Production of spherical mesoporous molecularly imprinted polymer particles containing tunable amine decorated nanocavities with CO2 molecule recognition properties


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Production of spherical mesoporous molecularly imprinted polymer particles containing tunable amine decorated nanocavities with CO₂ molecule recognition properties

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HIGHLIGHTS

- Spherical poly(AAm-co-EGDMA) particles with CO₂ recognition cavities were synthesized.
- Optimum ratio of acetonitrile to toluene in the organic phase was 30:70.
- Particles were thermally stable up to 240 °C as revealed by thermogravimetric analysis.
- CO₂ capture capacity reached 0.62 mmol/g at 25 °C and 0.15 bar CO₂ partial pressure.
- CO₂ capture capacity increased by increasing the density of CO₂-philic NH₂ groups.

GRAPHICAL ABSTRACT

CO₂ breakthrough curves

A B S T R A C T

Novel spherical molecularly imprinted polymer (MIP) particles containing amide-decorated nanocavities with CO₂ recognition properties in the poly(acrylamide-co-(ethylene glycol dimethacrylate)] mesoporous matrix were synthesized by suspension polymerization using oxalic acid and acetonitrile/toluene as dummy template and porogenic solvents, respectively. The particles had a maximum BET surface area, S_{BET}, of 457 m²/g and a total mesopore volume of 0.92 cm³/g created by phase separation between the copolymer and porogenic solvents. The total volume of the micropores (d < 2 nm) was 0.1 cm³/g with two sharp peaks at 0.84 and 0.85 nm that have not been detected in non-imprinted polymer material. The degradation temperature at 5% weight loss was 240–255 °C and the maximum equilibrium CO₂ adsorption capacity was 0.56 and 0.62 mmol/g at 40 and 25 °C, respectively, and 0.15 bar CO₂ partial pressure. The CO₂ adsorption capacity was mainly affected by the density of CO₂-philic NH₂ groups in the polymer network and the number of nanocavities. Increasing the content of low-polar solvent (toluene) in the organic phase prior to polymerization led to higher CO₂ capture capacity due to stronger hydrogen bonds between the template and the monomer during complex formation. Under the same conditions, molecularly imprinted particles showed much higher CO₂ capture capacity compared to their non-imprinted counterparts. The volume median diameter (73–211 μm) and density (1.3 g/cm³) of the produced particles were within the range suitable for CO₂ capture in fixed and fluidized bed systems.

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1. Introduction

Excessive concentration of atmospheric CO$_2$, mainly because of extensive utilization of fossil fuels, has significantly contributed to global warming and requires immediate remedies [1,2]. Carbon capture and storage is the most viable short- to medium-term approach for decreasing the amount of CO$_2$ released into the atmosphere [3,4]. Carbon capture technologies can be classified into three main groups: oxy-fuel combustion, pre-combustion, and post-combustion carbon capture [5–9], the latter being the most practical approach to retrofit carbon capture onto existing power generation systems [4,9]. Post-combustion carbon capture by amine scrubbing is the most established technique for CO$_2$ capture from flue gases [10]. However, amines are corrosive and undergoes degradation on exposure to CO$_2$, O$_2$, SO$_2$, and heat, leading to increased costs and generation of products that are harmful to human health and the environment [11–15]. In addition, amine-based scrubbing systems require large volume, because of their low interfacial area per unit volume and high energy consumption for the solvent regeneration, decreasing the power output typically by 20–25% [16,17].

Solid CO$_2$ adsorbents are more efficient and environmentally friendly than amine solutions, because of their lower regeneration energy, high surface area per unit volume, and low volatility and corrosivity [16,18]. Zeolites are the most explored sorbents for post-combustion CO$_2$ capture with a very high CO$_2$ uptake and selectivity in some cases [19–21]. However, the presence of moisture in flue gases dramatically reduces their adsorption capacity and the desorption step must be carried out at higher temperatures [22,23].

Silica-based porous materials are associated with low CO$_2$ uptake, because of the low affinity of silica towards CO$_2$, but can readily be functionalized with amine groups [20,24–26], which exhibit a high affinity towards the molecules with a high quadrupole moment and polarizability. Amongst all the molecules present in post-combustion flue gases, CO$_2$ has the largest quadrupole moment and polarizability, which results in much higher CO$_2$ adsorption capacity of amine-functionalized adsorbents compared to their unmodified counterparts [27]. Amine groups can be incorporated into the silica surface through physical impregnation or covalent tethering [28–31]. Although impregnated silica materials exhibit lower cyclic stability than adsorbents with covalently tethered amines, both methods lead to a significant increase in CO$_2$ capture capacity, especially in the presence of moisture [15,32,33].

Metal organic frameworks (MOFs) have gained a lot of interest because of their high specific surface area, ease of structural tuning, narrow pore size distribution, and high CO$_2$ adsorption at elevated pressures [34]. However, they mostly suffer from low CO$_2$ adsorption at low pressures [35,36], low CO$_2$ selectivity, and significant deterioration in the presence of moisture, NO$_x$ and SO$_x$ [20,37]. Carbonaceous materials are another attractive type of adsorbents, because of their high surface area, tunable pore structure, low heat of adsorption, and low sensitivity to moisture and gas impurities [38]. Since CO$_2$ is adsorbed by physisorption, they suffer from a low CO$_2$ adsorption capacity under low pressure [2], which can be improved by potassium intercalation [39–41] and introduction of nitrogen functional groups [42,43]. The main disadvantage of highly porous materials, such as activated carbon and some MOFs is their low density, which can limit their application in fluidized bed reactors [7,39].

