1. Introduction

The concentration of CO₂ in the atmosphere has increased from 340 ppm in 1980 to 402.8 ppm in 2016.1 2 According to generally accepted climate change scenarios, very serious environmental consequences can be expected if, as forecast, the atmospheric level of CO₂ continues to rise above 450 ppm.3 4 An overview of the global carbon budget indicates that anthropogenic CO₂ emission from fossil fuels, currently at ca. 9.9 Pg C/yr,1 5 continues to be the main source of emissions. The total level of emissions, including land-use emissions (1.3 ± 0.7 PgC /yr), far exceed the capacity of the planet’s natural CO₂ sinks in the ocean (2.4 ± 0.5 PgC /yr) and on land (3.0 ± 0.8 PgC /yr).1 According to international efforts to address climate change consolidated by the Paris agreement (2016), carbon capture and storage/utilization (CCS) is one of the important mitigation strategies for limiting and/or reducing the levels of atmospheric CO₂ in the medium-term.

The current technologies for CCS, mainly amine scrubbers, can be efficiently implemented at large point sources of CO₂ emission. Pre-combustion capture is typically associated with coal-fired integrated gasification combined cycle plants, i.e. conversion of coal to H₂, CO and CO₂, where the concentration of CO₂ is ~30%. For existing pulverized-coal, oil or gas fired plants, post-combustion CO₂ capture technologies are required, where CO₂ is to be removed from the typically dilute (<15% by volume) flue gas. Among the current trends in CCS there is a shift from absorption processes of CO₂ capture by liquid (aqueous) amines, to CO₂ adsorption by solid porous materials. Separation by solid adsorbents may provide an energy benefit of up to 50% on the adsorbent regeneration with respect to the energy penalty in the regeneration step in the liquid amine based process.6 7 8 It is considered that for solid sorbents to be competitive with the existing MEA scrubbing system, the CO₂ working capacity must be in the range of 3-4 mmol of CO₂ per gram of sorbent.8 9

The solid porous materials for CCS that have been tested within the last decade, can be classed into two large groups: (i) physiosorbents (zeolites, porous carbons, metal-organic frameworks) and (ii) porous supports grafted/impregnated with various bases (mainly various polyethyleneimines). Impregnation of porous carbon supports with polyamines completely shifts the CO₂ adsorption mechanism to chemisorption.10 Due to the increased heat of CO₂ adsorption on amine impregnated or functionalised porous materials, these materials demonstrate significantly higher selectivity11 12 13 14 with respect to CO₂ when compared to physisorbents. This, however, also implies that the heat of desorption needed to release the CO₂ and regenerate the sorbents is also high. Porous carbon supports have advantages over the more extensively investigated silica supports due to the possibility of incorporating an electrical swing process, making use of the electrical properties of the materials. The adsorption capacity is typically proportional to the amount of loaded amine, although the adsorption efficiency with respect to the amount of present amino groups is highly dependent on surface morphology, pore size and available pore volume, \(V_{surf}\), of the porous substrate.10 In earlier work, we observed a significant increase of CO₂ uptake, reaching a value of 2.3 mmol g⁻¹ at a CO₂ partial pressure of 0.1 bar,15 when the pore size and, in particular, pore volume of mesoporous carbons impregnated with polyethyleneimine (PEI) increased.
Impregnation of porous substrates with liquid amine results in a decrease of the free pore volume of the adsorbents. Analysis of the literature data of amine impregnated porous species demonstrates that the CO$_2$ capacity and efficiency of amine utilization drops when the volume of amine used for filling the pores approaches the total available pore volume, $V_{tot}$, of the original substrate$^{12,10,16}$ as a result of limited diffusion of CO$_2$ gas into the bulk of the adsorbent. Typical values of $V_{tot}$ for porous carbon substrates produced via resorcinol-formaldehyde condensation are in the range of 2-3.5 cm$^3$/g$^{13,17,18,19,20}$ (corresponding to a material porosity of 80-87%), although values of 5.35 cm$^3$/g$^{21}$ and 6 cm$^3$/g$^{22}$ (92% porosity) were also reported when special templating procedures were applied. This implies that for typical highly porous carbon substrates the maximum amount of amine which can be used for impregnation will not exceed 75-80% of the total weight of the adsorbent when all pores in the substrate are filled.

