Research Article

Controlling Foam Morphology of Poly(methyl methacrylate) via Surface Chemistry and Concentration of Silica Nanoparticles and Supercritical Carbon Dioxide Process Parameters

Deniz Rende, Linda S. Schadler, and Rahmi Ozisik

Department of Materials Science and Engineering and Rensselaer Nanotechnology Center, Rensselaer Polytechnic Institute, Troy, NY 12180, USA

Correspondence should be addressed to Rahmi Ozisik; ozisik@rpi.edu

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Polymer nanocomposite foams have received considerable attention because of their potential use in advanced applications such as bone scaffolds, food packaging, and transportation materials due to their low density and enhanced mechanical, thermal, and electrical properties compared to traditional polymer foams. In this study, silica nanofillers were used as nucleating agents and supercritical carbon dioxide as the foaming agent. The use of nanofillers provides an interface upon which CO$_2$ nucleates and leads to remarkably low average cell sizes while improving cell density (number of cells per unit volume). In this study, the effect of concentration, the extent of surface modification of silica nanofillers with CO$_2$-philic chemical groups, and supercritical carbon dioxide process conditions on the foam morphology of poly(methyl methacrylate), PMMA, were systematically investigated to shed light on the relative importance of material and process parameters. The silica nanoparticles were chemically modified with tridecafluoro-1,1,2,2-tetrahydrooctyl triethoxysilane leading to three different surface chemistries. The silica concentration was varied from 0.85 to 3.2% (by weight). The supercritical CO$_2$ foaming was performed at four different temperatures (40, 65, 75, and 85°C) and between 8.97 and 17.93 MPa. By altering the surface chemistry of the silica nanofiller and manipulating the process conditions, the average cell diameter was decreased from 9.62±5.22 to 1.06±0.32 μm, whereas, the cell density was increased from 7.5±0.5×10$^8$ to 4.8±0.3×10$^{11}$ cells/cm$^3$. Our findings indicate that surface modification of silica nanoparticles with CO$_2$-philic surfactants has the strongest effect on foam morphology.

1. Introduction

In nature, foams are found in the form of bone, natural sponge, coral, and natural cork. Inspired by these materials, processing of polymer foams has received considerable attention [1, 2]. Polymer nanocomposite foams are micro- and nanoporous materials that are used as thermal and sound barriers, shock absorbers, absorbents, cushions, and tissue engineering scaffolds [3, 4]. Recent advances in processing of polymer nanocomposite foams enabled new application areas in hydrogen storage [5], electromagnetic shielding [6], and sensing technology [7]. Various polymers are being used in foam applications such as polyurethane, polystyrene, polyethylene, polypropylene, poly(vinyl chloride), polycarbonate, and poly(methyl methacrylate) [8–14] as well as specialty polymers.

Foams are divided into various categories depending on their pore morphology (open versus closed) or density (low versus high). Open cell morphology consists of pores (bubbles) that are connected to each other, making the material softer and more absorbent. In the closed cell morphology, the pores are isolated from each other, which makes the foam more rigid. In addition to these categories, polymer foams can also be characterized according to their density, cell size, cell density, and wall thickness, all of which influence the properties of the foam. In general, polymer foams have low thermal conductivity, poor mechanical properties, and poor surface quality due to the underlying porous structure. However, their low density, low thermal conductivity, and sound barrier properties make them highly attractive for a variety of applications. For example, low density foams are primarily used in packaging and insulation applications,
and high density foams are used in structural applications [3, 15] as thermal or sound barriers. Polymer foams can also be categorized by their flexibility: flexible, semiflexible, and rigid. Flexible and semiflexible polymer foams are used to produce cushions, textiles, toy parts, and sporting goods. On the other hand, rigid polymer foams have applications in insulation, construction, durable goods, and infrastructure.

