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Synthesis and functionalisation of spherical meso-, hybrid meso/macro- and macro-porous cellular silica foam materials with regulated pore sizes for CO₂ capture†

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A variety of meso, meso/macro and macro-structured siliceous cellular foam (SCF) materials have been tailor-designed and fabricated using a modified microemulsion templating methodology with trimethyl benzene (TMB) as the pore expander and Pluronic™ block co-polymer (P123) as the surfactant for preparing polyethyleneimine (PEI)-impregnated adsorbents for CO₂ capture. The effect of preparation conditions, such as the TMB/P123 mass ratio, aging temperature and aging time, on the SCF morphology and pore structures and hence on the CO₂ adsorption performance of the PEI-modified SCF adsorbents was investigated comprehensively. BET measurements and morphological characterisation with SEM revealed that the SCF materials prepared using lower TMB/P123 ratios (≤ 1) and aging temperatures (≤ 100 °C) were typically meso-structured with relatively lower cell wall thicknesses but increasing the TMB/P123 ratio, aging temperature and aging times led to a transformation of the SCFs from being meso-structured into hybrid meso/macro or even purely macro-structured nano-cellular materials with increased wall thicknesses, pore volumes and window sizes. CO₂ adsorption characterisation of the PEI-impregnated SCFs demonstrated that while all the SCF materials exhibited higher capacities and faster adsorption kinetics compared to conventional meso-structured siliceous materials, the hybrid meso/macro and macro-structured SCF substrates were found to have the best CO₂ adsorption performance, with uptake capacities reaching 180.2 mg-CO₂ per g-adsorbent (5.85 mmol per g-PEI) for PEI-600 impregnation and 198.2 mg-CO₂ per g-adsorbent (6.44 mmol per g-amine) for the hybrid impregnation of PEI-600-TEPA at 75 °C and 0.15 bar CO₂, which are significantly higher than those previously reported under similar conditions. The macro- and hybrid meso/macro-structured SCF materials were found to be particularly suitable for preparing high molecular weight PEI-modified adsorbents for greatly improved thermo-stability. At 60 wt% PEI loading, the CO₂ capacity reached 126 and 97.3 mg-CO₂ per g-ads for PEI-10 000 and PEI-60 000, respectively, both showing extraordinary lifetime performance. Differing from previous findings, no particularly favourable pore diameters or window sizes for PEI impregnation are observed for the wide range of SCF materials examined, although close to linear relationships between the CO₂ uptake capacity and total pore volume appear to exist for the SCF materials with pore volumes below 2.2 cm³ g⁻¹ and pore diameters/window sizes ≤ 28 nm.

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1. Introduction

Carbon capture and storage (CCS) has widely been recognised as being a crucial part of worldwide low carbon energy portfolios required to meet the target of global climate change without compromising energy security, given the projected soaring

energy demand and the continual dominance of fossil energy in the worldwide energy landscape.^{1–4} However, the successful development and deployment of enabling capture technologies hold the key to the overall success of CCS as carbon capture represents the most costly single element of the whole CCS chain, which goes from CO₂ capture through compression and transport to geological storage. At present, aqueous amine scrubbing in its various forms is the state-of-art capture technology for use, but the high energy penalty together with a range of environmental and operational issues (e.g. solvent degradation, equipment corrosion and toxic and corrosive emissions) has become the major challenge for the large scale deployment

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of this technology for CCS.⁵ It has been estimated that post-combustion capture (PCC) with aqueous amine scrubbing could increase the cost of power generation by 32% and 65% for gas and coal-fired power plants, respectively. Consequently, the development of alternative cost-effective PCC technologies has received tremendous attention over recent years, with the adsorption-based solid adsorbent looping technology having become the persistent focus of many investigations due to the potentially significantly reduced energy penalty and the choice of a wide range of candidate adsorbent materials, including but not limited to zeolites,^{6,7} immobilized amines,^{8,9} activated carbons,^{10,11} metal-organic frameworks (MOFs),^{12,13} covalent-organic frameworks (COFs)^{14,15} and zeolitic imidazolate frameworks.^{16,17} Among the candidate sorbent materials, immobilized amines possess many desirable CO₂ adsorptive properties that may not be easily obtained with other sorbent materials, such as their extremely high CO₂ selectivity, fast adsorption kinetics and high adsorption capacity especially at very low CO₂ partial pressures, great moisture tolerance and favourable operating temperature window of 50–80 °C that minimizes or essentially eliminates the cooling need of typical flue gas streams from power plants.^{18–23}

The CO₂ adsorption of immobilised amine adsorbents follows similar reaction mechanisms to those in the aqueous amine absorption process, which involves the reversible formation of ammonium carbamate under anhydrous flue gas conditions and ammonium bicarbonate under wet conditions. Two typical preparation methodologies are used, namely surface grafting and direct impregnation. In general, amine-grafted silica sorbents benefit from improved amine efficiency and thermal stability but usually suffer from low amine loadings due to the limited density of accessible surface silanol groups, giving rise to CO₂ adsorption capacities being generally well below 3 mmol CO₂ per g sorbent (hereinafter abbreviated to mmol g⁻¹) or 13.2 wt%. The highest adsorption capacity of *ca.* 4 mmol g⁻¹ (17.6 wt%) was obtained by Dreser *et al.*²⁴ at 75 °C in 10% CO₂/N₂ for a hyperbranched aminosilica with a grafted amine loading of 9.78 mmol amine per g SBA-15, whereas three-dimensional macro-porous silica with a grafted amine density of 10.98 mmol N per g could only achieve a maximum CO₂ capacity of 2.76 mmol g⁻¹ under similar conditions.²⁵ In comparison, impregnation as a more convenient and less corrosive preparation methodology can achieve much larger amine loading capacity but the produced sorbents may suffer from the potential evaporation loss of the impregnated amines, an issue that can be improved by using higher boiling amines.²⁶ Builes and Vega²⁷ compared the performance of immobilised amines prepared from different methodologies and found that compared to grafting, impregnation persistently gives rise to significantly higher adsorption capacities particularly at low CO₂ partial pressures, due to the mobility of the functionalised amine chains that leads to the formation of microporosity-like micro-cavities inside the bulk phase of the impregnated amines. In contrast, similar structures cannot be formed within the densely grafted amine groups and this can limit the potential of achieving higher adsorption capacities by increasing the grafting density.

Numerous investigations have shown that for both impregnation and grafting, the performance of supported amine sorbents is not only determined by the chemical nature of the amines but also by the type of porous solid support, with mesoporous silica materials such as MCM-41, MCM-42, MCM-48, SBA-15, SBA-16 and hexagonal mesoporous silica (HMS) being the most widely used solid supports because of their tuneable structures and re-addressable surface chemistries. Son *et al.*²⁸ studied the CO₂ adsorptive properties of different mesoporous silica materials at the same maximum PEI loading of 50 wt% and found that dictated by the average pore diameter of the solid support, the CO₂ uptake of PEI-modified silica sorbent follows the order of KIT-6 > SBA-16 ≈ SBA-15 > MCM-48 > MCM-41, with the capacities varying from 2.5 mmol g⁻¹ for MCM-41 with a 1-dimensional (1D) pore diameter of 2.8 nm to 3.1 mmol g⁻¹ for KIT-6 with a 3D pore diameter of 6.5 nm. Over recent years, many efforts have been made to expand the pore diameter or increase the pore volume of mesostructured silica supports to increase the amine loading and hence CO₂ uptake capacity. It has been found that regardless the preparation techniques employed, the immobilised amine sorbents prepared using pore-expanded MCM-41,^{29–32} MCM-48,²⁹ SBA-15^{29–32} and HMS^{33,34} all showed considerably higher CO₂ adsorption capacities than their pore-unexpanded counterparts. In addition to the pore size and volume, the length of pore channels of support materials also appears to play a vital role in determining the CO₂ adsorption capacity and kinetics of amine-modified silica adsorbents due to the mass transfer limitations within extended meso-channel networks. Studies found that pore-expanded mesoporous silica supports with shorter meso-channels can outperform those with longer channels by a factor of 2.7 and 1.45 for PEI-impregnated MCM-41 and SBA-15 in pure CO₂ at 50 °C³⁰ and a factor of 2.2 for amine-grafted SBA-15 at 75 °C in 15% CO₂/N₂,³⁵ respectively. However, despite the efforts in recent years, very few of the reported amine-modified conventional mesoporous silica materials have CO₂ capture capacities greater than 3 mmol g⁻¹ or 13.2 wt% in 15% CO₂/N₂ at the desirable adsorption temperature range of 50–75 °C, due to amine loading limitations. Previous studies have demonstrated that for post-combustion capture, solid amine adsorbents with a CO₂ capture capacity of *ca.* 3 mmol g⁻¹ for 50 wt% PEI loading are equivalent to 20 wt% aqueous MEA solvent when 90% of the CO₂ in typical flue gas streams was captured.^{36,37} This suggests that more effective solid supports need to be explored to further improve the adsorption performance of immobilised amine adsorbents.