A promising class of materials which can provide larger values of internal volume available for filling with organic polyamines are aerogels.$^{23}$ Aerogels are ultralight materials with very high values of internal pore volume. Sol-gel chemistry methods result in cross linked hydro(organo)gels which, after drying and thermal annealing, were reported to produce aerogels with hierarchical pore structures.$^{24,25}$ Resorcinol and formaldehyde are commonly used precursors in the synthesis of all-carbon aerogels.$^{24}$ Building on this synthetic approach recent studies have reported the development of highly porous carbon aerogels that incorporate carbon nanotubes (CNTs) or graphene into the structure.$^{26,27,28,29}$ Carbon nanotube based aerogels were reported$^{22}$ to have extremely high pore volume and shown to be effective for sorption of organic$^{28,30}$ and inorganic$^{31}$ liquid spills. Cross-linking between carbon nanotubes or graphene flakes (GrOx) can occur via gelation of the carbon material with polyethylene oxide (PEO), PEI, surfactants, metal salts etc.$^{32}$, hydrothermal treatment$^{33}$ or even without additional cross-linking species just using carbon nanotubes as spacers during lyophilisation.$^{28}$ It has been shown that the gelation of carbon nanotubes/carbon nanofibres and graphene oxide may be assisted by small amounts of cross-linked PVA, which serves as a scaffold and provides larger pore size and available pore volume in the resultant gel.$^{30}$ These materials provide considerable scope for optimizing the pore structure to maximize the efficiency of CO$_2$ uptake as well as being possible to synthesize in the form of monoliths. The advantages of using CNTs/graphene in such structures can include enhanced internal pore volume, optimized electrical properties and a structure with greater mechanical stability.$^{27}$

In this work we provide the first study, to our knowledge, of the application of carbon nanotube/PVA aerogels as low weight solid supports for incorporating liquid amines in order to increase CO$_2$ uptake from dilute gas streams. Aerogels were prepared from various ratios of oxidised CNT and PVA by freeze drying. Their mechanical and electronic properties were determined and their stability with respect to organic and inorganic solvents was tested. The effect of impregnating aerogels with polyethyleneimine was studied and the CO$_2$ uptake was determined under dry conditions at a 0.1 bar CO$_2$ partial pressure which is in the range of partial pressures found in the flue gas of fossil fuel power stations. The material is shown to be competitive with the best CO$_2$ capture materials under conditions of atmospheric pressure and low CO$_2$ partial pressure with the additional advantage of allowing the development of an electrical swing desorption process due to the possibility of rapid ohmic heating.

2. Experimental Details

2.1. Aerogel Synthesis

Materials. Elicarb multiwall carbon nanotubes were purchased from Thomas Swan & Co Ltd; the carbon nanotubes have a diameter range of 8-12 nm, lengths of a few micrometres and contain 5-8 walls. Polyvinyl alcohol (PVA), 95% hydrolysed, M.W. 95000 Da was purchased from Acros Organics. Glutaraldehyde (GA), 25% solution in water was purchased from Sigma-Aldrich. Polyethyleneimine, branched, M.W. 600, 99% was purchased from Alfa Aesar; according to manufacturer information the material contains primary, secondary and tertiary amino groups in ratio 2:1:1.

Analytical grade methanol, sulfuric acid (>95%) and nitric acid (70%) were purchased from Fisher Scientific.

Oxidation of carbon nanotubes. Typically 1 g of carbon nanotubes was heated for 3-4 h in a 500 ml mixture of sulfuric (conc. 95%) and nitric (conc. 70%) acids, 3:1 v/v ratio. The oxidised carbon nanotubes were then isolated by vacuum filtration, and washed in sequence with dist. water, 3% NaOH, water, 3% HCl, water and methanol. As a result of oxidation a weight loss of 3-4% was observed. XPS analysis of the oxidised tubes showed 9 at. % of oxygen in the material in comparison to 0.5 at. % O in the purchased carbon nanotubes. The oxidation of carbon nanotubes was an essential step to ensure the formation of a stable aqueous dispersion.

Aerogel preparation procedure. Powdered oxidized multi-walled carbon nanotubes (further denoted as CNT), ca. 20 mg for all samples, were mixed with the calculated amount of stock solution of PVA in water (30 mg ml$^{-1}$) and diluted with water to the total volume of 3 or 5 ml in a graduated 5ml glass beaker. Three sample groups of aerogels were prepared with CNT-to-PVA weight ratios of 1:1, 1:2 and 1:3, i.e. 20 or 40 and 60 mg PVA were added to 20 mg of CNT. The beaker was placed in an ultrasonic bath and sonicated for 30 minutes and then stirred for 15 min at room temperature; this sequence was repeated at least 3 times until a stable dispersion was formed. Glutaraldehyde (25% in H$_2$O), 20 µl for each 20 mg of PVA, and HCl (2.5%), 35 µl for each 20 mg of PVA, were added to the mixtures, which were stirred at room temperature for an additional 30 minutes. The stirrer bars were removed and the CNT:PVA dispersions were left to gelate overnight at 50-55°C on a hot plate. The prepared hydrogels were placed in a cold bath at -20 to -25 °C until complete freezing, and then kept at a temperature of -20 °C for an additional 30 minutes. Frozen gels
were dried under vacuum, keeping a temperature of -20 °C for the first 15 minutes and then allowed to rise to room temperature; the complete removal of water and formation of aerogels was considered to have occurred when the residual pressure was 2-3 x 10⁻¹ mbar and required overnight pumping. Under these synthesis conditions a small reduction of aerogel volume with respect to the original volume of the slurry before freezing was observed for all series of produced aerogels. The final aerogel volumes were measured to be 4.25±0.35 ml for all CNT:PVA ratios for the '5ml' series and 2.36±0.15 ml for '3 ml' series of samples. There was no systematic effect of CNT:PVA ratio on the aerogel volume.