Polymer-based materials such as porous aromatic frameworks (PAFs), hyper cross-linked polymers (HCPs), and covalent organic polymers (COPs) [44–46] are a new class of CO$_2$ sorbents characterized by high selectivity and CO$_2$ uptake, hydrothermal stability and ease of structural modification. Polymeric adsorbents are usually produced as very fine particles that pack too tightly and can easily be entrained out of the bed. Therefore, they must be pel- letized to be used in fluidized bed systems, which can block their reactive sites and reduce CO$_2$ adsorption capacity. Recently, molecularly imprinted polymer (MIP) particles for CO$_2$ capture with a separation factor of up to 340, insensitive to moisture, SO$_2$, NO, and O$_2$, have been synthesized by bulk polymerization [47,48]. Molecular imprinting creates template-shaped cavities within the polymer matrix with molecular recognition properties towards a specific target molecule [49]. However, bulk polymerization is not suitable for large-scale production. After polymerization, the bulk polymer is crushed, ground and sieved to obtain particles of the desired size range, which is time consuming, laborious, and only 30–40% of the ground polymer can be recovered. In addition, the particles produced by this method are susceptible to mechanical attrition because of their irregular shape and broad size distribution [49].

In suspension polymerization each monomer droplet acts as a tiny batch reactor, which facilitates heat transfer and thus, the polymerization occurs faster and the final conversion of the monomer can reach higher values [50]. Due to their spherical shape, the particles have a lower tendency to break down and their size can be finely tuned by controlling the operating conditions and phase compositions during emulsification [50]. In addition, the process is scalable and higher particle yields can be achieved due to the smaller amount of waste (off-specification) material.

In this work, novel spherical acrylamide-based MIP particles for CO$_2$ capture were fabricated by suspension polymerization in an oil-in-water (O/W) emulsion. Amide-decorated nanocavities were created by crosslinking acrylamide in the presence of oxalic acid, followed by extraction of the dummy template to expose nanocavities. The particles were inherently amide-functionalized and there was no need for amine grafting to increase CO$_2$ adsorption capacity. Due to CO$_2$-philic moieties on the walls of the cavities and their shape that is complementary to CO$_2$ molecules, the particles are highly selective to CO$_2$. They are suitable for use in fluidized bed reactors, because of their tunable diameter greater than 100 µm and relatively high density. The emulsion formulation and operating conditions were optimized to maximize CO$_2$ adsorption capacity of the particles in realistic post-combustion CO$_2$ capture situations.

2. Experimental section

2.1. Materials

Oxalic acid (OA), acrylamide (AAm), acetonitrile (AN), toluene (TL), methanol, and 0.1 M hydrochloric acid were supplied by Fisher Scientific, UK. Ethylene glycol dimethacrylate (EGDMA), azobisisobutyronitrile (AIBN), and polyvinyl alcohol (PVA, M$_w$ = 13,000–23,000, 87–89% hydrolyzed) were purchased from Sigma Aldrich, UK. All the reagents were of analytical grade. Pure water was supplied using a Millipore Milli-Q Plus 185 water purification system. All the gases were supplied by BOC, UK with a purity higher than 99.999%.

2.2. Particle synthesis

O/W emulsions were prepared using the dispersed phase composed of OA (template), AAm (functional monomer), EGDMA (crosslinker), and AIBN (initiator) dissolved in a mixture of TL and AN (porogenic solvents). The continuous phase was a 0.1–1 wt% aqueous solution of PVA. Formulation parameters and polymerization conditions are shown in Table 1 and in the caption of Fig. 7. The particle synthesis involved four major steps, as follows:
2.2.1. Step 1. Monomer-template self-assembly

The first step is self-assembly of functional monomer and template in a mixture of TL and AN and creation of monomer-template complex (Scheme 1, step I). Direct use of CO₂ as a template is impossible due to low solubility of CO₂ in organic solvents at ambient conditions. An alternative approach is to use a dummy template, here oxalic acid, which is a structural analog to the CO₂ molecule. Oxalic acid has a spatial structure very similar to two CO₂ molecules with their C atoms sitting back-to-back and O atoms pointing in opposite directions. The distance between adjacent NH₂ groups in the organic phase prior to polymerization was 0.45–0.7 nm [47] which is ideal for hydrogen bonding with CO₂ molecules, which have a kinematic diameter of 0.33 nm. OA and AAm were dissolved in the solvents and left overnight to ensure that molecular self-arrangement was fully completed.

2.2.2. Step 2. Suspension polymerization

The PVA solution was poured into a 500-mL jacketed reactor equipped with a four-neck lid and agitated using a four-bladed impeller with a diameter of 50 mm. (Fig. 1). The organic phase was prepared by dissolving EGDMA and AIBN in a pre-mixed oxalic acid-AAm solution and emulsified in the reactor. After purging the reactor for 10 min with N₂ to remove oxygen, nitrogen blanketing was used to prevent oxygen from diffusing into the reactor. The temperature was raised using a water-recirculating heater/chiller system to initiate the polymerization (Scheme 1, step II). The same procedure was used for non-imprinted polymer particles (NIPs), except that no template was added in the organic phase.

