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COVER ARTICLE
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Bio-based macroporous polymer nanocomposites made by mechanical frothing of acrylated epoxidised soybean oil

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Mechanical frothing is one of the most commonly used methods to create gas-liquid foams. Until recently, the polymerisation of mechanically frothed gas-liquid foams was limited to the synthesis of quasi two-dimensional polymer structures, such as films. In this study we show that three-dimensional bio-based polymer foams can be created by microwave curing of gas-soybean oil foams created by mechanical frothing using lauril peroxide as the radical initiator. It was found that the introduction of air during the mechanical frothing was necessary to create the three-dimensional polymer foams. Using bacterial cellulose nanofibrils (BC) simultaneously as a foam stabiliser has potential because it obstructs the flow of liquid from the lamella region in these gas-soybean oil foams while simultaneously acting as nano-filler in the polymer foam. It was found that the stability of the gas-soybean oil foam templates and the mechanical properties of the polymer nanocomposite foams are enhanced upon the addition of BC in to the foams.

Introduction

Polymer foams, which represent a group of lightweight materials, were a £7.5 billion industry in 2010 and are expected to reach £8.6 billion by 2015. Their very high porosity and low density makes polymer foams attractive materials for both science and engineering applications. Generally, polymer foams can be classified into rigid or flexible foams, open- or closed-cell foams. Open-cell foams are used as scaffolds for tissue engineering or catalyst supports whereas closed-cell foams have found application in sandwich structures and thermal insulation. Generally, the interconnected nature of open-cell foams provides better permeability while closed-cell foams offer low or zero permeability, which results in better insulating properties.

Polymer foams can be produced in various ways; by using chemical or physical blowing agents, thermally induced phase separation (TIPS), sintering of particles/microspheres or polymerising the continuous phase of suitable emulsion templates. A novel technique to produce foams has been reported by Murakami and Bismarck. The authors utilised oligomeric tetrafluoroethylene (OTFE) particles to stabilise air bubbles in monomers and UV-polymerised this foam to produce macroporous closed cell polymers. A similar technique was also used to produce macroporous ceramics; a wet foam was produced by mechanically frothing a suspension of hydrophobic alumina powder. The alumina powder serves as foam stabiliser. After a drying step to remove the water followed by sintering of the alumina, Gonzenbach et al. were able to produce macroporous ceramics with porosities up to 88%. However, the polymerisation of mechanical frothed gas-monomer foams remain under studied. Marlin et al. first studied macroporous polymers produced from mechanically frothed air-urethane foams. The authors mechanically frothed a mixture of isocyanate, polyols, surfactants and catalyst with air, followed by thermal polymerisation of the monomer to produce polymeric foams. However, the authors only managed to produce quasi two-dimensional (2D) polymer foams of 6 mm in thickness. Greer and Wilkes mechanically frothed a highly viscous oligomer solution, polymerisation was initiated using a 175 keV electron beam to produce 2D thin (0.5 mm thick) polymer foams. They also used the same technique to produce polymeric foams from surfactant stabilised low viscosity monomers. However, the use of a high-energy electron beam reduces the commercial value of this technology.

In this study, a soybean oil-derived renewable monomer was first studied macroporous polymers produced from mechanically frothed air-urethane foams. The authors mechanically frothed a mixture of isocyanate, polyols, surfactants and catalyst with air, followed by thermal polymerisation of the monomer to produce polymeric foams. However, the authors only managed to produce quasi two-dimensional (2D) polymer foams of 6 mm in thickness. Greer and Wilkes mechanically frothed a highly viscous oligomer solution, polymerisation was initiated using a 175 keV electron beam to produce 2D thin (0.5 mm thick) polymer foams. They also used the same technique to produce polymeric foams from surfactant stabilised low viscosity monomers. However, the use of a high-energy electron beam reduces the commercial value of this technology.
hand, Bonnaillie et al.\textsuperscript{30} pressurised CO\textsubscript{2} to 60 bar and used it as the physical blowing agent to produce soybean oil-derived macroporous polymers.