Clearly, a desirable solid support should best have large pore volume coupled with superior interconnectivity among the pores to maximise amine loading whilst reducing the mass transfer limitations of adsorption. Spherical mesocellular siliceous foam (MCF) represents a relatively new class of mesoporous silica materials with a continuous network comprised of uniform micro-spherical cells interconnected by large window pore channels, giving rise to a rigid macro-spherical geometry with a 3-dimensional porous framework with large pore size, pore volume and surface area. Compared with other mesoporous materials which usually consist of non-intersecting



tubular pores, the unique geometrical configuration of MCF makes it a desirable category of support materials for preparing immobilised amine CO_2 sorbents. Earlier investigations have demonstrated that mesocellular silica foam-supported PEI sorbents can significantly outperform other mesoporous silica materials by up to 30% for CO_2 adsorption, with capacities reaching up to 3.5 mmol CO_2 per g at 75 °C and 0.15 bar CO_2 ,^{38–40} compared to virtually all other mesoporous silica-supported PEIs with capacities being generally below 2.5 mmol CO_2 per g under similar conditions.^{28,35} To date, however, the impact of preparation and precursor chemistries on CO_2 adsorption has not been examined adequately. In this study, a variety of siliceous cellular foam materials with varying cellular networks from meso-structured to macro-structured were prepared with different precursor and preparation chemistries, and their performance as the candidates for preparing high capacity CO_2 adsorbents was evaluated.

2. Experimental

2.1 Chemicals

Non-ionic tri-block copolymer surfactant Pluronic P123 (EO20-PO70-EO20, MWav = 5800), HCl (37%), 1,3,5-trimethylbenzene (TMB), tetraethyl orthosilicate (TEOS, 99.999%), ammonium fluoride (NH_4F , 99.99%), and different branched polyethylenimines with average molecular weights (MWs) varying from 600 to 60 000 were all obtained from Sigma-Aldrich. PQ silica was obtained from PQ Corporation.

2.2 Preparation of siliceous cellular foams and polyamine-impregnated CO_2 adsorbents

Porous siliceous cellular foams (SCFs) with variable pore and window sizes were synthesised using a modified version of the micro-emulsion method reported earlier by Schmidt-Winkel and co-workers,⁴¹ with the amphiphatic surfactant P123 as the template and TMB as the pore-expanding agent. Briefly, in a typical preparation with TEOS as the silicon source and a TMB/P123 mass ratio of 1 : 1, 8 g of P123 was first dissolved in 150 ml of 1.5 M HCl at 40 °C and then 8 g of TMB was added. Following 40 min of further stirring, 18.4 ml of TEOS and 10 ml of 2.5 M NH_4F solution (F/Si molar ratio 0.03) were added to the mixture under vigorous stirring and the resultant solution was kept at 40 °C for variable times with continuous stirring. During this process, the hydrolysis of the TEOS at the surface of P123-coated TMB droplets leads to the formation of a composite shell material with interconnected cells and windows at the areas of contact of adjacent droplets. The formed milky reaction mixture was then transferred into an autoclave and kept at 100 or 120 °C for variable times under static conditions. The as-prepared white precipitate was separated by filtration and dried at ambient temperature. The dried precipitate was then calcined in air at 550 °C for 8 hours to obtain the porous silica support. The effect of preparation parameters on the surface textural properties of silica supports and hence the CO_2 adsorptive properties of their supported polyamine sorbents was investigated, including the TMB/P123 ratio, aging time and aging

temperature. For clarity and easy comparisons, the SCF materials synthesised under different conditions are denoted as SCF- x - y - z where x , y and z represent the TMB/P123 ratio, aging temperature and time, respectively. Table 1 summarises the preparation parameters examined.

Amine functionalisation of the SCF materials was achieved using a modified wet impregnation method. In this modified methodology, water rather than alkanols^{30,38,42} was used for the first time as the more cost-effective solvent. In a typical preparation, a calculated amount of selected polyamines was first dissolved in 10 ml of water under continuous stirring for 20 min, and then the corresponding amount of SCF support was gradually added to the solution. After overnight stirring, the mixture was dried in a vacuum at 50 °C to yield PEI-functionalised SCF adsorbents. A branched PEI with an average molecular weight (MW) of 600, denoted as PEI-600, was used in most preparations but for selected SCF samples, other amines were also examined, including tetraethylenepentamine (TEPA) and other PEIs with variable MW from 1800 to 60 000. All the polyamines were obtained from Sigma-Aldrich. The PEI contents of the prepared SCF adsorbents were determined by TGA as the weight loss of the prepared SCF adsorbents when they were heated to 700 °C and burned in air (100 $\text{cm}^3 \text{ min}^{-1}$) at the temperature to a constant weight. It was found that the weight losses obtained as the PEI contents agreed well with the pre-defined contents used in the impregnation, confirming the validity of the methodology.

2.3 Characterisation of SCF materials

The textural properties of the synthesised SCF materials were determined using nitrogen sorption isotherms with a Micromeritics ASAP 2420 apparatus at 77 K. In a typical measurement, the silica was first degassed at 120 °C for 12 hours before analysis. Specific surface areas were determined using the Brunauer–Emmett–Teller (BET) equation. In addition, the pore

Table 1 Summary of preparation conditions for different SCF materials

Sample	Initial P123 concentration, wt%	TMB/P123 mass ratio	Aging temperature, °C	Aging time, hours
SCF-0-100-24	5.0	0	100	24
SCF-0.4-100-24	5.0	0.4	100	24
SCF-0.7-100-24	5.0	0.6	100	24
SCF-1-100-24	5.0	1	100	24
SCF-1-70-24	5.0	1	70	24
SCF-1-80-24	5.0	1	80	24
SCF-1-90-24	5.0	1	90	24
SCF-1-100-12	5.0	1	100	12
SCF-1-100-42	5.0	1	100	42
SCF-1-100-60	5.0	1	100	60
SCF-1-120-42	5.0	1	120	42
SCF-2-100-24	5.0	2	100	24
SCF-2-100-42	5.0	2	100	42
SCF-3-100-24	5.0	3	100	24
SCF-3-100-42	5.0	3	100	42
SCF-3-120-24	5.0	3	120	24



volume, pore size distribution, cell pore size and window size of the mesostructured silica were determined by using the Barrett–Joyner–Halenda (BJH) method.

A JEOL 7100F Field Emission Gun Scanning Electron Microscope (FEG-SEM) (JEOL USA, Inc.) was used to illustrate the morphology and microstructure of the SCF samples while high resolution transmission electron microscopy (TEM) images were taken on a JEOL 2100F FEG-TEM at 200 kV.

2.4 Characterisation of PEI-modified SCF adsorbents

The CO_2 adsorption–desorption performance of PEI-modified SCF adsorbents was evaluated with a TA500 thermogravimetric analyser (TA Instruments), using a simulated flue gas stream containing 15% CO_2 in N_2 . The amount of sample used in each TGA test was typically around 30 milligrams (mg) for different samples. Prior to any measurement, each individual sample was first subjected to pre-treatment at 120 °C in N_2 for 20 min to remove any remaining moisture and/or pre-adsorbed gases, and the sample was then allowed to cool down to the selected temperature for adsorption tests. Three different types of CO_2 adsorption measurements were performed, including (1) temperature-programmed CO_2 adsorption, (2) CO_2 adsorption isotherms at selected temperatures, and (3) the cycle lifetime performance testing of the sorbents.

For temperature-programmed adsorption, the sample was cooled down to 30 °C when the gas supply was switched from N_2 to 15% CO_2 in N_2 . The sample was first held at this temperature for 30 min to reach equilibrium CO_2 adsorption before the

temperature was allowed to ramp to 130 °C at an extremely slow heating rate of 0.1 °C min^{−1}. The optimal adsorption temperature determined from the temperature-programmed adsorption test protocol was then selected to evaluate the CO_2 -capture performance of individual sorbent samples prepared. CO_2 adsorption capacity was taken as the weight gain after the sample was exposed to the CO_2 -containing gas flows and reached equilibrium adsorption. The adsorption measurement was repeated typically three times for each sample or set of conditions and the average value was used.

Cyclic adsorption–desorption tests were also carried out. In each cyclic test, the sample was first allowed to reach equilibrium adsorption at 75 °C in the simulated flue gas and with the gas supply switched to N_2 , the sample was then heated up at a rate of 15 °C min^{−1} to 130 °C and held at this temperature for 20 min. The same procedure was repeated for 50 cycles. The gas flow rate was set at 100 ml min^{−1} for all TGA tests.