2.2.3. Step 3. Surfactant removal

The surfactant was removed from the product suspension using two agitation methods, mechanical and ultrasonic agitation. In both methods the temperature of the wash water was kept at 85–90 °C for 5 min, followed by water extraction and addition of fresh hot water for the next wash cycle. The particle morphology and size distribution were determined after 4, 8, and 10 wash cycles by scanning electron microscopy (SEM) and dynamic light scattering using a Malvern Mastersizer S (Malvern Instruments, UK) particle size analyzer.

2.2.4. Step 4. Template removal

The template was removed by washing the particles several times with a 1:9 (v/v) mixture of hydrochloric acid and methanol, until no oxalic acid was detected in the rinses by a Lambda 35 UV/VIS spectrometer (PerkinElmer, US). The extraction of the template from the particles resulted in formation of nanocavities decorated with covalently attached amide groups (Scheme 1, step III). In addition, mesopores were formed in the polymer matrix due to phase separation between the polymer and the solvents during the polymerization process.

2.3. Material characterization

2.3.1. Scanning electron microscopy (SEM)

The particle morphology was analyzed using a TM3030 benchtop scanning electron microscope (Hitachi, Tabletop Microscope Europe) operating at an accelerating voltage of 15 keV. The sam-

| Table 1 | Summary of emulsion formulation and polymerization conditions of samples synthesized for CO₂ capture capacity measurements. *
<table>
<thead>
<tr>
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<tbody>
<tr>
<td>Sample</td>
<td>AAm (g)</td>
<td>OA (g)</td>
<td>EGDMA (g)</td>
<td>ABIN (g)</td>
</tr>
<tr>
<td>s.b.1</td>
<td>2.04</td>
<td>0.9</td>
<td>9.51</td>
<td>0.12</td>
</tr>
<tr>
<td>s.b.2</td>
<td>5.1</td>
<td>2.27</td>
<td>23.8</td>
<td>0.3</td>
</tr>
<tr>
<td>s.b.3</td>
<td>10.2</td>
<td>4.54</td>
<td>47.6</td>
<td>0.6</td>
</tr>
<tr>
<td>s.b.4</td>
<td>10.2</td>
<td>–</td>
<td>23.8</td>
<td>0.3</td>
</tr>
<tr>
<td>s.b.5</td>
<td>5.1</td>
<td>2.27</td>
<td>23.8</td>
<td>0.3</td>
</tr>
<tr>
<td>s.b.6</td>
<td>5.1</td>
<td>2.27</td>
<td>23.8</td>
<td>0.3</td>
</tr>
<tr>
<td>s.b.7</td>
<td>5.1</td>
<td>2.27</td>
<td>23.8</td>
<td>0.3</td>
</tr>
<tr>
<td>s.b.8</td>
<td>5.1</td>
<td>2.27</td>
<td>23.8</td>
<td>0.3</td>
</tr>
<tr>
<td>s.b.9</td>
<td>10.2</td>
<td>4.54</td>
<td>23.8</td>
<td>0.3</td>
</tr>
<tr>
<td>s.b.10</td>
<td>5.1</td>
<td>4.54</td>
<td>23.8</td>
<td>0.3</td>
</tr>
<tr>
<td>s.b.11</td>
<td>5.1</td>
<td>–</td>
<td>23.8</td>
<td>0.3</td>
</tr>
<tr>
<td>s.b.12</td>
<td>5.1</td>
<td>–</td>
<td>23.8</td>
<td>0.3</td>
</tr>
<tr>
<td>s.b.13</td>
<td>5.1</td>
<td>–</td>
<td>23.8</td>
<td>0.3</td>
</tr>
<tr>
<td>s.b.14</td>
<td>10.2</td>
<td>–</td>
<td>47.6</td>
<td>0.6</td>
</tr>
<tr>
<td>s.b.15</td>
<td>5.1</td>
<td>–</td>
<td>23.8</td>
<td>0.3</td>
</tr>
</tbody>
</table>

* Tp = polymerization temperature; tp = polymerization time, φₚ = dispersed phase content; S_BET = specific surface area, V_p = total pore volume, d_p = average pore size. In all experiments the PVA concentration in the aqueous phase was 0.5 wt% and the stirring rate in the reactor was 200 rpm.

Scheme 1. A synthesis of molecularly imprinted polymer particles in this work. I: OA-AAm self-assembly via hydrogen bonding and formation of pre-polymerization complex in the organic phase; II: polymerization with EGDMA; III: template removal, IV/V: adsorption/desorption of CO₂ molecules within the nanocavity. Oxygen, carbon, and nitrogen atoms are shown in red, black, and blue, respectively, and green areas represent the hydrogen bonding interactions. (For interpretation of the references to colour in this scheme legend, the reader is referred to the web version of this article.)
samples were coated with gold/palladium (80/20) prior to the SEM to prevent accumulation of electrostatic charges on the particles. The sputtering rate was 0.85 nm per second at 2 kV and 25 mA of plasma current.