We also produced high porosity macroporous soybean-oil derived polymer nanocomposites by dispersing bacterial cellulose nanofibrils (BC) in the monomer phase. BC is produced by the \textit{Acetobacter} species\textsuperscript{22} and has the advantage of being free of wax, lignin, pectin and hemicellulose. It is highly crystalline in nature, with a degree of crystallinity of approximately 90%\textsuperscript{23}. BC inherently has nanometre dimensions (24–86 nm in diameter, see Fig. 1) and unlike nano-fibrillated cellulose (NFC) derived from plant-based cellulose sources, does not require nanofibrillation.\textsuperscript{24} Hsieh et al.\textsuperscript{25} predicted that BC possesses a Young’s modulus of 114 GPa. These properties make BC an attractive nanofiller to reinforce fine structures, such as fibres, foams\textsuperscript{26} and the matrix of composite materials.\textsuperscript{27,28} Our current study focuses on a solvent- and surfactant-free process operating at atmospheric conditions to produce bio-based monomer foam templates, which can be polymerised to yield polymer foams. By reinforcing soybean oil derived polymer foams with BC, a new type of green macroporous polymer nanocomposites can be manufactured via simple mechanical frothing of monomer.

![Fig. 1](image1.png) Scanning electron micrographs of bacterial cellulose nanofibrils. Adapted from reference 40 with permission from Elsevier.

**Results and discussion**

**Stability indices of the liquid foam**

The stability of gas-epoxidised acrylated soybean oil (AESO) liquid foam templates is very important for the resulting polymer foams. The stability index of gas-AESO liquid foams was determined by tracking the movement of the air-monomer interface due to creaming visually. Photographs of the mechanically frothed gas-AESO and gas-AESO-BC liquid foams are shown in Fig. 2. The ability of the gas-AESO and gas-AESO-BC liquid foams to flow under gravity is shown in Fig. 3. It can be seen from Fig. 3 that AESO 1 flows easier than AESO 2 and 3. This is due to the viscosifying effect of BC in AESO. BC is known as a thickening agent, stabiliser and texture modifier.\textsuperscript{31} The stability of the liquid foams as a function of time is shown in Fig. 4. The neat liquid AESO foams without BC (AESO 1) destabilised rapidly compared to the liquid AESO foams containing 0.5 wt\% and 1 wt\% of BC (AESO 2 and 3). In fact, AESO 2 and AESO 3 showed insignificant changes in terms of the liquid foam height even 70 days after the foams were prepared. In contrast to AESO 2 and 3, the liquid foam height of AESO 1 reduced to half of its original height just 5 days after preparation and fully phase separated after 70 days.

The stability of liquid foams is largely governed by the dynamics of the thin interfacial films, which formed between air bubbles approaching each other. Two main mechanisms are involved in liquid foam destabilisation: gravitational and capillary drainage.\textsuperscript{12} The liquid foam will first undergo gravitational drainage, whereby two adjacent bubbles approach each other, resulting in the reduction of the foam lamella thickness. The drained liquid will move to the edge of the planar foam lamella,
known as the Plateau border. A second mechanism known as capillary drainage will become dominant when the pressure in the foam lamella is larger than the pressure in the Plateau border. As the monomer drains from the foam lamella to the Plateau border, due to pressure differences, the air bubbles will approach each other resulting in bubble coalescence.\textsuperscript{13} The foam AESO \textsuperscript{1}, which does not contain BC, exhibited rapid destabilisation due to the expected fast kinetics of gravitational drainage, followed by capillary drainage, which ultimately resulted in the full phase separation of the foam by 70 days. When the monomer phase contains BC (AESO \textsuperscript{2} and AESO \textsuperscript{3}), the kinetics of destabilisation were significantly reduced. It has been proposed that nano-particles in the liquid phase of foams will aggregate in the Plateau border, obstructing the flow of the liquid from the lamella.\textsuperscript{14} Ultimately, this liquid flow obstruction hinders the coalescence of bubbles in foams.