The density of the aerogels was extremely low with calculated densities ranging from ~10 mg cm⁻³ (CNT-to-PVA ratio 1:1) to ~25 mg cm⁻³ (CNT-to-PVA ratio 1:3) for '5ml' samples, and from 20 to 35 mg cm⁻³ for '3ml' samples respectively. The physical characteristics of the obtained aerogels are presented in Table 1.

### Table 1. Physical characteristics of the obtained aerogels

<table>
<thead>
<tr>
<th>Series</th>
<th>CNT:PVA wt ratio</th>
<th>Volume (ml)</th>
<th>Weight (mg)</th>
<th>Density (mg ml⁻¹)</th>
<th>Porosity, % (est.* *)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 ml</td>
<td>1:1*</td>
<td>4.2±0.3</td>
<td>43±3</td>
<td>10±1</td>
<td>99.35±0.05</td>
</tr>
<tr>
<td>1:2*</td>
<td>4.3±0.4</td>
<td>66±5</td>
<td>15±3</td>
<td>99.0±0.1</td>
<td></td>
</tr>
<tr>
<td>1:3*</td>
<td>4.3±0.3</td>
<td>87±7</td>
<td>20±3</td>
<td>98.6±0.07</td>
<td></td>
</tr>
<tr>
<td>3 ml</td>
<td>1:1*</td>
<td>2.3±0.1</td>
<td>46±1</td>
<td>20.3±0.6</td>
<td>98.8±0.03</td>
</tr>
<tr>
<td>1:2</td>
<td>2.4</td>
<td>66</td>
<td>28</td>
<td>98</td>
<td></td>
</tr>
<tr>
<td>1:3</td>
<td>2.5</td>
<td>84</td>
<td>34</td>
<td>97.5</td>
<td></td>
</tr>
</tbody>
</table>

*: The presented data show average values with standard deviation tolerance obtained from 3 to 5 samples of each kind.

**: porosity was estimated from the average measured aerogel monolith volume and volumes of CNT (density 1.9 g ml⁻¹) and PVA (density 1.19 g ml⁻¹).**

with a platform resting on the top surface. Masses of 5, 10, 20 and 50 g were placed on the platform and the resulting change in height was recorded using a camera fixed on a tripod. The final 50g mass was then removed from the aerogel and the height of the platform was recorded after five minutes to assess if the aerogel returned to its original size.

### 2.2.2 Estimation of aerogel's surface area and available internal volume

N₂ adsorption isotherms were measured at 77K with a Quantachrome NOVA 3000 gas sorption analyser. The aerogel monoliths were cut into small pieces (~5 mm) in order to fill the measurement cell without compressing the aerogel structure. The samples were regenerated in a degas station under vacuum (<1 mbar) at 90 °C to remove any adsorbed volatile species prior to analysis. The pore volume available for impregnation was also estimated by filling the aerogels with methanol. The aerogels were placed on a PTFE platform in a test-tube filled with excess of methanol, so that the methanol level was slightly below the platform. The capped test-tube was inclined until the sample on the platform was in contact with the methanol. When it was visually determined that aerogels were completely filled with methanol, the test-tube was returned to the vertical position and gently shaken to remove any excess methanol from the outside of the aerogel and the sample was removed from the test-tube. The amount of the absorbed methanol was measured by the test-tube weight difference before and after the soaking procedure. The available internal volume of the aerogel was considered to be equal to the volume of methanol absorbed by the aerogel.

**2.2.3 Impregnation procedure**

For impregnation of aerogel monoliths with polyethyleneimine the following procedure was applied: the calculated amount of PEI-600 was dissolved in methanol, so that the total solution volume would be slightly smaller (~80% to 90%) than the measured methanol volume taken up by the structure. This solution was introduced by halves onto the opposite faces of the aerogel monolith. After exposure to the liquid, the filled aerogel was placed on a PTFE substrate and dried in a vacuum desiccator gradually decreasing the pressure from 200 to 10 mbar at room temperature. This procedure was found to ensure a homogeneous distribution of amine within the aerogel monolith and precise dosing of added amine (see section 3.5).