In general, the foam morphology is formed via the use of foaming agents, which undergo phase transition either due to physical or chemical changes creating a gas phase that expands forming gas bubbles inside the polymer matrix. The nucleation of the bubbles occurs via two different classical mechanisms: homogeneous or heterogeneous nucleation. In the homogeneous nucleation case, concurrent initiation and growth of bubbles are observed leading to a wide cell size distribution in the final foam structure. The heterogeneous nucleation, however, requires the existence of a secondary material that promotes simultaneous growth of bubbles inside the polymer matrix, resulting in a narrow cell size distribution. The addition of inorganic nanoparticles, which act as nucleating agents, induces heterogeneous nucleation and provides a large number of nucleation sites. Furthermore, the presence of micro- or nanosized fillers dramatically decreases the energy barrier for cell nucleation compared to that required for homogeneous nucleation [9, 13, 16, 17]. Existing models based on classical nucleation theories sometimes fail to explain the nucleation satisfactorily. Other models, for example, consider microvoids as nucleation sites. Microvoids lead to stress induced bubble formation, and the population of sufficiently large viable microvoids that overcomes surface and elastic forces can then be related to the cell density [18–20].

The synthesis of nanofillers is of recent interest because they provide a high density of nucleation sites at low concentrations. The highest nucleation efficiency is achieved when nucleation on the filler surface is energetically favorable and the filler is uniformly distributed and dispersed within the polymer matrix. Therefore, heterogeneous nucleation conditions could be controlled to some extent via filler type, geometry (size, aspect ratio, etc.), and surface chemistry [1, 21, 22]. Selection of the filler type is an important step to achieve the final desired properties. For instance, carbon based nanofillers (carbon nanotubes and graphene) enhance the strength and conductivity [8, 23]. The flat surfaces of clays offer an excellent surface for nucleation; however only when dispersion is controlled, uniform cell size and remarkable mechanical properties are achieved [24–29]. Spherical nanofillers are another class of fillers that are being used to create polymeric foams [21, 30], polymer blends [31], and copolymers [32]. Among spherical nanoparticles, silica nanoparticles have been widely used as bare or after surface modification with vinyl or fluorinated silane coupling agents to foam polymers. Studies using bare silica nanoparticles showed a reduction in the average cell size with increasing concentration [33]. The surface modification of silica nanoparticles was reported to lead to enhanced interaction between the foaming agent and the nanofiller, thereby reducing the average cell size further [34]. For example, Goren et al. reported an 8-fold decrease in average cell size after modifying silica nanoparticles with fluorinated silane coupling agents [21]. However, spherical nanoparticles do not improve properties as much as nanosized rods or sheets.

In addition to filler characteristics, foaming conditions also influence the final foam structure. Conventional foamed products can be produced either by chemical or physical blowing agents. Chemical blowing agents are mixed into polymer matrix and decompose when heated up yielding a gas release. This process requires an additional step to eliminate the residual chemical blowing agent [35]. Due to the challenges in the removal of side products, physical foaming is often preferred over chemical foaming. Physical foaming involves the saturation of the polymer matrix with a gas at high pressure followed by a rapid decrease in pressure. This rapid decrease in pressure leads to the expansion of the gas and, therefore, results in the formation of bubbles within the polymer matrix. The lack of hazardous chemical solvent use during physical foaming makes this technique the preferred method in the production of polymeric foams. Supercritical fluids have also become an attractive option for the production of polymer foams. Due to stable, nontoxic, inflammable, low cost and its easily attainable critical conditions, supercritical carbon dioxide has been widely used to create polymer foams [14, 21–23, 29]. Supercritical CO₂ exhibits relatively high solubility in polymers, acts as a plasticizer by lowering the glass transition temperature, easily diffuses into the polymer, reduces the viscosity and surface tension of the polymer melt, and assists in polymer processing. Foaming with supercritical CO₂ can be performed either as a batch process, where the samples are kept in a pressure chamber and saturated with supercritical carbon dioxide [14, 21, 36], or as a continuous process such as the inside of an extruder [37–39]. After the polymer is saturated with supercritical CO₂, the rapid decrease in pressure generates a thermodynamic instability, which leads to phase transition from supercritical to gaseous state and causes the nucleation of gas bubbles. Saturation (CO₂ soaking) pressure, temperature, time, and depressurization rate are critical parameters in determining the final morphology of the polymer foam [40]. At high pressures, more carbon dioxide is absorbed into the polymer, leading to an increase in cell density [41]. In addition, high pressure enhances the driving force for cell growth.