3 Results and discussion

3.1 Surface textural and morphological characterisation of SCF materials

Fig. 1 presents the nitrogen adsorption–desorption isotherms and pore size distributions of the porous SCF samples, which were synthesized under different conditions to reveal the effect of the precursor and preparation chemistries on the textural and morphological characteristics of the siliceous foams and hence their corresponding performance as candidate supports

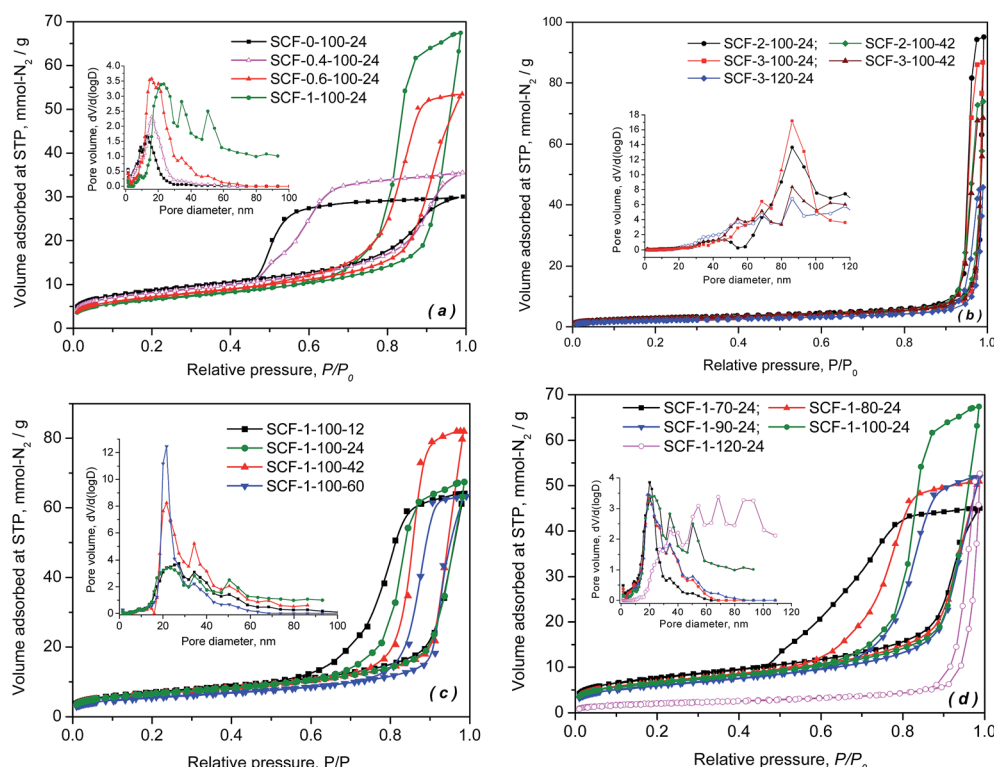


Fig. 1 N_2 adsorption isotherms and pore size distributions of the SCF samples prepared with different precursor chemistries under different conditions.



for preparing high capacity CO₂ adsorbents. As shown in Fig. 1, all synthesised cellular forms were found to have type IV isotherms with steep but variable types of hysteresis in the high relative pressure region, which is characteristic of typical meso-modulated materials that exhibit ink-bottle or thermodynamic effects associated with the condensation and evaporation of the adsorbate in the adjoining mesopores with smaller connecting channels or bottle necks. With increasing TMB/P123 ratio as shown in Fig. 1(a) and (b), the sorption hysteresis loops shifted dramatically toward higher relative pressures with an increasing amount of N₂ being adsorbed, which led to narrower pore size distributions (PSD) that also shifted toward larger pore diameters. This confirms the pore expansion behaviour of TMB. Meanwhile, with increasing amounts of TMB used, the pore hysteresis of the cellular materials also transformed from the wide H2 type hysteresis loops, which indicate the network nature of relatively uniform ink-bottle or channel-like pores, to the narrower H1 type hysteresis that is associated with the porous frameworks of higher pore size uniformity and facile pore connectivity. Clearly, the shift took place as a result of reduced curvatures or increased sizes of more uniform spherical composite micelles with increasing TMB concentration, due to the greater hydrophobic cores formed by the central hydrophobic polypropylene oxide chains of P123 dissolved into the non-polar TMB.

In general, the effect of aging time and temperature on pore hysteresis, as shown in Fig. 1(c) and (d), follows somehow the same trend as that of TMB. In general, the results demonstrate that an aging temperature of 100 °C and a TMB/P123 ratio of 1.0 give better control over the pore size distributions particularly in the mesoporous region, while the cellular materials synthesised at 120 °C or with higher TMB/P123 ratios (≥ 2), such as SCF-1-120-24, SCF-3-120-42 and SCF-2-100-42, appear to be almost entirely macro-structured with reduced volumes of nitrogen adsorbed and wider pore size distributions centred between 60 and 90 nm (Fig. 1(b) and (d)).

Table 2 summarises the surface textural properties of the cellular foam samples derived from the sorption isotherms, while Fig. 2 presents the variation of pore volume, pore diameter and window size with the synthesis parameters. The cell diameter or the sizes of cell cavities of the cellular foams were calculated using the adsorption branches of the N₂ isotherms, assuming that the pores are spherical, while the window size or the diameter of the channels connecting the pores was derived from the desorption branch of the isotherms (assuming cylindrical windows), using a modified BdB-FHH model proposed by Lukens *et al.*⁴³ It is evident that both the cell diameters and the window sizes increased significantly with increasing TMB concentration, aging time and aging temperature. Compared with the effect of aging time, however, the effect of TMB concentration and aging temperature appears to be more significant. At an aging temperature of 100 °C and aging time of 24 hours, increasing the TMB/P123 ratio from 1 to 3 dramatically expanded the cell cavity size from 21.2 to 74.2 nm and the window size from 11.2 nm to 39.1 nm, respectively, highlighting the novel pore expanding effects of TMB (Fig. 1(a)). Regarding the effect of aging temperature, although it appears to be quite modest at temperatures below 100 °C, a sharp increase in both the pore diameters and window sizes was observed when the aging temperature was increased slightly from 100 to 120 °C, as shown in Fig. 1(b). Closer examinations of the results shown in Table 2 and Fig. 2 also interestingly reveal that compared to the window sizes, the cell diameters increased more significantly with increasing TMB concentration, aging time and aging temperature. This suggests that the condensation of silica, which was responsible for the formation of the shells of TMB-cored micelles or micro-emulsion droplets, was significantly enhanced due to the accelerated hydrolysis of TEOS and aggregation of smaller micelles at higher aging temperatures, TMB concentrations and longer aging times, leading to the formation of larger cells with denser and thicker shells or walls.

Table 2 Surface textural properties of the synthesised SCF materials

Sample	BET surface area, m ² g ⁻¹	V _{total} , cm ³ g ⁻¹	V _{meso} , cm ³ g ⁻¹	V _{macro} , cm ³ g ⁻¹	Pore size, nm	Window diameter, nm
SCF-0-100-24	715	1.1	0.9	—	8.1	4.4
SCF-0.4-100-24	668	1.3	1.1	—	10.4	5.1
SCF-0.6-100-24	591	1.8	1.7	—	15.9	10.9
SCF-1-100-24	547	2.2	2.1	—	21.2	11.2
SCF-1-70-24	631	1.5	1.4	—	13.8	6.1
SCF-1-80-24	561	1.7	1.6	—	16.4	7.9
SCF-1-90-24	493	1.8	1.7	—	16.7	10.8
SCF-1-100-12	571	2.1	2.0	—	19.3	8.6
SCF-1-100-42	549	2.8	2.7	—	25.4	13.4
SCF-1-100-60	451	2.2	2.1	—	24.3	15.1
SCF-1-120-42	168	1.8	0.72	0.91	52.8	34.3
SCF-2-100-24	240	2.8	0.31	2.77	57.9	39.7
SCF-2-100-42	217	2.5	0.56	1.91	61.3	38.7
SCF-3-100-24	198	3.2	0.27	2.5	74.2	39.1
SCF-3-100-42	223	2.4	0.38	1.99	62.2	39.4
SCF-3-120-24	165	1.6	0.23	1.37	50.9	44.4
PQ meso-porous silica	269	1.7	1.7	—	24.7	18.6

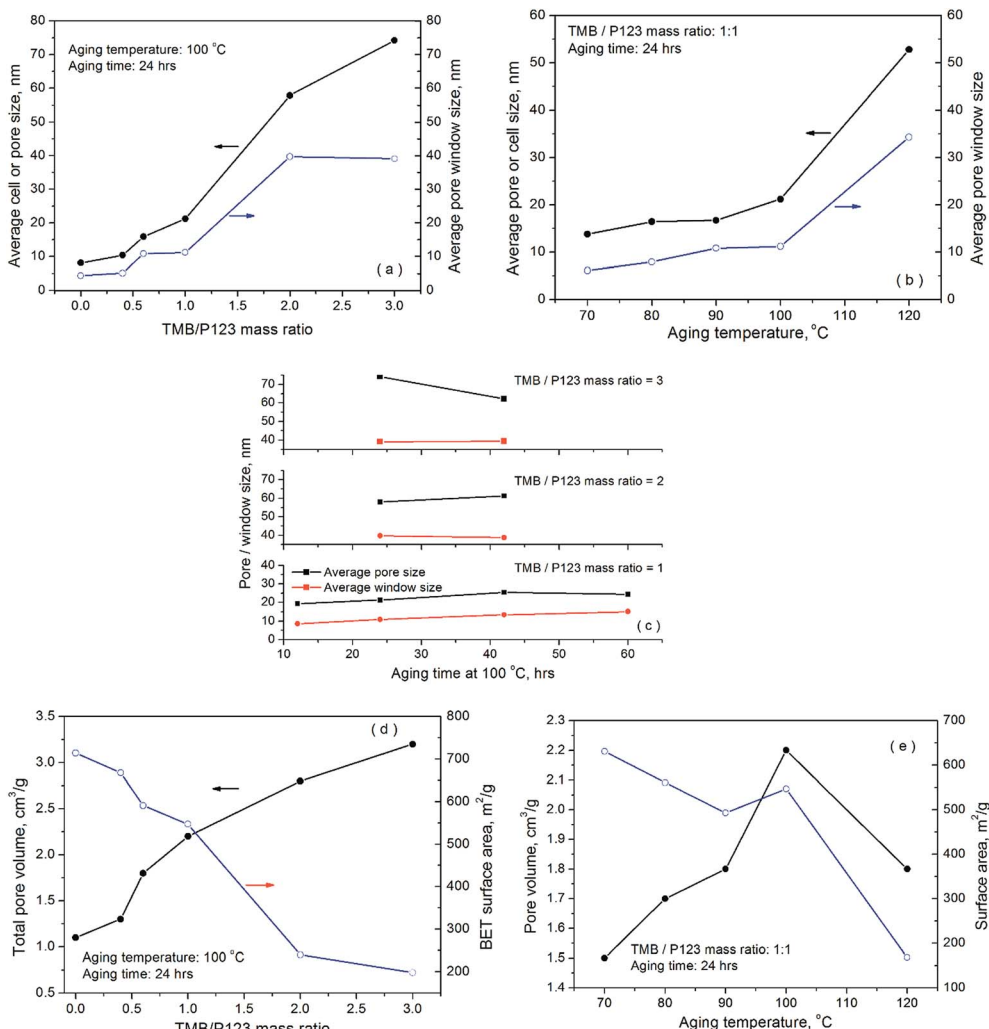


Fig. 2 Effect of precursor chemistry and preparation conditions on the surface textural properties of SCF material samples.