**Figure 1.** Photograph of a typical 1:1 CNT:PVA aerogel and the set-up used to determine mechanical stability and compressibility of aerogel materials: left - aerogel monolith before the test, centre – aerogel monolith under the load of 50g and right – aerogel 5 min. after removing the load.
2.2.4 CO$_2$ adsorption measurements

The material’s CO$_2$ uptake was assessed by thermal gravimetric analysis (TGA). Measurements were carried out on a Setaram Sensys Evo TG/DSC instrument. 12-40 mg of aerogel impregnated with polyethyleneimine was placed in a platinum crucible that was counter-balanced by an identical platinum crucible packed with an equivalent mass of lead balls. Experiments were carried out at a CO$_2$ partial pressure of 0.1 bar and at 75°C, which is considered as optimal temperature for PEI impregnated supports,$^{13,10,35,36}$ although high CO$_2$ uptake at lower temperatures was also reported.$^{37}$ Samples were regenerated at 90 °C under helium flow (50 sccm) for 3 h before the sample temperature was adjusted to the desired experimental value of 75 °C. After the microbalance had stabilized, the helium flow (50 cm$^3$ min$^{-1}$) was switched to a mixture of CO$_2$ (5 cm$^3$ min$^{-1}$) and helium (45 cm$^3$ min$^{-1}$) for 4 h. The change in the sample mass corresponded to the uptake of CO$_2$ by the sample. The capacity and heat of adsorption were then calculated. A baseline subtraction procedure was followed prior to data analysis. Desorption was carried out at 75 °C in a flow of pure He (50 sccm) for 4 h.

2.2.5 Scanning electron microscopy

Scanning electron microscopy (SEM) measurements were carried out with a Carl Zeiss SIGMA HD VP Field Emission Scanning Electron Microscope with 10 kV beam energy.

2.2.6 Electrical conductivity and Ohmic heating

Electrical conductivity and Ohmic heating measurements were carried out using a Keithley 2612A sourcemeter with a custom-made 4-probe setup (the scheme shown in fig. 5a), similar to the scheme for measuring the resistivity of bulk materials described by Tupta.$^{38}$ Electrical current was applied to the faces of the aerogel, in the form of cylindrical monoliths, via round copper electrodes and the voltage inside the aerogel was measured with 0.2 mm tungsten probes. The temperature was measured with an Omega K-type precision fine wire thermocouple (0.2 mm) inserted into the middle of the aerogel monolith.

3. Results and Discussion

There have been many reports on the use of mesoporous substrates, mostly highly porous carbon and silica materials, impregnated with organic polyamines,$^{19,20,21,39,40,41,42}$ The amount of amine that can be incorporated into such substrates is limited by the total pore volume. The typical measured values of total pore volume for porous carbons prepared by a templating method or by formaldehyde-resorcinol sol-gel condensation (‘carbon aerogels’) are in the range 1-3.5 ml g$^{-1}$,$^{13,18,43}$ although pore volumes as high as 5.3 ml g$^{-1}$,$^{21}$ and 6 ml g$^{-1}$,$^{22}$ have been reported. Ultra-light aerogels (foams) made of gelated organic polymers with carbon filler, such as the materials of interest here, show much higher values of total internal volume, with reported porosity values >98%.$^{27,30,44}$ For the purposes of selective carbon capture, requiring elevated temperatures of adsorption/desorption within the range 75 to 90 °C, the ultra-light aerogels (sponges) should exhibit good thermal stability. In addition, they require high chemical stability towards the organic amines used for impregnation and high mechanical stability to enable the use of monolithic materials to reduce the pressure drop in a practical carbon capture process. The materials synthesised from PVA/CNT mixtures reported here satisfy all these requirements: (i) crosslinked PVA forms stable hydrogels in aqueous solutions, (ii) PVA has a high melting temperature (185 °C) and (iii) PVA does not react with amines. Additionally, the presence of hydroxyl groups in the composite was considered to provide good wetting with amines and also could facilitate the reaction of amines with CO$_2$ via formation of alkyl carbonate salts$^{45,46}$ or carbamate esters.$^{47}$

The lyophilisation technique makes it possible to tune the resultant density of the aerogel by varying the initial concentrations of CNT and PVA, i.e. by changing the amount of water in the hydrogel. During synthesis, the mass of CNTs was kept close to 20 mg whilst the quantity of PVA was varied in order to synthesise aerogels with the desired CNT:PVA ratio. The dispersions of CNT in PVA solutions formed gels after 5-10 h of heating at 50-55 °C with negligible phase separation. The synthesized aerogels were mechanically stable. The aerogels with a 1:1, 1:2 and 1:3 wt ratio of CNT:PVA had dry masses of approximately 40, 60 and 80 mg respectively, i.e. freeze drying removed practically all the water from the hydrogel.

The presence of CNT is important for hydrogel formation. Pure PVA solutions at equivalent concentrations of 0.4 – 1.6% (20 – 80 mg PVA in 5 ml), which is below the critical PVA gelation concentration in water,$^{48,49,50,51}$ after adding the cross-linking agent and heating at 50-55 °C for 24 h, became cloudy, but did not form bulk hydrogels. Freeze drying of solutions of pure PVA after cross-linking did result in stable aerogels, in agreement with the literature.$^{49,52,53}$ All aerogel samples made of pure PVA underwent significant shrinkage during freeze-drying, the final volumes being 2.6 ml for the 20 mg sample (density 7.6mg ml$^{-1}$), 2.9 ml for the 40 mg sample (density 15.6 mg ml$^{-1}$), 3.3 ml for the 60 mg sample (density 20.6 mg ml$^{-1}$) and 3.05 ml for the 80 mg sample (density 26.8 mg ml$^{-1}$). An attempt to carbonise the CNT:PVA aerogels at 1000 °C in Ar flow did not produce rigid all-carbon aerogels and resulted in decomposition of the PVA component of the aerogel monolith and recovery of decarboxylated carbon nanotube powder.