Many researchers investigated the silica/PMMA nanocomposite systems foamed with supercritical CO₂ and a review article by Chen and coworkers [42] discusses many of the recent developments. Siripurapu et al. [33] studied the effect of concentration of untreated (bare) silica nanoparticles at 1, 5, and 8% silica concentrations. The foamed material was produced with CO₂ at 40°C and 6.83 MPa. An increase in nanosilica concentration led to reduced average cell diameter and increased cell density under isothermal and isobaric conditions. Yeh et al. [34] highlighted the importance of surface modification of silica nanoparticles, which might influence the interaction of filler with CO₂. For example, at various concentrations untreated and vinyl-modified silica nanoparticles containing PMMA composites were foamed at 150°C and 13.8 MPa. The cell sizes achieved in this process ranged from 10 to 23 μm. The surface
modification of silica nanoparticles was also performed with fluorinated silane coupling agents by our group [21], in which the silica nanoparticle sizes ranged from 60 to 150 nm. PMMA nanocomposites having 1% silica with 66, 80, 149, and 157 nm silica nanoparticles were foamed under supercritical CO₂ at 40°C and 17.93 MPa. In this study, the particle size was shown to have an important contribution to cell density and fluorination of the silica nanoparticles was shown to facilitate the dispersion of the silica nanoparticles inside the PMMA matrix.

Although nanofiller concentration and surface chemistry are known to have strong influence on the final cell size and cell density [14, 21, 43, 44], the extent of surface coverage (tethering density) of the functional groups on the foam morphology is yet to be reported in a systematic manner. In the current study, we aim to investigate the effect of tethering density of fluoroalkanes on the foam morphology of silica/PMMA nanocomposites. Fluorinated surface modifiers were chosen because they were shown to exhibit CO₂-phlic behavior [21]. The effects of silica concentration, foaming temperature, and saturation pressure on the final foam morphology were also investigated.

2. Experimental

2.1. Materials. To synthesize silica nanoparticles, tetraethyl orthosilicate (TEOS, 98% reagent grade, Sigma, 131903) and ammonia solution (NH₃, 28%, Sigma, 338818) were used as received. Tetrahydrofuran (THF, Fisher Scientific, T397-1) was used to purify the silica nanoparticles. The surface modification of the silica nanoparticles was performed with silane coupling agent tridecafluoro-1,1,2,2-tetrahydroxytriethoxysilane (F-TEOS, Gelest, SIT8175). Poly(methyl methacrylate), PMMA, was chosen as the matrix polymer because of its outstanding chemophysical properties [45] and relatively high affinity for CO₂. Commercial grade PMMA (Plexiglas V920-100) was donated by Altuglas International.

2.2. Preparation of Silica Nanoparticles. The method used in the synthesis of silica nanoparticles is an adaptation of the Stöber’s procedure [46]. To prepare 100 nm bare silica nanoparticles, 87.5 g ethanol, 4.37 g distilled water, and 4.12 g ammonia solution was mixed with 8.62 g of TEOS. The reaction mixture was stirred overnight at room temperature and unreacted solvents were removed by rotary evaporator at 60°C and 90 rpm for 2 hours. The sample was then kept in vacuum dryer overnight at 60°C. In order to remove any organic and inorganic impurities and unused reactants, the dried sample was first washed with THF and vacuum-filtered through 0.45 μm hydrophobic filtering paper, then washed with distilled water, and vacuum-filtered through 0.45 μm hydrophilic filtering paper. Finally, the sample was dried again in vacuum oven overnight at 60°C. This procedure approximately yields 1.5 g of bare silica nanoparticles with an average size of ~100 nm.