Structure and morphology of the macroporous polymer nanocomposites

SEM images showing the internal structure and morphology of the poly(merised)AESO (polyAESO) foams are shown in Fig. 5. It can be seen from the SEM images that the pores are randomly oriented with pore shapes that are spherical, semi-ellipsoidal or highly irregular, both from the top and side views of the materials. Pore throats can be seen in the pores, indicating that the pores are interconnected. However, the pore size is highly non-uniform throughout the samples (Fig. 5). The pores in the macroporous polymer are formed by the polymerisation of the liquid monomer foam templates produced by mechanical frothing; with some contribution from the additional CO\textsubscript{2} released by the thermal decomposition of the initiators (one mole of initiator produces two moles of CO\textsubscript{2}).

With increasing BC loading, the monomer phase became more viscous (Fig. 3) and the expansion of the bubbles in the gas-monomer liquid foams could not reach its equilibrium state within the time frame of the polymerisation of AESO. This quasi-equilibrium state of the bubbles results in the randomly orientated ellipsoidal pores (polyAESO \textsuperscript{1} and \textsuperscript{2}). Increasing the BC concentration to 1 wt\% (polyAESO \textsuperscript{3}) increases the viscosity of the continuous phase even further and results in polymer foam structures with very irregular and ill-defined pores.

Porosity of the macroporous polymer nanocomposites

Porosity is one of the important factors that govern the physical properties of macroporous polymers. Both the measured envelope and matrix densities along with the calculated porosity of the macroporous polymer and its nanocomposites are tabulated in Table 1. It can be seen that the envelope density increased from 0.44 g cm\textsuperscript{-3} to 0.58 g cm\textsuperscript{-3} and the porosity of the polyAESO decreases from 59\% to 45\% with increasing BC content. The introduction of BC into the monomer phase resulted in difficulties to mechanically froth the liquid at constant input of frothing energy. Even though the thermal degradation of the initiator could potentially contribute to the porosity (approximately 130 cm\textsuperscript{3} of CO\textsubscript{2} produced per 100 mL of AESO used), the images shown in Fig. 6 show otherwise. Polymerising AESO containing 1 wt\% of initiator could potentially result in a theoretical porosity of 56\%. However, the microwave irradiation also increased the temperature of the monomer to approximately

<table>
<thead>
<tr>
<th>Sample</th>
<th>$\rho_e$ (g cm\textsuperscript{-3})</th>
<th>$\rho_m$ (g cm\textsuperscript{-3})</th>
<th>$P$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>polyAESO \textsuperscript{1} (0 wt% BC)</td>
<td>0.44 ± 0.01</td>
<td>1.07 ± 0.01</td>
<td>59 ± 1</td>
</tr>
<tr>
<td>polyAESO \textsuperscript{2} (0.5 wt% BC)</td>
<td>0.49 ± 0.01</td>
<td>1.08 ± 0.01</td>
<td>54 ± 1</td>
</tr>
<tr>
<td>polyAESO \textsuperscript{3} (1 wt% BC)</td>
<td>0.58 ± 0.01</td>
<td>1.08 ± 0.01</td>
<td>45 ± 1</td>
</tr>
</tbody>
</table>

Fig. 4 The stability index of the mechanically frothed gas-monomer liquid foams. AESO \textsuperscript{1} : 0 wt\% BC, AESO \textsuperscript{2} : 0.5 wt\% BC and AESO \textsuperscript{3} : 1 wt\% BC.

Fig. 5 SEM images of the polyAESO. (a) and (b) are polyAESO \textsuperscript{1} (0 wt\% BC), (c) and (d) are polyAESO \textsuperscript{2} (0.5 wt\% BC), (e) and (f) are polyAESO \textsuperscript{3} (1 wt\% BC).
A comparison between non-mechanically frothed and air templated polyAESO. The photograph on the left shows polyAESO that was not mechanically frothed prior to its polymerization and the image on the right shows the air templated polyAESO 1 (0 wt% BC).