3.1. Aerogel appearance

The prepared aerogel monoliths are black spongy materials in cylindrical form with a diameter of ~1.8 cm (corresponding to the inner diameter of the slurry container) and height of 1.5-1.9 cm for an initial slurry volume of 5 ml. Fig. 1 shows a photograph of a typical aerogel sample prepared from a gelated 5 ml slurry. The internal pore structure of the materials was investigated by scanning electron microscopy (SEM). A selection of images is presented in fig. 2. From the low magnification images (top row), a well-defined macroporous structure was observed, with large micrometre size pores (up to a few tens of μm) distributed...
across the surface. At higher levels of magnification (bottom row), it is clearly seen that CNTs were well distributed throughout the polymer phase. From an examination of the SEM images it is possible also to conclude that the increase of polymer fraction in the aerogel from 50% (1:1 CNT:PVA ratio) to 75% (1:3 ratio) resulted in a dense local distribution of material and some growth of the size of internal voids.

3.2. Characterisation of aerogel pore structure.

Liquid nitrogen adsorption isotherms of aerogels prepared with different PVA content are shown in Fig. 3. The isotherm curves are all similar and correspond to Type III isotherms according to the IUPAC classification, typical for macroporous materials with pore size close to 100 nm. However this technique is not suitable for characterisation of pore structure and pore size distribution in macroporous materials. The volume of macropores in the aerogel samples was estimated by measuring the amount of methanol that could be absorbed by the aerogel when it was put in contact with the liquid. The volume of the absorbed methanol was found to be ~75% of the measured volume of aerogel cylindrical monoliths with 1:1 and 1:3 CNT:PVA ratio, and ~70% for 1:2 species, i.e. an aerogel monolith of 4 ml absorbs ca. 3 ml of methanol. Thus, the volume available for filling with liquid in the prepared CNT:PVA aerogel sponges ranged from 35 ml g⁻¹ for species with a CNT:PVA ratio of 1:3 to >70 ml g⁻¹ for 1:1 species.

3.3 Compression test.

A simple setup for testing the mechanical properties of CNT:PVA aerogels is shown in Fig. 1. After removing the weight all tested aerogels recovered to >90% of their original size within 5 minutes. The results of the compression test are presented in Fig. 4. The stiffness of the aerogels grows with an increase of PVA content in the material, corresponding to an increase of the material density. These simple experiments provided an estimation of the compressive Young’s modulus values of 5±0.4 kPa for aerogels with a CNT:PVA wt ratio of 1:1 and density ~10 mg ml⁻¹, 7.3 ±1.1 kPa for samples with ratio of 1:2 and density ~15.5 mg ml⁻¹, and 14.3±1.9 kPa with a CNT/PVA wt ratio of 1:3 and density ~21 mg ml⁻¹. These values are consistent with the
published Young’s modulus data for other ultra-light foams, e.g. 33 kPa for a polyurethane foam with density of 16 mg ml$^{-1}$ and >80 kPa for polyurethane foams with density of 90 mg ml$^{-1}$.

As expected, the resistivity of the aerogels dramatically increased with a decrease of the CNT content, ranging from 19 kΩ cm for materials with a CNT-PVA ratio of 1:1; 2 MΩ cm for a CNT-PVA ratio of 1:2 and 5 MΩ cm for species with a CNT-PVA ratio of 1:3. Non-impregnated aerogel samples with a CNT-PVA ratio of 1:1 and 1:2 could easily be heated to temperatures above 70 °C, and demonstrated a resistive heating dependence of 100 – 145 K g W$^{-1}$. Aerogels with a CNT-PVA ratio of 1:3 demonstrated a heating rate of 280 – 370 K g W$^{-1}$, although due to instrument limitations (max. compliance voltage of 200V) we were unable to reach temperatures above 40°C for these samples.

3.5 Impregnation and CO$_2$ capture

In order to investigate the behaviour of aerogels during the wet impregnation procedure$^{10,16,62}$ as well as to provide an estimate of the available internal volume, the uptake of methanol by the aerogels was investigated (sect. 3.2). The wet impregnation procedure, typically used for porous carbons$^{10,18}$ and silicas,$^{62}$ where the porous material powder is mixed with the calculated amount of amine dissolved in an arbitrary amount of solvent followed by solvent evaporation, cannot be applied directly to aerogel sponges because a uniform distribution of amine inside the aerogel cannot be achieved. Therefore, the impregnation method applied here consisted of complete absorption of a measured volume of amine solution into the aerogel sponge, so that the total volume of the absorbed liquid was roughly equal to 85% of the aerogel pore volume as estimated by methanol uptake. This amount of liquid was found to provide complete wetting of the aerogel throughout the bulk, but avoided excessive amine concentration on the surface of the monolith during the evaporation step. The applied vacuum drying scheme with slow methanol evaporation allowed significant shrinkage of the impregnated aerogels to be avoided. Fig. 6 shows the aerogel species before and after impregnation with 3-fold, 5-fold and 10-fold amounts of PEI600; the dimensions of impregnated aerogel species do not change significantly.