2.3. Surface Modification of Silica Nanoparticles. In order to increase the interaction of silica and CO₂, silica nanoparticles were modified by tethering fluoroalkane chains (F-TEOS) onto their surfaces. To prepare the surface modified nanoparticles, silica nanoparticles were synthesized as described before and each batch was divided into multiple parts; one was left as is (bare silica) and the other parts were modified with F-TEOS. This ensured that the starting silica nanoparticles on average had the same size. Two different amounts (0.1 and 0.7 g) of F-TEOS were added to two different silica batches to create two different surface tethering densities. The surface modification reaction took 24 hr at room temperature. Unreacted solvent was removed by rotary evaporator at 60°C and 90 rpm for 2 hours, and the samples were dried in a vacuum oven overnight at 60°C. To remove any residual solvent and impurities, each sample was subsequently washed with THF and water, filtered, and dried in vacuum oven overnight at 60°C.

2.4. Structural Analysis of Silica Nanoparticles. The size of the bare and surface modified nanoparticles was characterized by field emission scanning electron microscopy (SEM), transmission electron microscopy (TEM), and image processing. After the synthesis, the bare and surface modified silica nanoparticles were dried and the nanoparticles were first subjected to gold sputtering to form a 10–15 nm layer of gold on the sample surface. SEM images were obtained with a JEOL JSM-6332 using an accelerating voltage of 10 kV. TEM images were collected with a Philips CM12 with an accelerating voltage of 120 kV. Subsequently, both SEM and TEM images were analyzed with ImageJ [47] and diameter measurements of at least 100 nanoparticles were recorded in order to have statistically significant results.

The extent of the surface modification was determined by thermogravimetric analysis (TGA). Each silica nanoparticle sample was analyzed by TA Instruments TGA Q50 by heating up to 900°C at a rate of 20°C/min. The resulting percent weight changes between 200 and 800°C were used to calculate the percent coverage of the silica surface. Both bare and surface fluorinated samples were tested for at least three times.

The surface area of the silica nanoparticles was measured with Quantachrome BET Surface Analyzer Autosorb. Before BET analysis, the samples were degassed at 150°C under nitrogen atmosphere for 24 hrs.

2.5. Preparation of Polymer Nanocomposites. The polymer nanocomposite samples were prepared by melt mixing with a benchtop twin-screw extruder (Haake MiniLab). The extruder was equilibrated for 2 hrs at 220°C before each use. The rotation speed of the screws was set to 60 rpm. Before each operation, 5 g of neat PMMA was passed through the system to clean the screws. The samples with different concentrations of bare and surface modified silica nanoparticles were mixed with PMMA and each sample was cycled within the extruder for 4 min and flushed in 3 min. These parameters were chosen because they were previously shown to disperse and distribute silica nanoparticles effectively [21]. Neat PMMA samples were also processed in the same way as silica containing PMMA as control samples.
Although silica was added to PMMA in premeasured amounts, due to the chaotic nature of the twin-screw extruder used, silica concentration in silica/PMMA composites was subsequently measured again by thermogravimetric analysis (TA Instruments). For statistical accuracy, three different samples, each being approximately 40 mg, were analyzed with a TA Instruments TGA Q50 by heating to 900°C at a rate of 20°C/min. The silica concentration was determined from the remaining weight at 800°C, after which the weight remains constant. The melt mixing process led to several composite samples each having slightly different amount of bare or surface modified silica nanoparticles. In order to establish statistical significance, these samples were separated into two groups of “low” and “high” silica concentrations. The low silica concentration group contained 0.85–1.38% silica and the high concentration group contained 2.5–3.2% silica. All silica concentrations are reported as weight percentages. Another 2.9% silica/PMMA nanocomposite sample was prepared with highly surface modified silica nanoparticles. This sample was used specifically to investigate the effect of supercritical CO$_2$ process parameters on foam morphology while eliminating sample variations and effect of surface modification. The nanocomposite samples were labeled according to their silica surface tethering density (B: bare; F: fluorinated with low tethering density; and FF: fluorinated with high tethering density) and silica concentration (low or high). For instance, B-Low indicates a nanocomposite sample containing bare silica nanoparticles at low concentration (0.85%). The neat PMMA control sample was labeled as “PMMA.” This labeling convention was necessitated by the fact that during sample preparation neither the surface tethering density nor the silica concentration could be controlled accurately.