2.3.2. Particle size distribution

The particle size distribution was measured using a laser diffraction particle size analyzer, Malvern Mastersizer S (Malvern Instruments, UK). The mean particle size was expressed as the volume median diameter, $D_{v0.5}$, which is the diameter corresponding to 50% on the cumulative volume distribution curve. The degree of particle size uniformity was expressed as relative span:

$$\text{span} = \frac{D_{v0.9} - D_{v0.1}}{D_{v0.5}}$$

where $D_{v0.1}$ and $D_{v0.9}$ are the diameters on the cumulative volume distribution curve corresponding to 10% and 90%, respectively.

2.3.3. Pore size analysis

The pore size analysis was performed using a Micromeritics ASAP 2020 HD Accelerated Surface Area and Porosimetry System at 77 K in a liquid nitrogen bath. The samples were degassed at 80 °C overnight prior to the analysis. The specific surface area was calculated by the Brunauer–Emmett–Teller (BET) method over the relative pressure range, $P/P_0$, of 0.06–0.3. The mesopore size distribution was determined using the Barrett–Joyner–Halenda (BJH) model and the total pore volume was calculated from the amount of $N_2$ adsorbed at $P/P_0$ of 0.99. The size distribution of micropores and the total pore volume of micropores were measured using the Horvath–Kawazoe (HK) model at $P/P_0$ smaller than 10$^{-2}$.

2.3.4. Density measurement

The particle density was measured using a helium pycnometer 9200 (Micromeritics, US). The particles were dried overnight at 80 °C prior to the measurement. For each test, five runs were performed and the average value was reported.

2.3.5. Thermal analysis

The thermal stability of the CO$_2$-MIP particles was characterized using a thermogravimetric analyzer (TGA) Q5000IR (TA Instruments, US) under a nitrogen flow rate of 20 mL/min by heating 10–20 mg of sorbents from 50 to 600 °C at a ramp rate of 10 °C/min.

2.3.6. X-ray photoelectron spectroscopy (XPS)

XPS was performed on a K-ALPHA spectrometer (Thermo Scientific, UK) using monochromatic Al K$_\alpha$ X-rays (36 W, 400 μm spot size) and low-energy electron/ion flooding for charge compensation. Survey scan spectra for elemental analysis were obtained using a pass energy of 200 eV, a step size of 1 eV, a dwell time of 10 ms and 15 scans. High-resolution scans of the C 1s, O 1s and N 1s peaks were acquired using a pass energy of 50 eV and a step size of 0.1 eV with a dwell time of 50 ms and 10 scans.

2.3.7. Fourier transform infrared (FTIR) spectroscopy

The FTIR spectra were measured over the range of 500–4000 cm$^{-1}$ using a Thermo Scientific Nicolet iS50 ATR spectrometer with a monolithic diamond crystal. The samples were placed on the Universal diamond ATR top-plate and the spectrum was recorded within 32 s.

2.3.8. Dynamic CO$_2$ adsorption experiments

CO$_2$ adsorption on the synthesized particles was investigated in a fixed-bed column made of stainless steel (Fig. 1). In each test the column (15.8 mm O.D. and 9.25 mm I.D.) was loaded with 2.5–3 g of the particles sieved using a 0.075 mm stainless steel mesh. After loading, 0.035-mm stainless steel mesh and quartz wool plug were placed on either side of the bed to retain the particles. An in-house temperature controller was used to provide a uniform temperature along the column. The gas flow rate through the system was controlled using a mass flow controller (Alicat Scientific Inc, UK). Prior to each adsorption test, the particles were purged with $N_2$ for 2 h at 120 °C and 130 mL/min. The column was then sealed on both sides and left to cool down to the pre-determined adsorption temperature followed by blowing the feed stream composed of 15 vol% CO$_2$ and 85 vol% of $N_2$ through the bed at 130 mL/min. A CO$_2$ infrared analyzer (Quantek Instruments, USA) was used to continuously monitor the CO$_2$ concentration in the outlet stream. The CO$_2$ breakthrough curves were corrected by subtracting the mean resistance time of the gas stream (58 s at 130 mL/h and 40 °C), which was measured by flowing the feed gas mixture through the empty column.

3. Results and discussion

3.1. Production and characterization of particles

In O/W suspension polymerization, the organic phase, containing the template, the monomer, the cross-linker, and the initiator dissolved in the solvent(s), is dispersed in the aqueous surfactant solution. The droplets are converted to spherical polymer beads through a free radical polymerization. During the emulsification step, the surfactant forms a protective film at the interface preventing droplet coalescence. After polymerization, the surfactant must be removed to minimize particle agglomeration, since it results in a significant reduction in the surface area and CO$_2$ adsorption capacity. Fig. 2 shows the effect of two different particle agitation methods (ultrasound and mechanical) during washing on the morphology and size distribution of the particles after 4, 8, and 10 wash cycles. On the SEM images, it can be seen that after 8 and 10 wash cycles the particles agitated using ultrasound were considerably less agglomerated than mechanically agitated particles, and it was confirmed by the lower $D_{v0.5}$ and span values. After 10 ultrasound treatments, almost no agglomerated particles were detected. Therefore, particle washing using 10 repeated ultrasound treatments was implemented as a standard procedure in the subsequent work.