At this temperature, the viscosity of the monomer phase is low enough for the CO₂ produced to rise to the air-monomer interface and escape from the monomer phase before it gels and cured. Therefore, non-frothed AESO did not polymerise into porous polymers (see Fig. 6). By frothing the monomer phase, air bubbles are introduced. It is hypothesised that the CO₂ produced can escape from the monomer phase to the air-monomer interface at the top of the cylindrical vessel or into the air bubbles formed during mechanical frothing. The latter resulted in the expansion of the air bubbles during the polymerisation of the monomer. This might also explain why ellipsoidal pores are observed in polyAESO. The ellipsoidal pores could also be a result of buoyancy induced creaming.

**TGA of the macroporous polymer nanocomposites**

The thermal degradation behaviour of the macroporous polymer and its nanocomposites is shown in Fig. 7. It can be seen that all the samples undergo single step degradation in nitrogen atmosphere. Random polymer chain scission occurred during the degradation of polyAESO. The introduction of BC into the sample did not alter the degradation behaviour of polyAESO. This might be due to the low BC content in the sample. The onset degradation temperature determined from TGA analysis of the foams was found to be 380 °C for all samples. In addition to this, the residual carbon content for polyAESO 1 was found to be 0.98 wt% and approximately 2 wt% for both polyAESO 2 and 3. The increase in the residual carbon content of polyAESO 2 and 3 compared to polyAESO 1 can be explained by the carbonisation of BC in the nanocomposites.

**Compression properties of the macroporous polymer nanocomposites**

The mechanical properties of the macroporous polymer nanocomposites determine their potential applications. The Young's modulus and compression strength, along with the specific Young's modulus and compression strength (absolute compressive properties divided by the envelope density of the material) of the macroporous polymer and its nanocomposites are summarised in Table 2 and characteristic load-displacement curves of the linear elastic region are shown in Fig. 8. It should be noted that the polyAESO with different BC loadings possess different porosities and, therefore, we compare only the specific compressive properties of the polyAESO. By adding 0.5 wt% of BC (polyAESO 2), the specific Young’s modulus increased from 253 MPa kg⁻¹ m⁻³ to 339 MPa kg⁻¹ m⁻³. Even at such low nanofiller loading fraction, a 35% increase in the specific Young’s modulus was observed. This increase in foam stiffness can be attributed to the stiffness of BC, which has been estimated to possess a Young’s modulus of 114 GPa. However, when the nanofiller content was increased to 1 wt% BC (polyAESO 3), the specific modulus decreased by 60% to 90.2 MPa kg⁻¹ m⁻³ when compared to polyAESO 1. This decrease in the compressive modulus of polyAESO 3 is thought to be due the ill-defined

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† This temperature was obtained by measuring the temperature of the microwave-irradiated monomer without thermal initiator immediately after irradiation.
pore structure due to the inclusion of BC into the macroporous polymer (see Fig. 5). The specific compression strength of the polyAESO showed a different trend compared to the Young's modulus. It was observed that the compression strength of the polyAESO decreased with increasing BC content (Table 2). The compression strength of the polymer nanocomposite foams decreased by as much as 60% from 777 kPa kg⁻¹ m³ to 315 kPa kg⁻¹ m³ when the BC content was increased to 1 wt%. It is proposed that the poor specific strength of the nanocomposites is due to the poor compatibility between polyAESO and BC and hence, poor stress transfer between BC and the polymerised AESO.

The presence of unreacted AESO in the macroporous polymers could also affect the mechanical properties by the plasticising action of the monomer. It can be postulated that any unreacted monomer present only at the pore walls, where the monomer is exposed to oxygen during the polymerisation process. The oxygen-exposed surface will remain “tacky” as oxygen inhibits the polymerisation the surface in contact with air. However, the macroporous polymers were washed with ethanol to remove any unreacted monomer prior to mechanical testing. In addition to this, the mechanical properties of porous materials are governed predominantly by cellular morphology and the properties of the base polymer. Therefore, any remaining unreacted AESO should not have affected the mechanical properties measured in this study.