### 2.6. Foaming with Supercritical Carbon Dioxide

A batch foaming process was performed to prepare polymer nanocomposite foams using supercritical CO$_2$ as the foaming agent. A high-pressure reactor (Parr, 5512, 50 mL) was connected to a Teledyne ISCO high-pressure syringe pump. The polymer nanocomposite extrudates were cut into 1-2 cm long pieces and were placed in the pressure chamber. The samples were then saturated with supercritical CO$_2$ and were kept at predetermined temperature and pressure for 24 hrs (the exact processing conditions are provided in Section 3). After saturation with supercritical CO$_2$ for 24 hours, the pressure was decreased at a rate of 1.5 MPa/s. The samples were removed from the pressure chamber and were soaked in water that was maintained at 60–65°C for one minute.

### 2.7. Determination of Foam Morphology and Cell Size Distribution

The foam morphology was investigated using field emission scanning electron microscope (SEM, JEOL JSM-6332). Samples were freeze-fractured in liquid nitrogen and the fracture surfaces were sputter-coated with 10–15 nm of gold. The images were then collected under 15 kV accelerating voltage. SEM images of the fracture surfaces, both from the center and near the perimeter, were recorded with SEM for image processing with ImageJ [47].

Figure 1 shows the size distribution of bare and surface modified silica nanoparticles. Since the surface modified nanoparticles were prepared from the same batch as the bare silica nanoparticles, only one size distribution graph is presented. The average diameters were found to be 101.2 ± 0.8 nm for bare silica nanoparticles and 112.4 ± 1.3 nm for modified silica nanoparticles.

### 3. Results and Discussion

#### 3.1. Nanoparticle Size

The size of the silica nanoparticles was determined by image analysis of transmission electron micrographs (TEM). Figure 1 shows the size distribution of bare and surface modified silica nanoparticles. Since the surface modified nanoparticles were prepared from the same batch as the bare silica nanoparticles, only one size distribution graph is presented. The average diameters were found to be 101.2 ± 0.8 nm for bare silica nanoparticles and 112.4 ± 1.3 nm for modified silica nanoparticles.

#### 3.2. Tethering Density

As explained in Section 2, in order to ensure that the surface modified nanoparticles had the same starting size as bare silica nanoparticles, after the synthesis of the nanoparticles, the batch was divided into two groups and 0.1 and 0.7 g F-TEOS were added to prepare low (F) and high (FF) fluorinated surfaces. According to the molecular structure of the surface modifier, F-TEOS (Figure 2(a)), the reaction yields ethanol (Figure 2(b)). Fluorinated alkylsilsesquioxane derivatives react with the surface hydroxyl groups on silica surface and form self-assembled monolayers or multilayers [48]. The triethoxysilane group can be attached to the surface in three different configurations (Figure 2(b)). If the silane coupling agent is attached to the surface via one or two silane groups (as shown by mechanisms I and II in Figure 2(b)), the remaining silane group(s) may react with neighboring tethered or free F-TEOS molecules resulting in multiple layers on the surface.
A quantitative analysis of the F-TEOS coverage on the silica nanoparticle surfaces was performed by thermogravimetric analysis (Figure 3). Using untreated silica nanoparticle weight change ($\Delta w_{B-SiO_2}$) between 200 and 800°C as the reference, the following calculation provided the percent surface coverage of the silica nanoparticles [49]:

$$n_{\text{silane}} = \frac{\Delta w_{F-SiO_2} - \Delta w_{B-SiO_2}}{100 \cdot M_{\text{mod}} \cdot \sigma}$$