The physisorption data obtained from liquid nitrogen adsorption/desorption curves are presented in Figs. 3–5 and the corresponding specific surface areas, $S_{BET}$, specific pore volumes, $V_p$, and average pore sizes, $d_p$, are listed in Table 1. The majority of the samples follow a type IV nitrogen adsorption/desorption isotherm, featured by a hysteresis loop attributed to capillary condensation in mesopores (Fig. 3). Therefore, the material can be classified as a mesoporous adsorbent. However, for s.b.9 despite the presence of a slight hysteresis loop, the pore size distribution was wide and went beyond 50 nm implying a Type II isotherm profile. Compared to bulk polymerization [47], suspension polymerization led to more uniform mesopore size distribution. In addition, all samples synthesized by bulk polymerization followed a Type II isotherm with an average pore size of 10–24 nm, as compared with 5–15 nm found in this work (Table 1). The $S_{BET}$ values in this work were 316–457 m$^2$/g, compared with 80–258 m$^2$/g reported in bulk polymerization. Moreover, the maximum pore volume of 0.92 cm$^3$/g for the sample s.b.9 (Table 1) was considerably higher than the maximum $V_p$ value in bulk polymerization.

Since imprinted nanocavities are smaller than 2 nm, the Horvath–Kawazoe (HK) method was used to compare the size distribution of the micropores ($d < 2$ nm) in the MIP sample s.b.10 and the NIP material s.b.4, Fig. 5. Both samples were prepared under the same conditions, except that no template was used in s.b.4.
There were two sharp distinct peaks for s.b.10 at 0.84 and 0.85 nm and both were significantly larger than the ones for s.b.4, implying the presence of nanocavities in the MIP matrix. In addition, the total pore volume in the micro-range (pore diameter <2 nm) was 0.1 cm³/g for s.b.10 and only 0.06 cm³/g for s.b.4. The better structural properties of s.b.10 as compared with s.b.4 were confirmed by CO₂ capture measurements. The CO₂ capture capacity of MIP particles was 2.5 times higher than that of NIP particles (Fig. 8c), which can be attributed to the nanocavities in the imprinted polymer matrix.

To further verify the better performance of MIPs, five additional MIP/NIP pairs were analyzed. The performance of MIPs was assessed using the standard protocol based on the imprinting factor, IF, which is the ratio of equilibrium adsorption capacity of MIP to its corresponding NIP particles [51,52]. For the five MIP/NIP pairs (s.b.2 vs. s.b.12, s.b.3 vs. s.b.14, s.b.8 vs. s.b.13, s.b.9 vs. s.
b.15, and s.b.11 vs. s.b.12), the CO₂ capture capacity was measured (Fig. 8), and the corresponding IFs were found to be 2.3, 1.9, 1.5, 3.7, and 1.3 respectively. In all cases, IF was greater than 1, implying higher binding affinity of MIPs compared to NIPs. Since MIPs synthetized using bulk polymerization had an IF of 1.1 [47], it is clear that the current MIPs produced by suspension polymerization shows a higher binding affinity towards CO₂ molecules.

Although the pore volume in the micropore range for the MIPs is larger than that for the NIPs, the total pore volume and S BET surface area can be mainly attributed to the presence of mesopores formed due to phase separation between the polymer and the solvents during the polymerization process, particularly for polymeric particles with large pore volumes. Although the role of template in the creation of nanocavities is known, the effect of template on the structure of mesopores is not well understood and is beyond the scope of this study. Comparing the total pore volume and S BET surface area of MIP samples and their NIP counterparts in Table 1, no specific trend was observed, and both smaller and larger pore volumes of NIPs were found in some cases. The same observation was reported by Dirion et al. [51], who developed a library of
MIP sorbents for use in pure aqueous environments. Therefore, it might not be possible to evaluate the separation performance of MIP and NIP particles by comparing their total pore volume and SBET surface area.

Fig. 6 shows the TGA curves for selected samples. All TGA curves have similar patterns with no weight loss up to 220°C and the thermal decomposition temperature corresponding to 5% weight loss occurring at 240–255°C. The average density of the particles measured using a multivolume helium pycnometer was 1.3 g/cm³.