The surface area of the synthesised materials was found to depend heavily on the preparation conditions, varying from $714 \text{ m}^2 \text{ g}^{-1}$ for the sample SCF-0-100-24 where no TMB was used to just $165 \text{ m}^2 \text{ g}^{-1}$ for the sample SCF-3-120-24 prepared at 120°C and a TMB/P123 ratio of 3. As expected, the surface areas were found to decrease significantly as the pore volumes increased with TMB concentration and aging temperature. This is, in principle, because the resultant enlarged windows cost disproportionately a larger fraction of the surface areas of the enlarged cells. Indeed, all the samples synthesised at 120°C and those at 100°C but with high TMB/P123 ratios (≥ 2) were found to have the lowest surface areas, generally well below $250 \text{ m}^2 \text{ g}^{-1}$ but with largest pore diameters and window sizes (Table 2). In particular, the sample (SCF-3-120-24) prepared at an aging temperature of 120°C with a TMB/P123 ratio of 3 was found to have the largest window size of 44.4 nm but the lowest surface area of only $165 \text{ m}^2 \text{ g}^{-1}$ and the smallest pore volume of $1.6 \text{ cm}^3 \text{ g}^{-1}$. This further highlights the enhanced silica condensation as a result of accelerated hydrolysis of TEOS at high aging temperature and TMB concentrations, leading to increased cell diameters with reduced cell cavities and enlarged pore opening.

Schmidt-Winkel *et al.*⁴¹ found that the cell cavity or pore size has a linear relationship with the cubic root of the TMB/P123 mass ratio, which is in contrast to the direct linear relationship observed in a later investigation by Sridhar *et al.*⁴⁴ However, no such linear relationships were obtained across the whole wider range of TMB/P123 and aging temperatures examined in our investigation while our synthesised cellular foams were found to have significantly larger pore diameters and much lower surface areas compared to those obtained in previous investigations.^{41,44,45} Regarding the effect of precursor chemistry and preparation conditions on SCF wall thickness, van Grieken *et al.* and Schmidt-Winkel *et al.* both found that enhanced silica condensation is directly responsible for increased wall thickness,^{41,45} which contradicts the findings of Sridhar *et al.*⁴⁴ that increased wall thickness is not a result of enhanced silica condensation but shrunken pore size and a transition of pore geometry from being spherical to cylindrical. However, the findings revealed in Fig. 2, Table 2 and the imaging analyses below (Fig. 3) indicate that a combination of enhanced silica condensation and pore geometry transition coupled with the aggregation of smaller TMB-cored micelles due to reduced

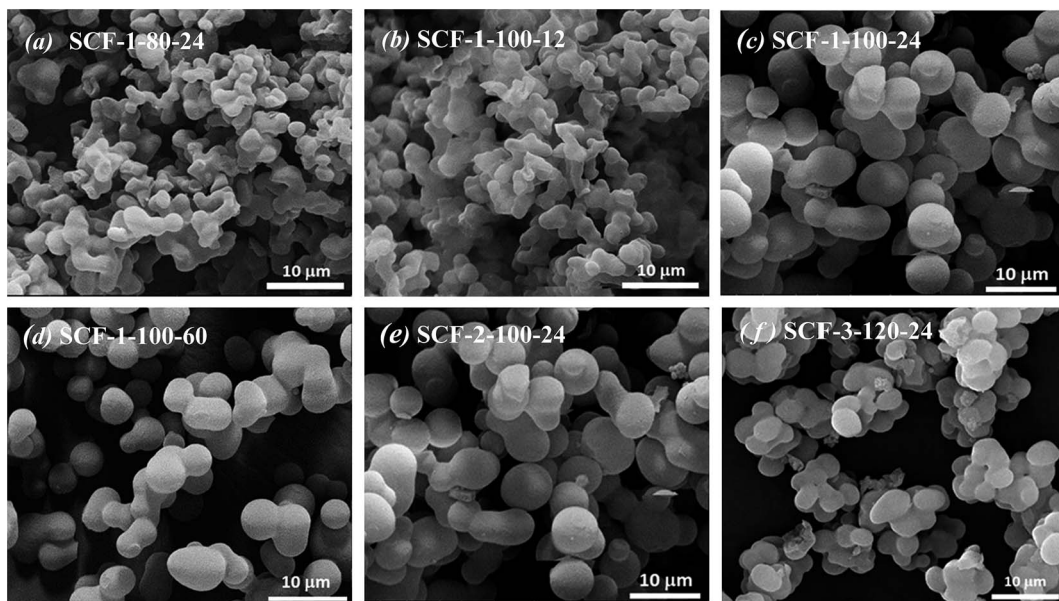


Fig. 3 FEG-SEM images of representative SCFs synthesized under different preparation conditions.

surface tensions is the most likely cause of the formation of larger cells with reduced cell cavities and enlarged window sizes at higher aging temperatures and TMB concentrations. The transition of pore hysteresis from H2 to H1 type as shown in Fig. 1 also corroborates the analyses above.

Fig. 3 and Fig. S1† show the morphological features at different microscopic scales of the typical cellular foams synthesised. As shown in Fig. 3(a) and (b), due to the extensive random aggregation observed among the smaller spheres of variable sizes, the siliceous particles prepared at lower aging temperatures and/or shorter aging times exhibit assembled coral-like cylindrical morphologies, which are somewhat similar to those of SBA-15.³⁸ In contrast, the formation of significantly larger, more uniform and less packed spherical particles is evident when increasing the aging temperature from 80 to 100 °C (Fig. 3(a) and (c)) or the aging time from 12 to 60 hours (Fig. 3(b) and (d)). A further transformation from spherical to ellipsoidal particles that tend to assemble into larger spherical aggregates can be seen with further increase in aging temperature from 100 to 120 °C and/or TMB/P123 ratio from 1 to 3. Closer examinations of the particle morphologies, as shown in Fig. S1,† reveal the open polygonal networks framed by silica struts, being characteristic of the structural features of siliceous cellular foams.^{41,46} Some deformations of the spherical cells are evident, probably caused by the unfavourable packing of the spherical micelles or the collapse of micellar cores during the packing due to their polydispersity. It can be seen that with increasing TMB concentration and aging temperature, the polygonal network structures became more developed and interconnected with increasingly more dense silica struts or thicker framework walls, while their sizes were found to be considerably less uniform with the formation of large empty voids or much greater cellular structures (Fig. S1(d)–(f)†). The FEG-TEM images shown in Fig. S2† tend to suggest that the

dense silica struts or walls of the SCF materials prepared at high aging temperatures with high TMB/P123 ratios were composed of highly interconnected and open-celled nano-foamed structures, highlighting the significantly enhanced silica condensation at nano-porous scales. By and large, the morphological transformation is consistent with the variation of surface areas and pore size distributions with TMB concentration and aging temperature as shown in Table 2 and Fig. 2(d) and (e).

3.2 CO₂ adsorption–desorption of the amine-modified sorbents

3.2.1 Quasi-static CO₂ adsorption study. To examine the variation of CO₂ uptake with temperature and hence to identify the optimal adsorption temperature, quasi-static CO₂ adsorption tests were first conducted for PEI-modified SCF adsorbents at temperatures between ambient temperature and 120 °C with a heating of 0.5 °C min^{−1}. The use of such a low heating rate is to facilitate near equilibrium CO₂ adsorption at individual adsorption temperatures in the range. Fig. 4 shows the results from the quasi-static tests for selected samples. As can be seen from Fig. 4, the CO₂ uptake first increased and then decreased with adsorption temperature for all the samples examined, and the differences between the CO₂ uptakes were significantly larger at lower adsorption temperatures. Meanwhile, the optimal adsorption temperature giving rise to the highest CO₂ uptake also varied considerably for different samples, despite their same level of PEI loading at 60 wt%. The sample SCF-3-120-24 was found to have the lowest optimal adsorption temperature of *ca.* 71 °C, compared to 75 °C for SCF-1-100-24 and SCF-2-100-24 and the highest optimal of 80 °C observed for the sample SCF-0-100-24, respectively.