Conclusions

Studies on the mechanical frothing of liquid monomers to produce non-aqueous foam templates followed by their polymerisation to produce polymeric foams had been conducted in the 1970s. However, this technique was limited to the production of 2D foams, which might be due to the instability of the air-monomer foam. In this study however, we succeeded in producing macroporous polymers through microwave heating of mechanically frothed gas-AESO liquid foams containing lauroyl peroxide as thermal initiator due to the high viscosity of the monomer phase and the obstruction of the flow into the Plateau border in the presence of BC during capillary drainage of the monomer liquid foams. The incorporation of 0.5 wt% BC into the polymeric foams resulted in a significant increase of the specific Young's modulus when compared to the neat polymer foams. This was attributed to the reinforcing effect of the highly crystalline and stiff BC. However, a further increase of BC to 1 wt% resulted in a decrease of the specific Young’s modulus when compared to the neat polymeric foams. This is thought to be due to the ill-defined cell structure of the macroporous polymer nanocomposites at 1 wt% BC loading. It was hypothesised that the poor compatibility between BC and polymerised AESO has resulted in the decrease of the compression strength of the polyAESO with increasing BC content.

Experimental

Materials

Epoxidised acylated soybean oil, AESO (Aldrich, density = 1.04 g cm⁻³, inhibited with 8300 ppm monomethyl ether hydroquinone), was purchased from Sigma-Aldrich and used as the monomer in this study. Lauroyl peroxide, also known as dodecanoyl peroxide (Alfa Aesar, purity ≥ 97%) and ethanol (GPR, purity ≥ 99.7%) were purchased from VWR International UK. Lauroyl peroxide was selected as the thermal initiator for this study because of its low initiation temperature (10 h half-life at 65 °C). De-ionised water was used throughout this study and was obtained by passing water through a reverse osmosis water filtration unit (Triple Red Laboratory Technology, Bucks, UK). BC nanofibrils were extracted from commercially available nata-de-coco (CHAOKOH coconut gel in syrup, Ampol Food Processing Ltd, Nakorn Pathom, Thailand) following previously described work and used in its freeze-dried form in this work.

Macroporous polymer nanocomposites preparation

Polymer foams of neat AESO (polyAESO 1) were prepared by polymerising the gas-AESO liquid foams (AESO 1) using lauroyl peroxide as the radical initiator. Firstly, 1 wt% (2.5 mol-%) of lauroyl peroxide (relative to the monomer phase) was mixed with AESO and air was introduced into the monomer by using a hand mixer operating at its maximum power output (100 W) for 5 min. The resulting gas-AESO liquid foam (AESO 1) was shaped into cylindrical glass vials (25 mm in diameter and 60 mm in height) using spatula and polymerised for 2 min using microwave irradiation, operating at a frequency 2.45 GHz with 180 W power output. The polymer foams were allowed to undergo a slow cooling process at room temperature followed by a slow drying process at 60 °C for 3 days. The resulting foam was then removed from the vial and was microwaved for 2 min at 1.5 W using a microwave oven (Ito, Inaba Co., Japan) with a plate size of 170 mm × 170 mm × 3 mm containing a distance of 150 mm between the microwaves and the foams. The resulting foam was then removed from the vial and was microwaved for 2 min at 1.5 W using a microwave oven (Ito, Inaba Co., Japan) with a plate size of 170 mm × 170 mm × 3 mm containing a distance of 150 mm between the microwaves and the foams.