The resistivity was measured in the low power region; when the power applied to the aerogel grew beyond the values where the materials were subjected to Ohmic heating (>0.5 W g$^{-1}$ for all species), the resistivity exhibited non-linear behaviour. The summary plot with resistivity of the prepared CNT:PVA aerogels is shown in fig. 5b.

**3.4 Electrical measurements**

In contrast to pure PVA aerogels, all CNT containing aerogels were electrically conducting. The I-V characteristics of aerogels were measured using a 4-probe scheme (fig. 5a) and electrical resistivity was calculated using the formula:$^{38}$

$$\rho = \frac{V}{I} \times \frac{A}{d}$$

where $\rho$ is the bulk resistivity of the material in Ω cm, $V$ is the measured voltage, $I$ is the electrical current, $A$ is the surface cross-section and $d$ is the distance between probe leads.

![Diagram](image_url)

Figure 5. a) schematic drawing of the experimental set-up for measuring electric resistivity of aerogel moulds; b) resistivity of the aerogels with different CNT:PVA weight ratio.

![Diagram](image_url)

Figure 6. Photographs of CNT-PVA (1:3 wt ratio): left – pieces of aerogel cut from the monolith before impregnation and right – the same pieces of aerogel after impregnation with 3-fold, 5-fold and 10-fold amounts of PEI600.

This approach allowed the amount of added amine to be controlled within a few percent and provided uniform distribution of amine in the substrate. Fig. 7 shows SEM images of aerogels impregnated with various amounts of PEI. It is obvious that species containing 75% (x3) and 83.3% (x5) of...
amine still retain the 3D morphology inherent to the initial aerogel. Further increase of amine content in the material (amine to substrate ratio x10, amine content in the sorbent 91%) resulted in the formation of continuous layers of amine inside the porous substrate.

Figure 7. SEM images of aerogel (CNT-PVA ratio 1:1) (a) before impregnation, (b) after impregnation with 3-fold amount of PEI (x3), (c) 5-fold amount of PEI (x5), (d) 10-fold amount of PEI (x10).

The volume available for impregnation in CNT:PVA aerogels (35-70 ml g⁻¹) significantly exceeds the values of 2.5 -6 ml g⁻¹ that have been reported for typical porous materials used as supports for liquid amines.13,20,22 This implies that, contrary to typical highly porous carbon substrates with complete pore filling (blocking) at 70-80% of amine content, leading to a decrease in amine utilization efficiency,15,18 the maximum amine loading in the impregnated aerogels can be higher than 85% wt without blocking the pores, thus, providing gas diffusion through the bulk of the adsorbent. Note that the theoretical capacity for CO₂ uptake with polyethylenimine of the general formula H-(NH-CH₂CH₂)ₙ-NH₂ with n =13.6 (PEI600) under dry conditions with formation of carbamate salt according to the generally accepted mechanism93 (scheme 1) is ca. 12.2 mmol g⁻¹.

For materials with a substrate impregnated with 3 fold wt amount of amine (corresponding to amine content in the material of 75%) the theoretical CO₂ capacity is 9.15 mmol g⁻¹, for x5 impregnation (83.3%) the theoretical CO₂ capacity is 10.17 mmol g⁻¹ and for x10 impregnation (91%) it is 11.1 mmol g⁻¹ (fig. 8), assuming 100% utilization of the NH units, i.e. without taking into account the reverse reaction at 75°C.

For example, for solid ammonium carbamate the rates of formation and decomposition reactions are the same at ~70°C64 (Gibbs energy change of the reaction, ΔG = 0). Fernandes et al.65 investigated the formation of carbamate salts for various primary and secondary amines in solution by ¹H NMR, measured equilibrium stability (formation) constants and determined standard molar enthalpy and entropy values for this reaction. We calculated equilibrium constants for carbamate salt formation at 75 °C using data from Fernandes et al. 65 and obtained the values of K_carbamate = 5.2 for monoethanolamine (MEA), ~1.5 for secondary amines (morpholine and piperazine), and 0.5 for ammonia. These values correspond well to the analysis of Gupta et al.66, who showed stability constants at 345 K of 2.7 and 1.55 for carbamate salts of MEA and diethanolamine respectively, and a decomposition equilibrium constant of ~1 for ammonium carbamate at 70 °C.64 These equilibrium constants correspond to conversion to carbamates of ~60% of secondary amines and ~75% of primary amines under equilibrium conditions. The expected CO₂ equilibrium uptake by a substrate impregnated with polyethylenimine with ~25% of primary amino groups and ~50% of secondary amino groups will be ~65-70% of the theoretical value (fig.8).