Thermodynamically, low temperature favours the CO₂ adsorption of amine-modified sorbent materials, but increasing adsorption temperature can help overcome the kinetic barrier



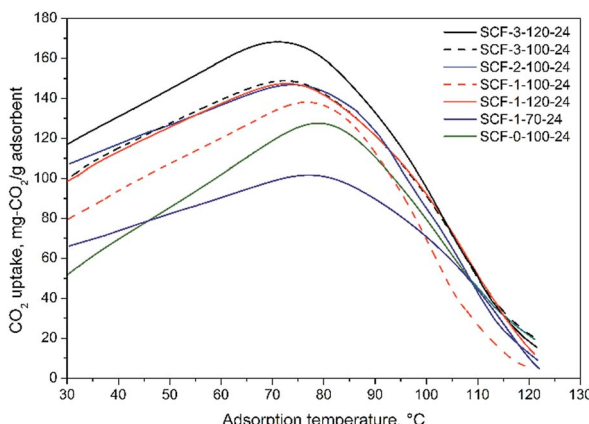


Fig. 4 Quasi-static CO_2 adsorption tests of selected PEI (60%)-modified SCF adsorbents in simulated flue gas (15% CO_2 in N_2). Conditions: heating rate of $0.5\text{ }^\circ\text{C min}^{-1}$ from ambient temperature to $120\text{ }^\circ\text{C}$; content of PEI-600: 60 wt%.

of CO_2 adsorption due to the accordingly improved mobility and hence diffusivity and accessibility of the impregnated PEI molecules.^{27,42,47,48} The considerable differences observed in the optimal adsorption temperatures and CO_2 uptakes reveal the different mass transfer limitations or diffusivities of CO_2 in the amine sorbents. It is evident that irrespective of the pore volumes, the SCF materials synthesised at higher TMB concentrations or aging temperatures, which usually had larger pore diameters and wider window sizes, also have lower optimal adsorption temperatures in general. In addition, the results also show that although the differences between the variations of CO_2 uptake with temperature diminished significantly at higher-than-optimal temperatures, the CO_2 uptake of the samples with higher optimal adsorption temperatures generally decreased more rapidly with temperature, being indicative of the lower diffusional resistance of the desorbed CO_2 when diffusing out through the pores and the bulk phase of immobilised amines during the desorption at higher temperatures. Presumably, for a given level of PEI loading, this is because larger pore and window sizes in an interconnected porous network can facilitate greater thermo-mobility and hence the kinetic behaviour of the immobilised amines through the pores such as to enhance the exposure of unreacted amine groups to CO_2 or facilitate the CO_2 diffusion deeper into the immobilised polymer amines, leading to higher amine efficiency or CO_2 uptake of the sorbents. By and large, the results from the quasi-equilibrium adsorption tests are generally in good agreement with the findings of Sakwa-Novak *et al.*⁴⁹ that improved amine mobility by adding additives can significantly increase the temperature sensitivity of immobilised amines for CO_2 capture and hence the amine efficiency due to the resultant higher probability of intra-chain CO_2 adsorption events. Builes and Vega²⁷ also found that the high mobility of immobilised amines can facilitate the creation of a diffusive network of micro-cavities within the bulk phase of the amines, which favours the diffusion of CO_2 in and out of the amines within the pores. Taking into account the surface textural properties (Table 2 and Fig. 2), the above results indicate that the formation of the

diffusive network of micro-cavities can be better facilitated in the cellular materials with larger window sizes and somewhat greater pore diameters.

3.2.2 CO_2 adsorption capacity of SCF-supported amine adsorbents. Fig. 5 shows the adsorption profiles of representative PEI-modified SCF adsorbents for each series of SCF materials synthesised. Despite the considerable variation of optimal adsorption temperatures for different SCFs, the CO_2 adsorption capacities were all measured at the same temperature of $75\text{ }^\circ\text{C}$ for the sake of consistent comparison. As can be seen from Fig. 5, all SCF samples showed significantly faster adsorption kinetics and higher CO_2 capacities than the commercial benchmark silica (PQ Silicas), which is a highly meso-structured siliceous material with its pore volume, pore diameter and window size being similar to or even greater than those of the SCFs (Table 2). The times taken for the SCF adsorbents to achieve 80% and 90% of equilibrium CO_2 capacity were found to be just *ca.* 1.5 min and 3 min, which is nearly 8 times faster than the *ca.* 12 and 23 min obtained for the PQ silica, respectively. This highlights the novel accessibility of the immobilised amines to CO_2 due to the high levels of pore network connectivity in the SCF materials.

Table 3 compares the CO_2 adsorption performance for all PEI-600-modified SCF materials at different PEI loading levels, while Fig. 6 shows the variation of CO_2 uptake with the precursor chemistry and preparation conditions used. All PEI-modified SCF adsorbents showed increased CO_2 uptake capacities with increasing levels of PEI but with greater variations in CO_2 capacity, determined by the preparation conditions. Among all the materials, SCF-3-120-24 gave rise to the highest CO_2 adsorption capacity of $180.5\text{ mg per g-adsorbent}$ at a PEI content of 70 wt%, followed by SCF-2-100-42 (178.4 mg CO_2 per g) and SCF-3-100-42 (176.5 mg CO_2 per g), which are all significantly higher than those obtained previously for amine-modified mesoporous cellular foam materials³⁸⁻⁴⁰ and virtually any other 2D mesoporous silica materials.^{28,35,50-52} At 50 wt% and 60 wt% PEI, however, the highest CO_2 uptake was instead obtained for SCF-1-100-60 at 150.4 and 168 mg g^{-1} , respectively. It is noteworthy that with a total pore volume of just $1.1\text{ cm}^3\text{ g}^{-1}$,

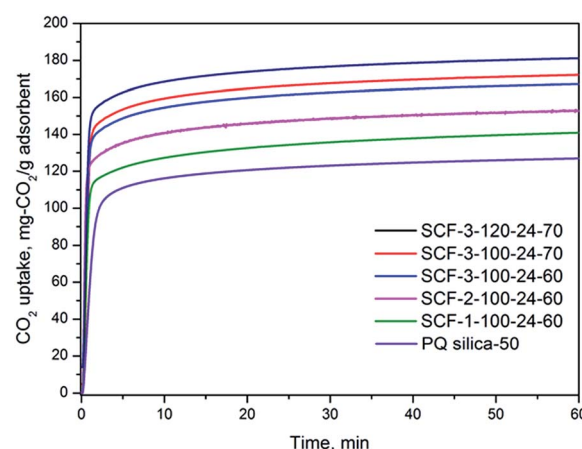


Fig. 5 CO_2 adsorption profiles of representative PEI-impregnated SCF adsorbents in $15\% \text{CO}_2/\text{N}_2$ at $75\text{ }^\circ\text{C}$.



Table 3 CO_2 uptake capacities of PEI-impregnated SCF adsorbents with different PEI-600 contents at 75 °C in simulated flue gas (15% CO_2 in N_2)

Samples	CO ₂ uptake, mg-CO ₂ per g-adsorbent			Max. CO ₂ uptake	
	50 wt%	60 wt%	70 wt%	mg CO ₂ per g-ads	mg CO ₂ per g-PEI
SCF-0-100-24	118.2	139.4	107.1	139.4	232
SCF-0.4-100-24	125.2	143.2	140.2	143.2	239
SCF-0.6-100-24	134.9	157.9	143.7	157.9	263
SCF-1-100-24	128.7	148.9	162.7	162.7	271
SCF-1-70-24	85.3	103.5	67.8	103.5	173
SCF-1-80-24	91.6	128.4	119.5	128.4	214
SCF-1-90-24	100.6	138.4	154.7	154.7	221
SCF-1-100-12	113.3	139.2	151.0	151.0	215
SCF-1-100-42	137.3	151.3	173.9	173.9	248
SCF-1-100-60	150.4	172.1	144.4	172.4	287
SCF-1-120-24	134.4	156.7	166.2	166.2	237
SCF-2-100-24	113.1	156.2	167.3	167.3	239
SCF-2-100-42	101.3	146.2	178.4	178.4	255
SCF-3-100-24	115.9	146.4	172.5	172.5	246
SCF-3-100-42	112.4	149.8	176.5	176.5	252
SCF-3-120-24	116.7	156.4	180.8	180.8	258
PQ mesoporous silica	127.4	113.8	104.7	131.4	255