Table 2 Mechanical properties of the macroporous polymer. E, σ, E' and σ' denote Young’s modulus in compression, compression strength, specific Young's modulus and specific compression strength, respectively. The errors tabulated are the standard errors

<table>
<thead>
<tr>
<th>Sample</th>
<th>E (MPa)</th>
<th>σ (kPa)</th>
<th>E' (MPa kg⁻¹ m⁻³)</th>
<th>σ' (kPa kg⁻¹ m⁻³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>polyAESO 1 (0 wt% BC)</td>
<td>111 ± 11</td>
<td>343 ± 34</td>
<td>253 ± 25</td>
<td>779 ± 79</td>
</tr>
<tr>
<td>polyAESO 2 (0.5 wt% BC)</td>
<td>166 ± 33</td>
<td>254 ± 19</td>
<td>338 ± 68</td>
<td>519 ± 39</td>
</tr>
<tr>
<td>polyAESO 3 (1 wt% BC)</td>
<td>52 ± 9</td>
<td>183 ± 26</td>
<td>90 ± 15</td>
<td>315 ± 46</td>
</tr>
</tbody>
</table>

It was observed that the compression strength of the polyAESO decreased with increasing BC content (Table 2). The compression strength of the polymer nanocomposite foams decreased by as much as 60% from 777 kPa kg⁻¹ m⁻³ to 315 kPa kg⁻¹ m⁻³ when the BC content was increased to 1 wt%. It is proposed that the poor specific strength of the nanocomposites is due to the poor compatibility between polyAESO and BC and hence, poor stress transfer between BC and the polymerised AESO.

The presence of unreacted AESO in the macroporous polymers could also affect the mechanical properties by the plasticising action of the monomer. It can be postulated that any unreacted monomer present only at the pore walls, where the monomer is exposed to oxygen during the polymerisation process. The oxygen-exposed surface will remain “tacky” as oxygen inhibits the polymerisation the surface in contact with air. However, the macroporous polymers were washed with ethanol to remove any unreacted monomer prior to mechanical testing. In addition to this, the mechanical properties of porous materials are governed predominantly by cellular morphology and the properties of the base polymer. Therefore, any remaining unreacted AESO should not have affected the mechanical properties measured in this study.

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Studies on the mechanical frothing of liquid monomers to produce non-aqueous foam templates followed by their polymerisation to produce polymeric foams had been conducted in the 1970s. However, this technique was limited to the production of 2D foams, which might be due to the instability of the air-monomer foam. In this study however, we succeeded in producing macroporous polymers through microwave heating of mechanically frothed gas-AESO liquid foams containing lauroyl peroxide as thermal initiator due to the high viscosity of the monomer phase and the obstruction of the flow into the Plateau border by the addition of BC. Through this route, we were able to produce 3D macroporous polymers. The resulting bio-based macroporous polymers possessed a porosity of approximately 60% but it was found that the stability of the gas-AESO liquid foam template was poor in comparison to the air/AESO foams containing BC. This liquid foam destabilises within 7 days of preparation at room temperature. In order to enhance the stability of the gas-AESO liquid foams and the mechanical properties of the macroporous polymers, BC was added as a nanofiller to produce macroporous polymer nanocomposites. With the introduction of BC, the stability of the liquid AESO foams was significantly enhanced. No phase separation (e.g. creaming) was observed even 70 days after the preparation of the liquid foams. This was thought to be due to the obstruction of the Plateau border in the presence of BC during capillary drainage of the monomer liquid foams. The incorporation of 0.5 wt% BC into the polymeric foams resulted in a significant increase of the specific Young's modulus when compared to the neat polymer foams. This was attributed to the reinforcing effect of the highly crystalline and stiff BC. However, a further increase of BC to 1 wt% resulted in a decrease of the specific Young’s modulus when compared to the neat polymeric foams. This is thought to be due to the ill-defined cell structure of the macroporous polymer nanocomposites at 1 wt% BC loading. It was hypothesised that the poor compatibility between BC and polymerised AESO has resulted in the decrease of the compression strength of the polyAESO with increasing BC content.

