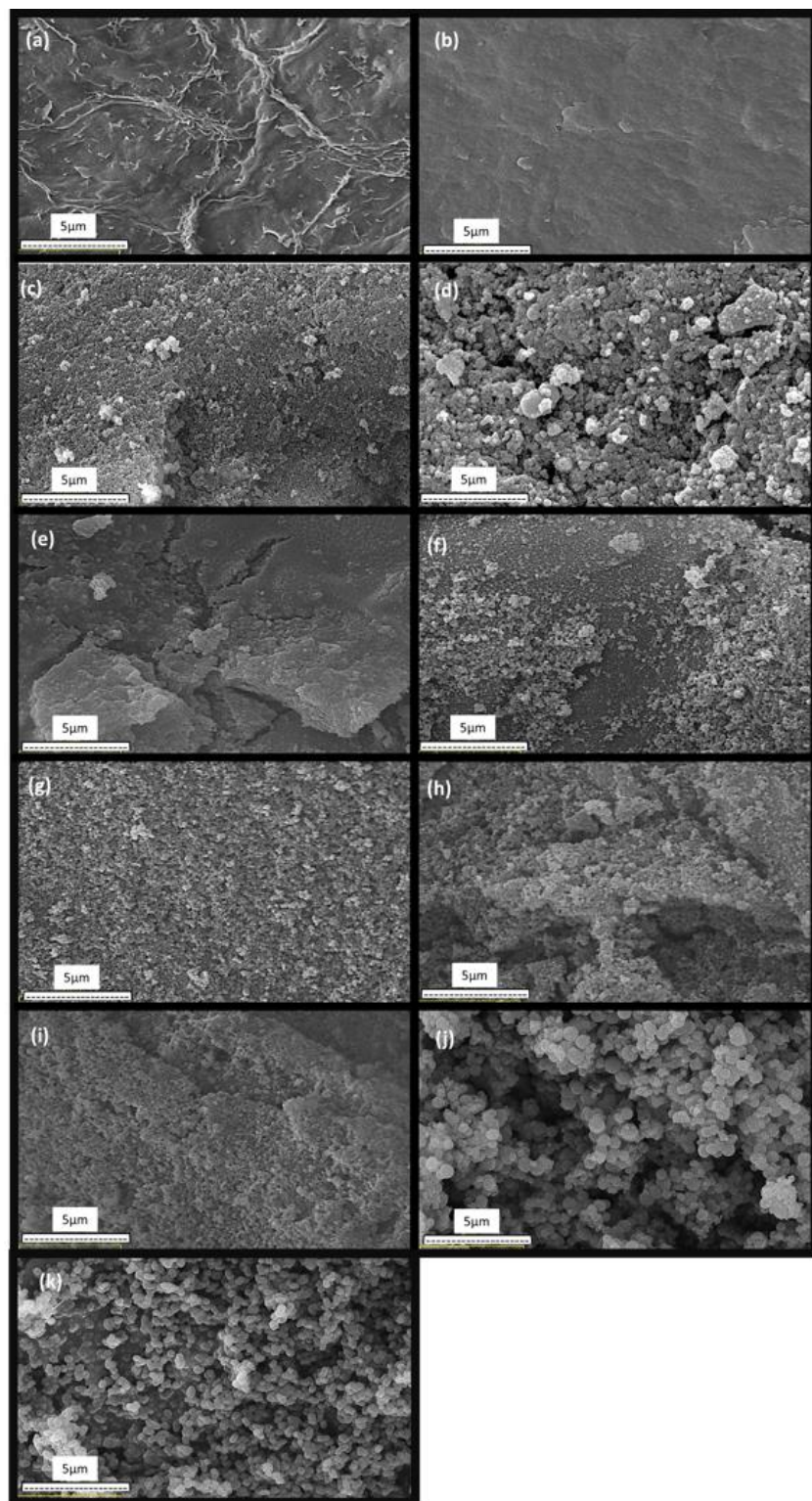


Figure 5 - SEM images of (a) ST-M1, (b) ST-M2, (c) SSMMP-M1, (d) SSMMP-M2 (e) SSMMP-20%-Im(Me)-NTf₂, (f) SSMMP-10%-Im(Me)-NTf₂, (g) SSMMP-5%-Im(Me)-NTf₂, (h) SSMMP-5%-Im(nBu)-I, (i) SSMMP-5%-Im(nBu)-NTf₂, (j) MCM-41 and (k) MCM-48

Figure 5 highlights that morphology appears to play a role in CO₂ capture. It is possible to observe that samples ST-M1 and ST-M2 (entries (a) and (b)) have a compact and ordered surface compared to all other studied samples (SSMMP-M1, SSMMP-M2, all SSMMP-IL samples, MCM-41 and MCM-48). The two latter have relatively smaller particles and rougher surface. ST-M1, Talc Luzenac, ST-M2, and SSMMP-20%-Im(Me)-NTf₂ have low CO₂ sorption capacity, less than 1 mmol CO₂/g of sorbent even at 30 bar of CO₂ pressure. All other samples have sorption capacity equal to or greater than 1 mmol CO₂/g- of sorbent at 30 bar.

4.1. SSMMP-M1 recycle test

Figure 6 presents the CO₂ sorption/desorption test results for SSMMP-M1 and SSMMP-5%-Im(Bu)-I. As seen for SSMMP-M1, sorption capacity ranged from 1.94 to 2.08 mmol of CO₂/g during the 10 cycles at 25°C, presenting an average sorption capacity of 2.00 mmol of CO₂/g. For SSMMP-5%-Im(nBu)-I the sorption capacity variation during the 10 sorption/desorption cycles was 0.83 to 0.97 mmol CO₂/g and its average sorption capacity 0.89 mmol CO₂/g.