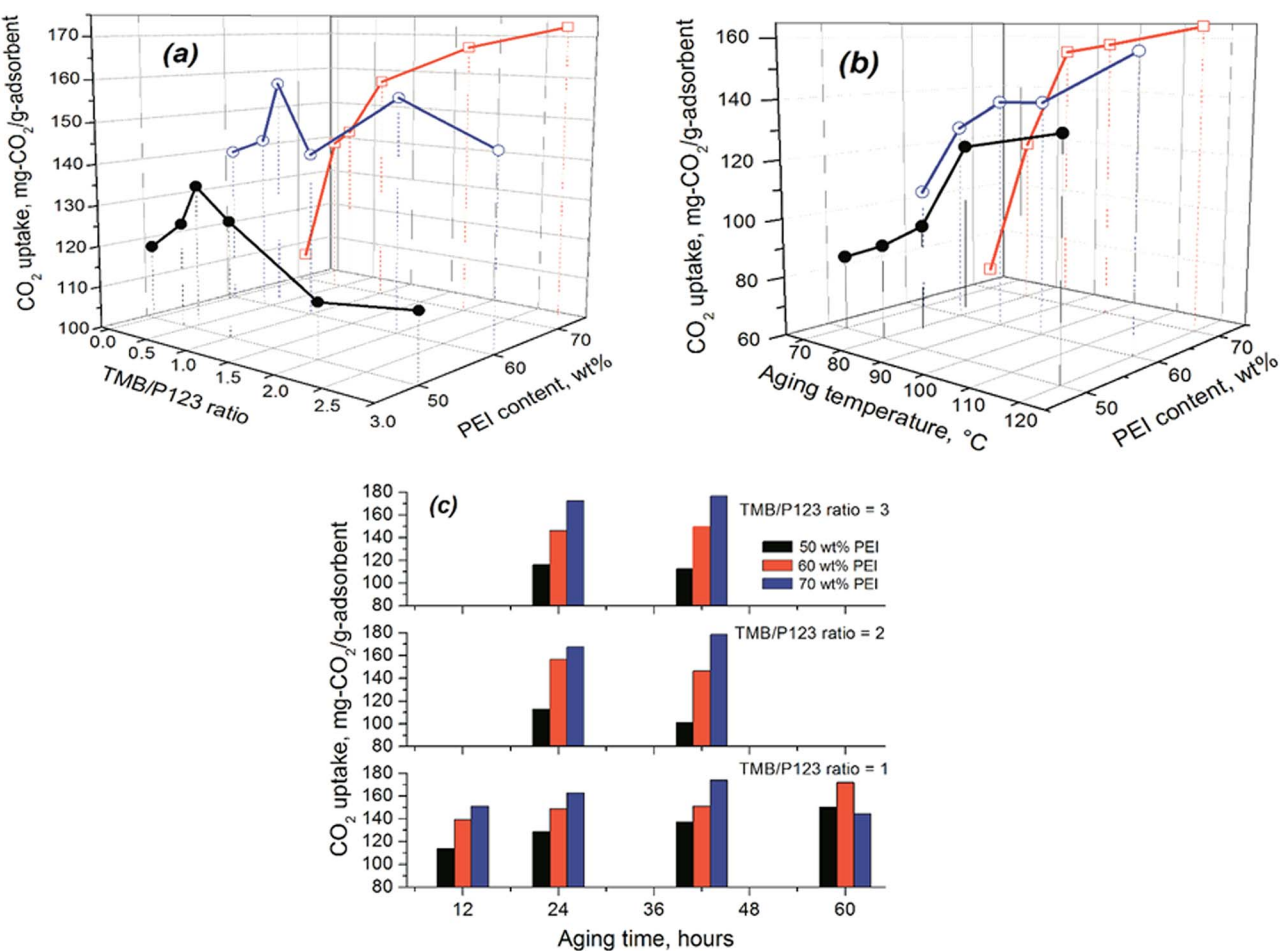


Fig. 6 Variation of the CO_2 uptake of SCF materials with precursor chemistries and other preparation conditions.

the SCF material prepared without the pore-expanding agent (SCF-0-100-24) also achieved a higher CO_2 capacity of 138 mg- CO_2 per g-ads than that of the baseline PQ silica material (127.4 mg CO_2 per g-ads) with an over 50% higher meso-pore volume ($1.7 \text{ cm}^3 \text{ g}^{-1}$), highlighting the importance of the 3D pore connectivity of the SCF materials. It appears that for the SCF materials prepared with lower TMB concentrations, there generally exists a maximum PEI loading level beyond which the CO_2 adsorption capacity decreases with further increase in PEI loading, being consistent with previous findings on amine-modified MCFs.^{38,53} However, the same trend was not observed for those prepared with higher TMB concentrations or aging temperatures, which showed much lower CO_2 uptake capacities but with a sharp continuous increase with increasing PEI loading up to 70 wt%. Nevertheless, no further appreciable increases in CO_2 uptake were obtained when levels of PEI impregnation were increased to higher than 70 wt% and the prepared sorbents started to become pasty.

Fig. 6 indicates that increasing TMB concentration and aging temperature/time led to increased CO_2 adsorption capacity for all SCF materials, but the trend varied significantly for different individual conditions and PEI loading levels. As can be seen from Fig. 6(a), the effect of TMB concentration appears to depend on the amount of PEI immobilised in the SCF materials. At relatively lower PEI loading levels (≤ 60 wt%), the CO_2 uptake first sharply increased with increasing the TMB/P123 ratio from 0 to 1.0 and then decreased significantly when the ratio was further increased from 1 to 3. This instead led to lower CO_2 capacities or amine efficiencies for the best-performing SCF materials that were synthesised all with high TMB/P123 ratios (≥ 2), compared to those prepared with lower TMB concentrations, as shown in Fig. 6(a). However, at a higher PEI loading level of 70 wt%, no such trend was observed as can be seen from the continuous increase of the CO_2 uptake with TMB concentration. This is clearly because of the variable maximum PEI-accommodating abilities, which were lower for the SCF materials synthesised with lower TMB concentrations and aging temperatures (Table 3). A similar trend was also observed on the effect of aging time, namely an increase in aging time led to reduced CO_2 capacity or amine efficiency at lower PEI contents for the SCF substrates synthesised using high TMB/P123 ratios. In comparison, all SCF materials showed a continuous increase in CO_2 uptake with aging temperatures at all PEI loading levels, as indicated in Fig. 6(b). It is noteworthy that despite the variable trends observed, the SCF materials prepared with higher TMB/P123 ratios (≥ 2) and aging temperatures ($> 100^\circ\text{C}$) all exhibited greater CO_2 adsorption capacities.

In theory, given the similar nature of the SCF materials all with interconnected 3D porous structures, the CO_2 capacity of the range of SCF-based adsorbents should be proportional to the amount of PEI impregnated onto the SCF substrates, which is in turn determined by the total pore volume of the substrates. Previous investigations have found that the total pore volume was the decisive factor affecting the CO_2 uptake at high amine loading levels particularly for meso-structured support materials with 2D hexagonal symmetries.^{38,42,54,55} Fig. 7 shows the

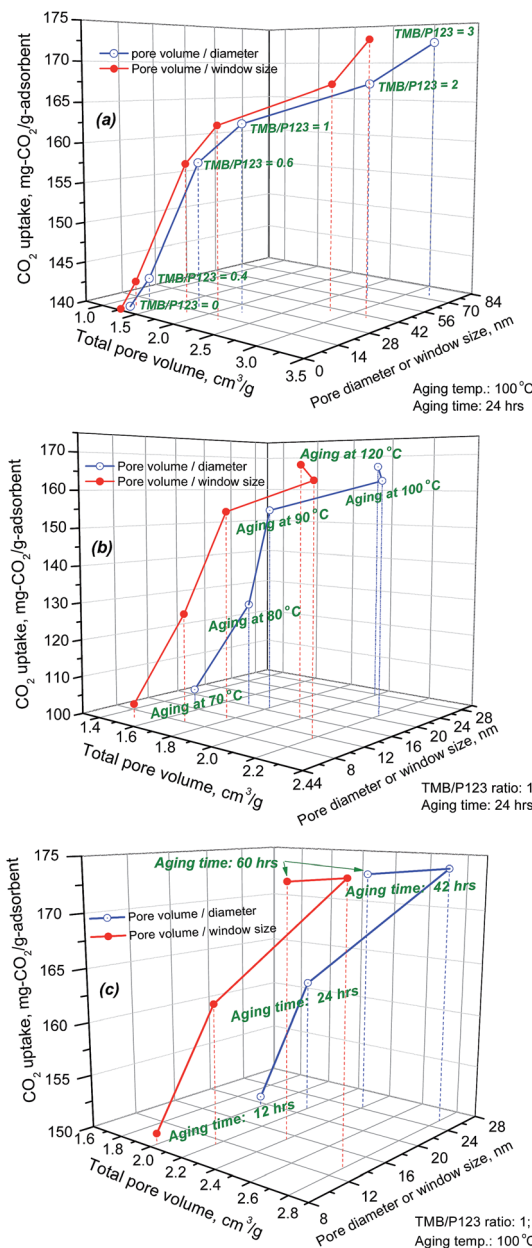


Fig. 7 Effect of total pore volume, pore diameter and pore window size on the maximum CO_2 uptake capacity of the SCF materials synthesised under different conditions.

variation of CO_2 uptake capacity as a function of the surface textural properties of the SCF materials. It is evident that the CO_2 uptake capacity increased dramatically with the total pore volume, pore diameter and window size of the SCF substrates, particularly for the SCF materials synthesised at lower TMB concentrations and/or aging temperatures. In addition, close-to-linear relationships can also be observed between the CO_2 uptake and the total pore volume for the SCF substrates with pore volumes lower than $2.2 \text{ cm}^3 \text{ g}^{-1}$ and pore diameters/window sizes below 28 nm as shown in Fig. 7(a) and (c). For the SCFs with larger pore volumes or pore diameter/window sizes, the CO_2 uptake deviated from the linear relationship toward lower CO_2 uptakes, indicating the lower amine efficiencies or

that the pores could not possibly be fully impregnated with PEI to give higher CO_2 capacity.