3.2. Particle size control

Inappropriate particle size can considerably limit the applicability of solid sorbents. Particles with very small sizes can cause large pressure drops in fixed bed reaction systems [53]. In fluidized bed reactors, small particles belonging to Group C according to the Geldart classification [54] are prone to cohesion, which may result in channeling at high flue gas flow rates, and reduced capture efficiency [55]. On the other hand, large particles (Group D) have a low surface area-to-volume ratio, resulting in a lower diffusion rate of CO₂ within the particle. Therefore, particles for CO₂ capture in fluidized beds should be large Group A particles or small Group B particles [55,56]. Based on the particle density in this work (1.3 g/cm³), their smallest size must be about 120 μm to fall under these categories. The particle size was controlled by changing the agitation rate in the reactor and the concentration of surfactant in the aqueous phase. The effect of agitation speed on the mean droplet size at two different dispersed phase contents, φ₀, is shown in Fig. 7a and b. At φ₀ = 9%, an increase in stirring rate from 100 to 600 rpm caused a decrease in the droplet size from 171 to 73 μm. At φ₀ = 20%, an opposite trend was observed and an increase in agitation speed led to an increase in the droplet size. In stirred vessels, there are two factors that determine the particle size of an emul-
sion: droplet breakage due to viscous and/or inertial forces, and droplet coalescence, due to droplet collision. At low \( \phi_v \) values, the droplet breakup is a dominant factor controlling the droplet size, and an increase in the agitation speed results in higher shear forces exerted by the impeller on the droplets and their fragmentation into smaller droplets. At higher \( \phi_v \) values, droplet coalescence dominates over droplet breakup and consequently leads to larger droplets at higher stirring rates. The latter behavior is common in industrial suspension polymerization processes, where the dispersed phase content is relatively high [50]. At \( \phi_v = 9 \text{ vol\%} \), the particles fabricated at 100–300 rpm were rather uniform in size with a span of 0.6–0.91 (Fig. 7a). On the other hand, at 400–600 rpm the particles were highly polydispersed with a span of up to 2.2. At \( \phi_v = 20 \text{\%} \), the particle size distribution was independent of the stirring rate and the span varied between 0.77 and 1.29.

The effect of the concentration of PVA in the continuous aqueous phase on the particle size is shown in Fig. 7c. As expected, the median particle size decreased on increasing the PVA concentration from 0.1 to 1 wt\%. At the higher PVA concentration, the amount of surfactant adsorbed on the surface of the droplets is higher, which reduces the interfacial tension and lowers the required energy for droplet disruption. Therefore, for a constant energy input, smaller droplets were formed at higher PVA concentration. The most uniform droplets (span = 0.77) were produced at 0.5 wt\% PVA, which is a typical surfactant concentration in suspension polymerization. Also, at 1 wt\% PVA in the aqueous phase it is more difficult to remove PVA after polymerization. Therefore, 0.5 wt\% was considered as the optimum PVA concentration.

3.3. CO\textsubscript{2} adsorption capacity of synthesized particles

The CO\textsubscript{2} adsorption capacity, \( q \), of the samples from Table 1 was measured in the fixed bed column shown in Fig. 1 and calculated using the following equation [57]:

\[
q = \frac{QC_{t_{ad}}}{m_s} \quad (2)
\]

where \( q \) (mmol/g) is the CO\textsubscript{2} adsorption capacity, \( Q \) (mL/min) is the feed gas flow rate upstream of the column, \( m_s \) (g) is the mass of the particles in the column, and \( C_i \) (mmol/mL) is the concentration of CO\textsubscript{2} in the feed stream calculated using Eq. (3):

\[
C_i = \frac{n_i P}{RT} \quad (3)
\]

where \( n_i \) is the molar fraction of CO\textsubscript{2} in the feed gas (\( n_i = 0.15 \)), \( P \) (kPa) is the total pressure of the gas stream (~102 kPa), \( T \) (K) is the adsorption temperature, \( R \) is the universal gas constant, 8.314 J/(K\textsubscript{mol}), and \( t_{ad} \) is the stoichiometric time, which is equal to the breakthrough time in an ideal adsorption column with a vertical breakthrough curve, i.e., in a column operating at 100% CO\textsubscript{2} removal efficiency until the particles are fully saturated with CO\textsubscript{2}. The \( t_{ad} \) value corresponds to the area between the breakthrough curve and the vertical axis \((C_o/C_i)\) and can be calculated as:

\[
t_{ad} = \int_0^t \left(1 - \frac{C_o}{C_i}\right)dt \quad (4)
\]

where \( C_o \) (mmol/mL) is the CO\textsubscript{2} concentration in the effluent stream and \( t \) (min) is the time. The CO\textsubscript{2} adsorption capacity for different samples over the range of \( C_o/C_i \) from 0 to 1 is shown in Fig. 8. The particles were evaluated based on their equilibrium CO\textsubscript{2} adsorption capacity, \( q_{eq} \), which is a \( q \) value corresponding to \( C_o/C_i = 1 \). It occurs when the effluent concentration reaches the feed gas concentration, and the material is no longer capable of adsorbing CO\textsubscript{2}.

The effect of polymerization temperature over the range of 50–70 °C is shown in Fig. 8b. To achieve a sufficient decomposition rate of AIBN, the polymerization temperature should be at least 50 °C. At the start of the polymerization process, individual polymer particles (nuclei) grow separately inside organic phase droplets. These nuclei eventually agglomerate via inter-nuclei crosslinking and the final porous bead is formed. The higher the polymerization temperature, the higher the polymerization rate [58] and the larger the number of nuclei formed. The size of individual polymer particles in the bead decreases on increasing number of nuclei, which...
should result in higher surface area of the bead. Indeed, the highest surface area in Table 1 (457 m²/g) corresponds to the beads synthesized at 70 °C. However, higher temperatures can weaken monomer-template interactions and reduce the quality of active sites in the polymer network. As a result, \( q_{eq} \) had similar values for the particles synthesized at 60–70 °C (0.26–0.29 mmol/g).