To confirm SSMMP-M1 and SSMMP-5%-Im(nBu)-I stability, BET and FTIR analyzes were performed after 10 sorption/desorption cycles. According to the results (Figures S7 and S8), no structural and surface area changes were observed after recycling.

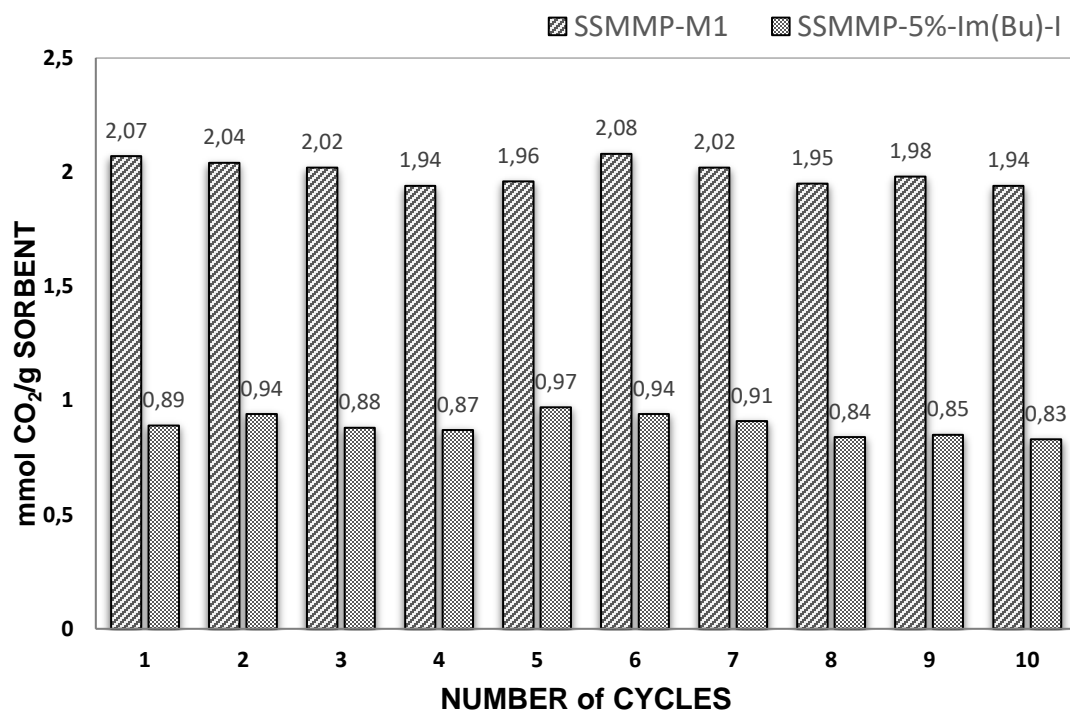


Figure 6 - CO₂ sorption/desorption test for SSMMP-M1 and SSMMP-5%-Im(nBu)-I at 1 bar and 25°C.

4.2. CO₂/N₂ selectivity

CO₂/N₂ selectivity was evaluated for SSMMP-M1 and SSMMP-5%-Im(nBu)-I. Results indicated that the presence of ILs in the sample increased 124.67% the CO₂ removal (selectivity of 16.94 ± 0.87) compared to pristine SSMMP-M1 (selectivity of 7.54 ± 0.64). The greater CO₂ affinity presented by the sample functionalized with IL is in agreement with other published studies (Harvey et al., 2016; Duczinski et al., 2018; Mohamedali et al., 2018; Polesso et al., 2019). These results indicate that the presence of the IL on the SSMMP surface increased the sample affinity to CO₂ (Nkinahamira et al., 2017; Zhu et al., 2018; Polesso et al., 2019). The obtained selectivity results proved to be promising when compared to other materials reported in the literature. Mohamedali et al. 2018 analyzed the selectivity of pure ZIF-8 and incorporated with different concentrations of acetate-based ILs samples in CO₂/N₂ mixtures at a pressure of 1 bar and a temperature of 30°C. For pure ZIF-8, CO₂/N₂

selectivity was 5. For all samples with different concentrations of incorporated IL, the selectivity was 10, corroborating the important effect of ILs in CO₂/N₂ selectivity.

5. Conclusion

This work reported the synthesis, characterization, CO₂ sorption capacity, and CO₂/N₂ selectivity of different synthetic materials. For SSMMP-IL, the decrease in the specific surface area resulting from the increase in the IL content decreased CO₂ sorption capacity due to reducing reactive groups present on the SSMMP surface. Pristine SSMMP-M1 presented an excellent sorption capacity at low pressures (2.07 mmol CO₂/g sorbent at 1 bar) compared to synthetic talc and other solid sorbents presented in the literature. Unlike CO₂ sorption tests, CO₂/N₂ selectivity was improved in the presence of the IL showing an increase of 124% in CO₂ selectivity (CO₂/N₂ mixture) compared to pristine SSMMP-M1. SSMMP-M1 and SSMMP-5%-Im(nBu)-I stability was also verified after 10 consecutive CO₂ sorption/desorption cycles. SSMMP-M1 CO₂ sorption capacity and the selectivity of SSMMP-5%-Im(nBu)-I are promising for industrial use in post-combustion capture and separation processes due to the ease of synthesis (one-pot), low cost of reagents, and low energy consumption. There are no heating steps for synthesis or the need for calcination as well, unlike most solid sorbents presented in the literature as suitable materials for capturing CO₂ at low pressures. Concerning material disposal, after many cycles of SSMMP used for CO₂ capture, it can still be used in other applications (hydrothermal process, polymer filler, among others).

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