However, it needs to be emphasised that all the best-performing SCF materials, which were the macro-structured siliceous foam materials synthesised with high TMB/P123 ratios (≥ 2) and aging temperatures (120°C) or long aging times (> 24 hours), all interestingly exhibited the lowest CO_2 capacities or amine efficiencies at low PEI impregnation levels but their CO_2 uptake capacities were found to increase much more sharply with PEI impregnation, instead giving rise to the highest CO_2 capacities at high PEI loading levels among all the materials as shown in Table 3 and Fig. 6(a and b). These anomalies were found to be largely responsible for the significant data scattering observed in the above correlations (Fig. 7) where SCF-2-100-42, SCF-3-100-42 and SCF-3-120-42 were excluded. For instance, SCF-3-120-24, which had the lowest total pore volume of only $1.6 \text{ cm}^3 \text{ g}^{-1}$ and surface area of $165 \text{ m}^2 \text{ g}^{-1}$, had one of the lowest CO_2 uptakes at $117 \text{ mg-CO}_2 \text{ per g-ads}$ (233 mg per g-PEI) at 50 wt% PEI but it achieved the highest CO_2 uptake of $180.8 \text{ mg-CO}_2 \text{ per g-adsorbent}$ ($258 \text{ mg-CO}_2 \text{ per g-PEI}$) with 70 wt% PEI loading among all the SCF materials, and this compares to that of the SCF-1-100-24 sample, which has a higher capacity of $128.7 \text{ mg-CO}_2 \text{ per g-ads}$ ($257.4 \text{ mg-CO}_2 \text{ per g-PEI}$) at 50 wt% PEI but has a much lower capacity of just $162.7 \text{ mg-CO}_2 \text{ per g-ads}$ ($232 \text{ mg-CO}_2 \text{ per g-PEI}$) at 70 wt% PEI. Meanwhile, SCF-3-100-24, which had the largest pore volume of $3.2 \text{ cm}^3 \text{ g}^{-1}$, gave rise to a CO_2 uptake of just $172.5 \text{ mg CO}_2 \text{ per g-ads}$, which was considerably lower than those of SCF-3-120-24 and other materials with much lower pore volumes. Similar comparisons can also be drawn between SCF-3-100-42 and SCF-1-100-42, SCF-2-100-42 and SCF-1-100-60 and SCF-3-120-24 and SCF-1-100-24.

The findings that macro-structured siliceous foams outperform their meso-structured counterparts in CO_2 uptake have not been observed in previous investigations, which mostly focused on meso-structured MCFs^{38–40} and other silica materials.^{28,35,50–52,55} In addition, Yan *et al.*³⁸ found that the pore window size, when larger than 8.6 nm , is no longer an important factor affecting the impregnation and distribution of PEIs and hence the CO_2 capacity of PEI-modified MCFs. This clearly cannot be generalised to the SCFs in this investigation probably because of the small number of samples or conditions examined previously.^{29,38–40,52,53}

Clearly, the exceedingly high CO_2 uptake performance of the highly macroporous SCF materials cannot be accounted for solely by their greater porosities and pore geometries. Closer examinations of the surface textural results shown in Table 2, Fig. 2, 3 and S1[†] reveal that the SCF materials prepared at higher aging temperatures, longer aging times and higher TMB concentrations all expectedly had increased pore diameters and/or window sizes but with drastically reduced surface areas (Table 2). In addition, it seems that the effect of aging temperature and time on the surface textural properties of the SCF materials depends on the TMB concentrations used in the synthesis. Further, as shown in Table 2 and Fig. 2, the effect of aging temperature and time on the surface textural properties at high TMB concentrations was also found to differ from the

effect at low TMB concentrations in that at high TMB concentrations, further increase in aging time from 100 to 120°C or aging times from 24 to 42 hours led to significantly reduced, rather than increased, pore volumes and macro-pore sizes (*e.g.* SCF-3-120-24 *vs.* SCF-3-100-24 and SCF-3-100-42 *vs.* SCF-3-100-24) with boosted silica struts or cell wall thickness as shown in Fig. S1(e) and (f).[†] This essentially led to the formation of mixed meso/macro-cellular SCFs with relatively wider but regular pore size distributions as shown in Fig. 1(b) and (d). It is therefore believed that the formation of the ordered meso/macro-porous SCF materials with enhanced silica strut structures, which serve as the framework platforms for impregnation, can facilitate higher PEI impregnation levels and more importantly, greater mobility or diffusivity of the immobilised bulk amines, which is vital in boosting the accessibility to CO_2 molecules. This hypothesis is strongly supported not only by the high amine efficiencies achieved by the SCFs at high PEI impregnation levels but also their lower optimal adsorption temperatures, which act as an indicator of the amine mobility, and the faster adsorption kinetics at high PEI loading levels, which indicates the low CO_2 diffusion resistance within the bulk phase of the supported PEI amines, as shown in Fig. 4. However, the significantly lower surface areas of the highly meso/macro-cellular SCFs, which decreased from typically over $500 \text{ m}^2 \text{ per g-ads}$ to below $250 \text{ m}^2 \text{ per g-ads}$, may lead to lower CO_2 uptake capacities at low PEI loading levels due to the disproportionately reduced larger number of available top surface amine sites due to the accordingly increased coating thickness, which cannot be offset by the increased amine mobility of the bulk impregnated PEI at low loading levels.

3.2.3 CO_2 adsorption capacity of SCF materials modified with varying molecule weight PEIs. Binary impregnation was examined as a means to further improve the CO_2 adsorption performance of the amine-modified SCF adsorbents. The results above suggest that the accessibility of the bulk phase of impregnated PEIs plays a vital role in determining the CO_2 uptake capacity of the PEI-SCF adsorbents, which is affected heavily by both the surface textural and morphological properties of the SCF materials that govern the distribution state of the impregnated PEIs. Previous investigations have confirmed that the use of surfactants in amine impregnation can facilitate additional CO_2 pathways within the bulk phase of immobilised amines and hence improve the mobility and accessibility of the amines for higher amine efficiencies and CO_2 capture capacities, and CO_2 -unreactive surfactants were typically used, such as sodium dodecylsulfate,⁵⁶ P123, STAB and SPAN80⁵⁷ and CTAB and polyethylene glycol of different molecular weights.⁴⁹ Instead of using CO_2 -unreactive surfactants, tetraethylenepentamine (TEPA), which is a type of low molecular weight PEI, was used as the substitute for surfactants to prepare the binary PEI adsorbents *via* co-impregnation, and two best performing SCF support materials, namely SCF-3-120-24 and SCF-3-100-24 were selected for the preparations. Fig. 8 presents the CO_2 uptake performance of the binary SCF adsorbents.

As can be seen from Fig. 8, the binary amine-modified SCF adsorbents achieved significantly higher CO_2 uptake for both



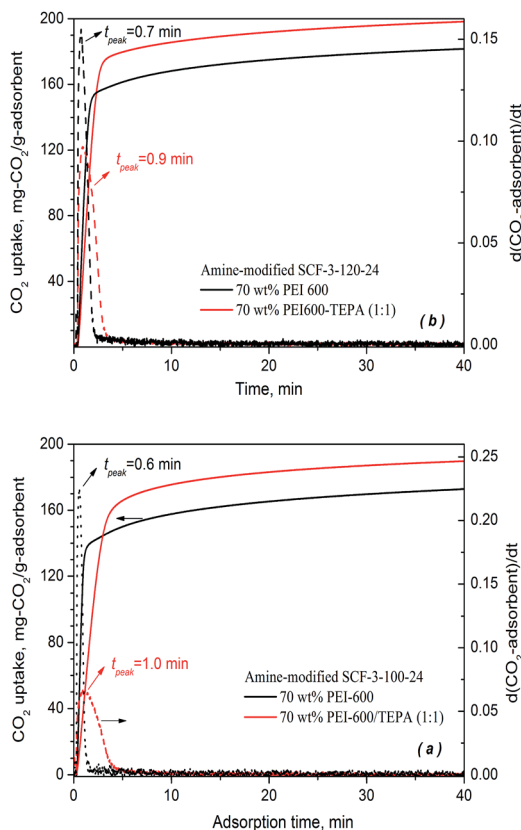


Fig. 8 CO₂ adsorption profiles of binary amine-modified SCF adsorbents at 75 °C in a simulated flue gas of 15% CO₂ in N₂.