The effect of molecular imprinting on CO₂ adsorption capacity is shown in Fig. 8c for samples s.b.10 and s.b.4 prepared under the same conditions except that oxalic acid was added in the organic phase in sample s.b.10. As a result, \( q_{eq} \) for s.b.10 was 0.42 mmol/g, as compared to 0.17 mmol/g for the NIP sample s.b.4, which confirms that molecular imprinting improves CO₂ adsorption capacity. The high affinity of MIP particles towards CO₂ can be attributed to a combination of size exclusion effect on gas molecules, which is based upon their size and shape, and dipole-dipole interactions between CO₂ molecules and CO₂-philic NH₂ moieties inside the cavities. The cavity architecture (size and shape) facilitates the entrapment of CO₂ molecules in the polymer matrix due to structural compatibility of the cavities and CO₂ molecules. In addition, larger quadrupole moment and polarizability of CO₂ molecules compared to N₂ molecules (2.85 to 1 and 1.5 to 1, respectively) enhance the affinity of protic NH₂ groups on the cavity walls towards the CO₂ molecules leading to higher CO₂ uptake.

The effect of template concentration and solvent ratio in the organic phase on CO₂ adsorption capacity can be seen in Fig. 8c. When the monomer-to-template ratio was reduced from 2.25:1 to 1.12:1, a decrease in \( q_{eq} \) was observed (samples s.b.11 and s.b.2), probably due to a smaller number of NH₂ groups per cavity. When the volume ratio of AN to TL was reduced from 50:50 to 30:70, an increase in \( q_{eq} \), \( S_{BET} \), and \( V_p \) was observed (samples s.b.2 and s.b.9). It is known that apolar, non-protic solvents, such as toluene, stabilize hydrogen bonds and maximize the likelihood of complex formation between the template and the monomer, which leads to a larger number of cavities and enhanced affinity of MILPs towards the target molecules [59]. A polarity index of TL (2.4) is lower than that of AN (5.8), which allows stronger template-monomer interactions in a pre-polymerization solution. On the other hand, toluene is considered as a bad solvent for AAm and a certain amount of good solvent (AN) must be added to ensure that the organic phase is homogeneous. Increasing the level of bad solvent can cause early phase separation between the polymer and the solvents which results in aggregation of copolymer nuclei into larger clusters [59,60], hence, the pore size and pore volume increase (Fig. 4c and Table 1). Increased particle porosity can facilitate the diffusion of CO₂ through the polymer matrix, resulting in a higher CO₂ adsorption capacity for sample s.b.9 compared with s.b.2 (Fig. 8c).

Fig. 8d shows the effect of polymerization time on the CO₂ adsorption of synthesized particles over the range of 1–6 h. The difference in \( q_{eq} \) for the samples produced after 3 and 6 h was negligible (0.29 and 0.30 mmol/g, respectively). However, \( q_{eq} \) for the particles prepared after 1 h was 0.22 mmol/g, due to low conversion of AAm and a loss of active sites. Since the difference in \( q_{eq} \) for the particles synthesized after 3 and 6 h was negligible, 3 h was considered sufficient time for polymerization and used for the majority of samples in Table 1.

The effect of dispersed phase content, \( \phi_o \) on CO₂ adsorption capacity is shown in Fig. 8a. An increase in \( \phi_o \) from 9 to 33% resulted in an increase in \( q_{eq} \) from 0.17 to 0.32 mmol/g. It can be explained by the lower amount of acrylamide diffused from the organic phase droplets to the aqueous phase during polymerization, leading to the higher amount of NH₂ groups in the polymer network and the higher number of cavities in the polymer matrix. The diffusion of AAm was suppressed at higher \( \phi_o \) values, because of the smaller amount of aqueous phase in the reactor. The mass fraction of nitrogen on the surface of the particles was quantified using XPS, as shown in Fig. 9a. In the samples prepared under the same operating conditions and using the same organic phase (s.b.1, s.b.2, and s.b.3), the nitrogen content increased on increasing \( \phi_o \) due to an increasing amount of NH₂ groups on the particle surface. Therefore, the main parameter affecting the CO₂ adsorption capacity is the density of NH₂ active sites on the surface, rather than specific surface area or total pore volume. This conclusion can be confirmed by comparing the nitrogen content and \( q_{eq} \) for two samples, s.b.10 and s.b.2, prepared under the same conditions, but the concentration of template and monomer was doubled in s. b.2. As shown in Fig. 9, the nitrogen fraction of s.b.10 was 2.4 times larger than that in s.b.2 and as a result, \( q_{eq} \) was 1.5 times larger than that of s.b.10 (Fig. 8c). The same trend was reported in bulk polymerization with higher surface density of NH₂ groups leading to higher CO₂ adsorption [47].