where $n_{\text{silane}}$ is the number of moles of silane, $\Delta w_{F-SiO_2}$ is the percent weight change of fluorinated silica nanoparticles (between 200 and 800°C), $M_{\text{mod}}$ is the molecular mass that is lost (as volatile gas) between 200 and 800°C per mole of used silane, and $\sigma$ is the specific surface area of the silica nanoparticles and was measured with BET surface analysis, which yielded specific surface areas of 32.6 m$^2$/g for bare nanoparticles and 33.9 m$^2$/g for the surface modified nanoparticles. For reaction mechanisms shown in Figure 2(b), the molecular weights of the remaining silane group on the silica ($M_{\text{mod}}$) were 481.36, 452.36, and 423.36 g/mol corresponding to single, double, and triple ethoxysilane attachments. In order to assess surface coverage, $n_{\text{silane}}$ obtained from (1) needs to be compared to $n_{\text{hydroxyl}}$. According to Zhuravlev [50], silica nanoparticles with an average size of 100 nm contain 5.85 hydroxyl groups per square nanometer; therefore, if (1) yields over 5.85 silane groups (on 100 nm sized silica), the surface coverage would be over 100%, which can be explained only by the presence of a multilayer silane coverage as suggested by reaction mechanisms I and II in Figure 2(b).

Table 1 presents surface coverages calculated via (1) for each possible reaction mechanism presented in Figure 2(b). This calculation showed that low fluorinated silica nanoparticles (F) have a surface coverage around 73%, whereas this value exceeds 100% for the highly fluorinated silica nanoparticles (FF) indicating multilayer formation.

The silica nanoparticles prepared in the current study can be summarized as follows. (i) The average size of silica nanoparticles prepared in the current study can be summarized as follows.
nanoparticles are 101.2 ± 0.8 nm for bare (B) silica and 112.4 ± 1.3 nm for fluorinated (F and FF) silica. (ii) The fluorinated silica nanoparticles have an average surface coverage of ∼73% for F and ∼120% for FF indicating multilayer fluorination at the surface of these samples.

3.3. Silica Concentration in Polymer Nanocomposites. The silica concentration of the samples was determined by thermogravimetric analysis and the results are presented in Figure 4. The final silica concentrations were grouped into “low” and “high” concentrations containing 0.85–1.38% and 2.5–3.30% silica, respectively. Recalling that the surface tethering densities are represented with “B” for bare silica, “F” for low tethering density, and “FF” for high tethering density, each silica/PMMA is labeled with both of these markers, such as B-Low or B-High, in the following sections.

3.4. Effect of Silica Surface Modification and Concentration on Foam Morphology. Polymer nanocomposite samples were processed with supercritical CO₂ and the resulting foam morphologies obtained via scanning electron microscopy (SEM) are presented in Figure 5. Visual observation of the SEM images clearly shows the difference between the foam morphologies of nanocomposites containing bare and fluorinated silica nanoparticles. At low and high concentrations of bare silica nanoparticles, the foam morphology (average cell size, cell density) looks similar to that of neat PMMA even though the nucleation mechanism between these two types of samples are completely different: homogeneous in neat PMMA versus heterogeneous in filled PMMA. The similarity of the foam morphologies is mainly attributed to the poor interaction of CO₂ with bare silica and also to the formation of silica agglomerates inside the polymer matrix (micrograph not shown). On the other hand, all four samples containing fluorinated silica nanoparticles (F-Low, F-High, FF-Low, and FF-High) have remarkably low (average) cell sizes and high cell densities. In order to quantify these visual observations, the foamed samples’ cell density and cell size distributions were characterized by image processing.

Cell density calculations were based on a method described by Kumar and Suh [51]. Typically, a micrograph showing more than 100 bubbles (cells) was chosen for the analysis. The number of bubbles (n) was counted and the number of bubbles per unit volume (N) was calculated using the following equation:

\[ N = \left( \frac{nM^2}{A} \right)^{3/2}, \]

where \( M \) is the magnification factor and \( A \) is the true area of the SEM image. According to this method, the cell density of the samples was calculated from five different images collected from the center and the edges of the fracture surface.