Under adsorption conditions with excess of CO₂, the carbamate salt formation reaction (Scheme1) will be shifted to the products, and, therefore, the real uptake values will be in between the theoretical maximum uptake and the equilibrium uptake values.

Figure 8. Theoretical maximum CO₂ uptake (solid squares) and estimated equilibrium CO₂ uptake at 75°C (hollow circles) by porous species impregnated by PEI600 under dry conditions as a function of amine to substrate wt ratio.
The CO₂ capacities of PEI impregnated CNT:PVA aerogel monoliths are very competitive with respect to other porous supports impregnated with branched PEI600 and measured with the same method (table 3). Impregnated aerogels with a CNT:PVA wt ratio of 1:1 demonstrated on average better values of CO₂ capacity per gram than more dense species with higher PVA content in the aerogel (3.3 mmol g⁻¹ vs 2.6 mmol g⁻¹). The volumetric uptake at ca. 0.3 mmol CO₂ per ml of sorbent is a about a factor of two or three less than that reported for impregnated carbon black materials at a partial pressure of 1 bar. The efficiency of utilization of amino groups under dry conditions for 1:1 samples was 33-36%, which is significantly higher than ≤25% efficiency that we observed earlier for mesoporous carbon impregnated with PEI600. CO₂ uptake curves for aerogels impregnated with polyethyleneimine at 75 °C, 0.1 bar CO₂ show fast initial sorption followed by a second, slower process (fig. 9a), which is typical for porous substrates impregnated with organic amines (see references given in table 3).

The second, slow sorption stage is associated with slow diffusion of CO₂ through the amine and formation of reaction products. This potentially could be improved by further optimising the pore structure and thickness of the amine layer or by using other schemes to increase the CO₂ diffusivity through the layer of PEI, e.g. by using various additives. The CO₂ uptake behaviour of aerogels loaded with PEI600 naturally depends on the amount of amine in the substrate. Fig. 9a shows CO₂ uptake curves for aerogels with a CNT:PVA ratio of 1:1 loaded with 3- and 5-fold weight amounts of PEI600. Although the CO₂ capacity values after 4 h of sorption are fairly close (3.66 and 3.54 mmol g⁻¹ for x5 samples vs 3.55 mmol g⁻¹ for x3 sample) the slope of the curve for the x5 sample is steeper indicating that the process is still far from equilibrium after 4 h. When the CO₂ uptake for a x5 sample (different batch) was measured for 10h the slow sorption step after 3h of experiment exhibited behaviour close to linear with a CO₂ uptake rate of 0.075 mmol g⁻¹ h⁻¹. Fig. 9b shows the extended region of the CO₂ uptake TGA curves. For both x5 samples, despite a slightly different total CO₂ capacity, the sorption behaviour is similar, while for the x3 sample the slope beyond 3 h is much lower, indicating that the stock of unreacted amino groups is small and the reaction of CO₂ with amino groups may be close to equilibrium at 75°C.

![Figure 9](https://example.com/figure9.png)

Table 3. Literature data reporting high TGA CO₂ uptake for various silica and carbon based porous substrates impregnated with PEI600.

<table>
<thead>
<tr>
<th>Porous support</th>
<th>PEI 600 content, %</th>
<th>P CO₂, bar</th>
<th>T, °C</th>
<th>qCO₂, mmol g⁻¹</th>
<th>Ref.</th>
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<tr>
<td>Hexagonal mesoporous silica</td>
<td>65</td>
<td>1</td>
<td>75</td>
<td>4.18</td>
<td>Chen 67</td>
</tr>
<tr>
<td>Silica microcapsules</td>
<td>83</td>
<td>1</td>
<td>75</td>
<td>&gt;5</td>
<td>Qi 68</td>
</tr>
<tr>
<td>Silica foam</td>
<td>70</td>
<td>0.1</td>
<td>75</td>
<td>2.3</td>
<td>Subagyono 69</td>
</tr>
<tr>
<td>MCM-48</td>
<td>70</td>
<td>1</td>
<td>80</td>
<td>3.1</td>
<td>Sharma 12</td>
</tr>
<tr>
<td>Carbon black</td>
<td>50</td>
<td>1</td>
<td>75</td>
<td>3.1</td>
<td>D Wang 16</td>
</tr>
<tr>
<td>Hierarchical porous silica</td>
<td>70</td>
<td>1</td>
<td>75</td>
<td>4.1</td>
<td>J Wang 18</td>
</tr>
<tr>
<td>Silicalgel</td>
<td>50</td>
<td>0.1</td>
<td>75</td>
<td>2.9</td>
<td>Zhang 20</td>
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<tr>
<td>Mesoporous carbon spheres</td>
<td>50</td>
<td>1</td>
<td>75</td>
<td>2.9</td>
<td>M Wang 20</td>
</tr>
<tr>
<td>Mesoporous Carbon</td>
<td>73</td>
<td>0.1</td>
<td>75</td>
<td>2.3</td>
<td>Gibson 21</td>
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<tr>
<td>CNT:PVA aerogel sponge</td>
<td>75</td>
<td>83</td>
<td>0.1</td>
<td>75</td>
<td>3.25±0.3</td>
</tr>
</tbody>
</table>