SCF materials examined at an amine loading level of 70 wt%. The TEPA-PEI modified SCF-3-120-24 reached an exceedingly high CO₂ capacity of 198.2 mg-CO₂ per g-ads, 10% higher than its PEI-modified counterpart (180.8 mg-CO₂ per g-ads). A similar increase in CO₂ uptake by approximately 10% was also obtained for SCF-3-100-24, from 172.5 mg-CO₂ per g-ads for PEI impregnation to 190 mg-CO₂ per g-ads for PEI-TEPA co-impregnation. The CO₂ capacities obtained for the binary SCF adsorbents represent the highest ever reported under similar conditions.^{58–60}

Nevertheless, it appears that at the early stage of adsorption, the binary co-impregnation resulted in somewhat slower adsorption kinetics, as opposed to the faster kinetics expected for the lower viscosity and more reactive amine groups of TEPA compared to the higher molecular weight PEI-600. For SCF-3-100-24, the time taken to achieve 80% of equilibrium capacity increased from 1.4 min for PEI impregnation to 3.2 min for the PEI/TEPA co-impregnation whereas for the SCF-3-120-24 material, the time increased from 1.6 to 2.4 min, respectively. Meanwhile, the peak adsorption time (t_{\max}) increased from 0.6 to 1.0 min for SCF-3-100-24 and slightly from 0.7 to 0.9 min for SCF-3-120-24, respectively, when the impregnation changed from PEI-600 to the mixture of TEPA and PEI-600. The results differ from the previous findings that the use of blended amines, such as TEPA and DEA, for impregnation resulted in faster rates of adsorption kinetics in addition to higher capacities. Presumably, the lower rate of adsorption kinetics of the

binary SCF adsorbents is because of the potentially more dense bulk phase packing of the binary amines as the much less viscous and smaller TEPA molecules can penetrate into the micro-cavities created by the larger PEI-600 molecules, leading to steric crowding within the bulk phase of the impregnated mixed amines and hence increased diffusion resistance for CO₂ migration.²⁷ This kind of effect may not be significant in the case of the co-impregnation of TEPA and DEA, because of their similar molecular sizes and when the adsorption takes place at low temperatures.^{58,59}

In addition to the co-impregnation of amines of different molecular weights, attempts were also made to examine the suitability of the synthesised SCF materials for impregnating higher molecular mass PEIs as a measure to improve the thermal stability of the adsorbents. The development of amine-impregnated adsorbents for CO₂ capture has so far been mostly limited to the use of low molecular weight amines, such as TEPA, DEA and PEIs with a molecular mass typically below 1800, because they are less viscous and can facilitate faster adsorption kinetics and higher adsorption capacity at the same impregnation levels. However, the supported lower molecular mass amines usually have lower thermal stability, giving rise to inevitable amine loss due to evaporation and thermal decomposition loss in long term operations.⁶¹ To this end, higher molecular mass PEIs do offer some considerable advantages. However, in terms of adsorption capacity and kinetics, the CO₂ adsorption performance of higher molecular mass PEIs can be sluggish due to their higher viscosities and tertiary amine contents,⁶² and thus more robust porous support materials are required to mitigate the negative impact. To further investigate the ability of the synthesised SCF materials to accommodate higher molecular weight PEIs as a means to improve the thermal stability of supported PEI sorbents, the meso-structured SCF-1-100-24 was selected as the support for the impregnation of higher molecular mass PEIs. Fig. 9 shows the CO₂ adsorption performance in comparison with the baseline 2D meso-structured PQ silica material.

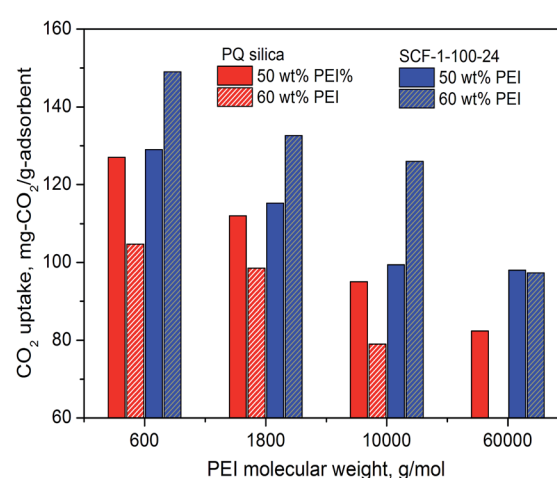


Fig. 9 Comparison of the CO₂ adsorption performance of the SCF materials modified with different molecular mass PEIs measured at 75 °C in simulated flue gas (15% CO₂ in N₂).



As expected, the results demonstrate that the CO_2 capacity decreased considerably for both the SCF and the baseline 2D meso-porous PQ silica with the increase of PEI molecular weight. At 50% PEI content, the CO_2 adsorption capacity of the PQ silica supported PEI dropped from 127.4 to 95 mg $\text{mg-}\text{CO}_2$ per g-ads as the molecular mass of PEI increased from 600 to 60 000, whereas the capacity of SCF-1-100-24 with 60 wt% PEI loading decreased from 148.9 mg- CO_2 per g-ads to 127 mg- CO_2 per g-ads, respectively. It is important to highlight that the meso-structured siliceous cellular foams facilitated significantly higher CO_2 capacities than the benchmark PQ silica for all the higher molecular mass PEIs at all impregnation levels examined, with their CO_2 capacity increasing significantly with increasing levels of impregnation. At 60 wt% PEI content, the PEI-modified SCF adsorbent gave rise to impressive capacities of 126 and 97.3 mg- CO_2 per g-ads for PEI-10 000 and PEI-60 000, respectively, which are even significantly higher than those reported for PEI-1800 modified PQ silica⁶³ and PEI-2500 modified MCF.⁴⁰ In contrast, the CO_2 adsorption capacity of the PEI-impregnated mesoporous PQ silica decreased sharply with increasing levels of the PEIs and no appreciable capacities could be measured at an impregnation level of 60 wt% for PEI-60 000. This highlights that the SCF materials are particularly suitable for preparing high molecular mass PEI adsorbents *via* impregnation with significantly improved thermo-stability only at a modest cost of CO_2 capacities.

3.2.4 Cyclic adsorption–desorption performance testing of PEI-modified SCF materials. The regenerability or cyclic adsorption–desorption performance of a CO_2 adsorbent is critical in determining their life-time importance in practical applications. Therefore, some of the best-performing PEI-600-modified SCF adsorbents are selected for cyclic performance testing under simulated flue gas conditions. As shown in Fig. 10, no appreciable changes in CO_2 uptake were observed from the 50 cycles of adsorption–desorption tests for different selected samples, highlighting the novel thermo-stability and

fully reversible CO_2 adsorption of the adsorbents. The negligible losses observed in the first few cycles were believed to be mainly associated with the evaporation loss of small quantities of lower molecular mass PEI components present in the PEI. In addition, it seems that the SCF materials prepared at higher aging temperatures or longer aging times present considerably higher thermal stability, which is attributable to the presumably increased holding strength of the materials for the impregnated PEI due to their denser and thicker shells as shown in Fig. 3 and S1.[†]

4. Conclusions

A wide range of meso, hybrid meso/macro and macro-structured siliceous cellular foam (SCF) materials have been tailor-designed and fabricated for preparing PEI-modified highly efficient adsorbent materials for CO_2 capture. The results demonstrate that while all the SCF materials exhibited higher capacities and faster adsorption kinetics in comparison with conventional meso-structured materials with 2D hexagonal pore geometries, all best-performing SCF materials are the hybrid meso/macro and macro-structured materials prepared using high TMB concentrations, high aging temperatures and longer aging times despite their drastically reduced surface areas. The CO_2 uptake capacity of the SCF materials reached up to 180.2 mg- CO_2 per g-adsorbent (5.85 mmol per g-amine) for PEI impregnation and was further improved to 198.2 mg- CO_2 per g-adsorbent (6.44 mmol per g-amine) for the binary impregnation of PEI–TEPA at 75 °C in a simulated flue gas of 15% CO_2 in N_2 , which are significantly higher than the previously reported under similar or comparable conditions. The macro- and hybrid meso/macro-structured SCF materials were also found to be particularly suitable for preparing high molecular weight PEI-modified adsorbents with significantly improved thermo-stability. At 60 wt% PEI loading, the CO_2 capacity reached 126 and 97.3 mg- CO_2 per g-ads for PEI-10 000 and PEI-60 000, respectively, which are the highest ever reported. The PEI-modified adsorbents displayed superior thermo-stability with fully reversible CO_2 adsorption as no appreciable changes were observed during the 50 cycle lifetime performance testing.

Relating the CO_2 adsorption performance to the surface textural and morphological properties of the SCF materials reveals that the pore volume, pore diameter and window size of the SCF substrates collectively determine the CO_2 uptake capacity of the PEI-modified adsorbents while for the macro- and hybrid meso/macro SCFs, the morphology particularly the thickness of cell walls or framework silica struts also plays a vital role in facilitating higher levels of PEI impregnation with enhanced PEI mobility and accessibility for higher CO_2 uptake capacities. Differing from previous findings, no particularly favourable pore diameters or window sizes for PEI impregnation are observed for the wide range of SCF materials examined, although close to linear relationships appear to exist between CO_2 uptake capacity and total pore volume for the SCF materials with pore volumes lower than $2.2 \text{ cm}^3 \text{ g}^{-1}$ and pore diameters/window sizes below 28 nm.

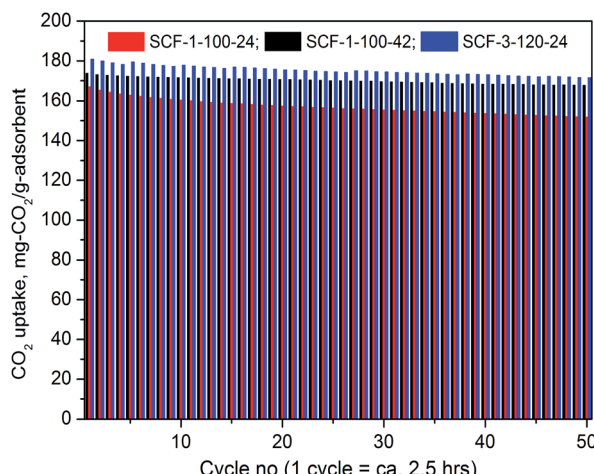


Fig. 10 Cyclic adsorption–desorption profiles of PEI-600-modified SCF materials for CO_2 uptake under simulated flue gas conditions (15% CO_2 in N_2). Conditions: adsorption temperature: 75 °C; desorption temperature: 130 °C; heating rate: 15 °C min^{−1}.



Conflicts of interest

Acknowledgements

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