The FTIR spectra of s.b.1, s.b.2, s.b.3, and s.b.10 samples are shown in Fig. 9b. The peak at around 3440 cm⁻¹ on all spectra Fig. 9. Surface chemical analysis and FTIR analysis of samples b.1, b.2, b.3 and b.10: (a) XPS spectra and the mass percent of carbon, oxygen, and nitrogen; (b) FTIR spectra.
can be attributed to amine N–H stretching vibrations. The most intense N–H stretching peak was obtained for s.b.3, followed by s.b.10, s.b.2, and s.b.1, which corresponds to the fraction of nitrogen on the surface found by XPS. Almost the same sequence of peaks was observed at 1663 cm$^{-1}$ which can be attributed to N–H bending vibrations. The two peaks at 1150 cm$^{-1}$ and 1250 cm$^{-1}$ are due to C–N stretching vibration. Therefore, the results imply the presence of amine groups on the surface of synthesized adsorbents. The peak at 1730 cm$^{-1}$ can be attributed to C=O bonds in AAm and EGDMA, and the peaks at 2850–3000 cm$^{-1}$ are due to C–H stretching vibration. There is no peak in the range of 1680–1640 cm$^{-1}$ corresponding to C=C stretch in EGDMA and AAm, meaning there is no monomer or crosslinker left in the polymer particles and all carbon-carbon double bonds are broken. The peaks over the range of 910–665 cm$^{-1}$ could be attributed to N–H wagging vibration.

Although the CO$_2$ adsorption capacity is higher at higher AAm concentration in the organic phase, AAm partitions between aqueous and organic phase. Excessive concentration of AAm in the aqueous phase results in gel formation in the continuous phase and failure of particle synthesis. For each $\phi_c$ value it is vital to find the highest possible concentration of AAm in the organic phase that will not cause gelation of the aqueous phase. A phase diagram of $\phi_c$ vs. $C_{AAm}$ is shown in Fig. 10, and the safe region, where no gelation occurred, is highlighted in green.

The effect of temperature on $q$ for s.b.9 is shown in Fig. 11. As expected, a considerable decrease in $q_{eq}$ from 0.62 mmol/g to 0.28 mmol/g was observed, as the temperature increased from 25 $^\circ$C to 80 $^\circ$C, which can be attributed to the exothermic nature of the adsorption process and weakened hydrogen bond interactions between CO$_2$ and NH$_2$ groups at higher temperatures. In comparison with the existing polymeric adsorbents at low CO$_2$ partial pressures, the current MIPs showed very good CO$_2$ capture capacity, which was larger than the maximum CO$_2$ capture capacity of conjugated microporous polymer (CMP) networks developed by Dawson et al. [61] (0.3 mmol/g at 0.15 bar and 25 $^\circ$C), microporous organic polymers (MOPs) produced by Kaliva et al. [62] (0.2 mmol/g at 0.15 bar and -5 $^\circ$C), and azo-covalent organic polymers (azo-COPs) synthesized by Patel et al. [63] (0.4 mmol/g at 0.15 bar and 25 $^\circ$C). The CO$_2$ capture capacity of the current MIPs was comparable to that of nanoporous organic polymers (NOPs) reported by Chen et al. [64] (0.63 mmol/g at 0.15 bar and 25 $^\circ$C), but lower than that of sulfonate-grafted porous polymer networks (PPNs) reported by Lu et al. [65] (1.12 mmol/g at 0.15 bar and 25 $^\circ$C).

![Phase diagram of acrylamide gelation based on AAm concentration in the organic phase and volume percent of the dispersed phase in emulsion.](image)

### 4. Conclusions

Novel molecularly imprinted poly(AAm-co-EGDMA) spherical particles were synthesized by suspension polymerization in an oil-in-water emulsion and used for CO$_2$ adsorption from a CO$_2$/N$_2$ gas mixture. The ability of the particles to adsorb CO$_2$ was closely related to their hierarchical porous structure characterized by the mesopores formed by phase separation and the nanocavities smaller than 1 nm with CO$_2$ recognition properties formed by molecular imprinting. The nitrogen adsorption-desorption isotherms for most of the samples were of type IV, featuring uniform pore size distribution and a BET surface area up to 457 m$^2$/g, which facilitated the diffusion of CO$_2$ through the polymer matrix and improved accessibility of the cavities to CO$_2$ molecules. The poor correlation between CO$_2$ capture capacity and total volume of the mesopores reflected the fact that the CO$_2$ adsorption capacity depended primarily on the quality and number of nanocavities, which were too small (d < 1 nm) to be detected by the BET analysis but their presence was revealed by employing the HK method.

Higher contents of functional monomer (AAm) and low-polar solvent (TL) in the organic phase prior to polymerization led to higher CO$_2$ capture capacity of the particles due to stronger hydrogen bonding interactions between the template and the monomer during complex formation and larger number of CO$_2$–philic NH$_2$ moieties in the polymer network. The fraction of nitrogen on the surface of the particles increased by increasing the dispersed phase content in the reactor and the concentration of acrylamide, but excessive acrylamide concentration and/or dispersed phase content led to gelation of the continuous phase and failure of the polymerization process.

The particle size was controlled by changing the agitation rate in the reactor and the surfactant concentration in the aqueous phase. The particles exhibited good thermal stability with a 5% weight loss at 240–255 $^\circ$C and the maximum equilibrium CO$_2$ adsorption capacity of 0.56 and 0.62 mmol/g at 40 $^\circ$C and 25 $^\circ$C, respectively, and 0.15 bar CO$_2$ partial pressure. The particle size and density were within the range suitable for applications in fixed bed and fluidized bed adsorption systems. In addition, suspension
polymerization approach significantly facilitates the production scale-up, compared to bulk polymerization.

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Appendix A. Supplementary data

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References