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and power output of 2.45 GHz and 700 W, respectively. After polymerisation of the gas-AESO foams, the samples were left to cool to room temperature before washing them with ethanol to remove unreacted monomer, followed by de-ionised water. The polymeric foams were then dried at 40 °C in an air oven overnight. Gas-AESO liquid foams with 0.5 wt% BC (AESO 2) and 1 wt% BC (AESO 3) were prepared by mixing BC and 1 wt% of lauroyl peroxide (relative to AESO) followed by the introduction of air into the BC-monomer dispersion using the hand mixer as previously described. The gas-AESO-BC liquid foams (AESO 2 and 3) were then polymerised and washed following the previously described methods. The macroporous polymer nanocomposites of AESO 2 and 3 are termed polyAESO 2 and polyAESO 3, respectively.

Characterisation of the gas-AESO liquid foams and macroporous polymer nanocomposites

Stability indices of gas-AESO liquid foams. The stability indices of gas-AESO liquid foams were assessed by monitoring the upward movement of the air-in-AESO boundary as a result of creaming at 20 °C. The movement of the air bubbles in AESO boundary was visually monitored every 24 h. The stability index was calculated by taking the ratio between the creaming heights at the time of assessment and the initial height of the gas-AESO liquid foams.

Structure and morphology of the macroporous polymer nanocomposites. The internal structure and morphology of the macroporous polymers and nanocomposites were characterised using variable pressure scanning electron microscopy (SEM) (JSM 5610 LV, Jeol Ltd, Herts, UK) using an accelerating voltage of 10 kV. The macroporous polymer nanocomposites were cut using a scalpel into approximately 0.5 cm³ pieces and stuck onto aluminium SEM stubs using carbon tabs. Prior to SEM, the samples were Au coated (K550 sputter coater, Eimettech Ltd, Ashford, Kent, UK) for 2 min at 20 mA.

Density and porosity of the macroporous polymer nanocomposites. He pycnometry (AccuPyc 1330, Micromeritics Ltd, Dunstable, UK) was used to measure the matrix (true) density of the polymer. Prior to the measurements, the polymer foams were crushed into powder using a pestle and mortar. The envelope (foam) density of the cylindrical macroporous polymers was calculated by taking the ratio between the mass and the volume of the macroporous polymer, which was determined from the diameter and the height of the material. The porosities of the macroporous polymer nanocomposites were calculated as follows:

\[ P = \left(1 - \frac{\rho_p}{\rho_n}\right) \times 100\% \]  

where \( P \), \( \rho_p \), and \( \rho_n \) are the porosity of the macroporous polymer, envelope (foam) density and matrix (true) density, respectively. A total of 5 specimens were measured for each type of sample.

Thermal gravimetric analysis (TGA) of the macroporous polymer nanocomposites. The degradation behaviour of the macroporous polymers and nanocomposites was characterised using TGA (TGA Q500, TA Instruments, UK). A piece of a sample with an approximate mass of 20 mg was heated from room temperature to 800 °C at a heating rate of 10 °C min⁻¹ in nitrogen atmosphere.

Compression properties of the macroporous polymer nanocomposites. The macroporous polymer nanocomposites were tested in compression using an Instron universal material testing machine (Instron 4505, Instron Corporation, MA, USA) equipped with a 1 kN load cell. The test was conducted in accordance to BS ISO 844 : 2009. The cylindrical test specimens had the same diameter and height of 25 mm. Strain gauges (FLA-1-11, Techni Measure, Studley, UK) were glued onto the middle portion of the test specimens using cyanoacrylate glue (EVERBUILD Building Products Ltd, Leeds, UK). In order to avoid the glue filling the pores, only a very thin layer of glue was applied on the strain gauges. The samples were compressed between two flat and parallel thick polished plates coated with Teflon. Specimens were tested at a crosshead speed of 1 mm min⁻¹. A total of 5 specimens were tested for each type of sample. The errors tabulated are standard deviations.

Acknowledgements

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22 A. J. Brown, 
23 W. Czaja, D. Romanovicz and R. M. Brown, 
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