![Table 3](https://example.com/table3.png)

**Table 2.** CO₂ capacities of CNT:PVA aerogels loaded with PEI600, measured at 75 °C, 0.1 bar CO₂ under dry conditions, are presented in table 2.
At a reaction time of 4h the amine utilization efficiency is slightly higher for the x3PEI sample (0.36 vs 0.33 for the x5PEI sample). This difference is significantly higher for the initial fast step and after the first 10 min (0.167 h) of sorption (fig. 9c), where the x3 PEI sample demonstrated a CO₂ capacity of 2.46 mmol g⁻¹ with an amine utilization efficiency of 27%, i.e. 0.2 g PEI per g of the sorbent reacted with CO₂, compared to 1.91 mmol g⁻¹ with an efficiency of 19% for the x5 PEI sample (0.16 g PEI per g of the sorbent). Assuming a uniform distribution of amine on the surface of the aerogel with a surface area of 60 m² g⁻¹, the PEI layer should have a thickness of 50 nm for the x3 PEI species and 83.5 nm for the x5 PEI species. In the case of 100% utilisation of amino groups for reaction with CO₂, it is possible to estimate that in the x3 PEI sample after 10 min of exposure to CO₂, only the first 14 nm of the amine layer are involved in the reaction. The access to PEI which is deeper under the surface is hindered by diffusion of CO₂ through the layer of liquid reaction products. For x5 PEI such an estimate gives a similar value of 15.5 nm of PEI thickness involved in the reaction during the fast initial step.

Conclusions

A series of ultra-light aerogels made of oxidized carbon nanotubes and PVA has been prepared by freeze drying of hydrogels, characterised, and tested as amine impregnated solid supports for CO₂ capture. The aerogel spongy materials were prepared with CNT:PVA wt ratios of 1:1 to 5 M CNT:PVA values of the compressive Young’s modulus of 5 to 14.3 kPa. Published data for other ultra-light aerogels have rather low surface areas (43-61 m²/g), but were shown have very high values of internal volume available for filling (impregnation), 35 to >70 ml g⁻¹, depending on the density of species. Aerogels with all tested CNT:PVA weight ratios are mechanically stable due to good component homogeneity; the mechanical properties of the prepared aerogels are consistent with published data for other ultra-light foams and demonstrate values of the compressive Young’s modulus of 5 to 14.3 kPa. CNT:PVA aerogels are electrically conducting; a decrease of the CNT content in the aerogel resulted in a dramatic increase of the material bulk resistivity from 19 kΩ.cm for species with a CNT:PVA wt. ratio of 1:1 to 5 MΩ.cm for species with a CNT:PVA wt. ratio of 1:3. Aerogels with a CNT:PVA wt. ratio of 1:1 and 1:2 are suitable for electrical swing adsorption (ohmic heating) but the 1:3 ratio materials are impractical as they require voltages >200V for reaching temperatures feasible for solid sorbent regeneration.

Using a modified wet impregnation procedure, samples were prepared with a homogeneous distribution of amine (PEI600) within the aerogel monolith and precise dosing with an amine content of 75 to 91%, corresponding to 3-fold (x3) to 10-fold (x10) amount of added amine with respect to the weight of the substrate. In x3 and x5 impregnated species the amine layer coats the initial surface texture inherent to the original aerogel, while in x10 samples the amine layer is thick and forms a smooth and continuous surface. PEI600 impregnated aerogels were tested under 10% CO₂ partial pressure conditions in order to simulate the flue gas of a fossil fuel power plant. The obtained CO₂ capacity values significantly exceed our previous results obtained with impregnated mesoporous carbons, and are competitive with the highest reported values for porous substrates impregnated with similar amines. This makes ultra-light CNT:PVA aerogel sponges very promising supports for amine impregnated solid sorbents for CO₂ capture. The value of the CO₂ capacity was dependent on the structure of the carbon support; the species with highest CNT content (CNT:PVA ratio 1:1) demonstrated on average the highest uptake values of ~3.3±0.3 mmol g⁻¹ for x3 and x5 samples and an efficiency of amino groups utilisation of >0.35. The analysis of TGA adsorption curves showed that the initial fast sorption step occurs during the first 10 minutes and involves the outer PEI600 layer to a depth of ~15 nm, beyond which further CO₂ uptake is limited by diffusion through the top layer of the products.

Conflicts of interest

There are no conflicts to declare.

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Notes and references