The cell density measurements confirmed that neat PMMA, and the polymer nanocomposite samples prepared with bare silica nanoparticles have very low cell densities compared to the foamed samples containing fluorinated silica nanoparticles regardless of the tethering density and silica concentration (Figure 6(a)). The neat PMMA foam has an average cell density of 7.5 ± 0.5 \( \times 10^8 \) cells/cm\(^3\). The foamed samples containing bare silica nanoparticles formed agglomerates (micrograph not shown), which evidently affected the cell density. A remarkable difference is observed in cell densities between the samples containing bare and fluorinated silica containing PMMA. This is because surface chemistry affects both the dispersion and distribution of silica nanoparticles in the PMMA matrix and the interaction between silica nanoparticles and CO₂ molecules. It is well known that supercritical CO₂ exhibits high affinity for fluorocarbons, fluoroethers, siloxanes, and polycarbonates [52]. Fluorination of silica decreases the surface energy and leads to a decreased critical activation energy required for nucleation of CO₂ bubbles [21]. On the other hand, among the fluorinated samples, the highest cell density was achieved for high concentration samples (FF) indicating that the silica concentration is the determining factor.

The immediate change in cell density upon surface modification (going from B to F) is quite impressive showing almost a 100-fold increase. This suggests that there is a tremendous effect of surface chemistry on foam morphology. On the other hand, increasing surface tethering density from ∼73% to over 100% did not change the cell density values significantly. It is, therefore, important to have the proper surface chemistry but having too much of CO₂-philic chemical groups on the nanofiller surface does not further improve cell density values. This result can be explained by the selective placement of CO₂ molecules at the filler/polymer interface. Even a modest selectivity achieved through surface modification leads to improved results. Unfortunately tethering densities lower than ∼70% were not available in the current study; therefore, it is not possible to state if there is
Increasing tethering density
Bare Fluorinated
LowHigh
Increasing concentration

Figure 5: SEM images of the foamed samples with varying surface modification and concentration.

Figure 6: (a) Cell density and (b) average cell size of the neat PMMA, nanocomposites containing bare nanosilica (B), nanocomposites with surface treated silica at low fluorination (F) and high fluorination (FF). Error bars represent standard error of five samples and error bars smaller than symbol size are not shown.
an optimum tethering density that would lead to the best results (high cell density, small average pore size).

The cell sizes of the samples were measured from SEM images. The results of the average cell diameters are presented in Figure 6(b). The average cell diameter of the neat PMMA polymer foam was calculated to be 9.62 ± 5.22 μm. The cell size measurements of the samples containing bare silica nanoparticles showed that these samples have comparably lower average cell diameters. However, the high standard deviation in neat PMMA and nanocomposites containing bare silica nanoparticles indicate the presence of a wide distribution of cell sizes in these samples. This similarity in the cell size distribution between foamed PMMA and B samples is attributed to the fact that all of these samples had homogeneous nucleation. Bare silica containing samples additionally had heterogeneous nucleation; however this secondary mechanism must have been quite ineffective due to the similarities in the foam morphologies between PMMA and B samples. Previous studies reported that homogeneous nucleation leads to a wide cell size distribution whereas heterogeneous nucleation leads to a narrow cell size distribution [13]. Obviously, the bare silica containing samples’ cell size distribution is similar to that of neat PMMA because both systems undergo homogeneous nucleation, but the average cell size in bare silica containing samples is relatively lower than that of neat PMMA because these samples also experience heterogeneous nucleation. Contrary to neat PMMA and bare silica containing PMMA (B samples), the surface fluorinated silica nanoparticle containing samples (F and FF) have significantly lower average cell sizes and smaller standard deviations. The smallest average cell sizes were achieved at higher concentrations of silica (“high” samples). Surface tethering density had a minor effect on the results. These observations are in agreement with the results obtained for cell densities (Figure 6(a)).

3.5. Effect of Processing Conditions on Foam Morphology. During supercritical CO$_2$ saturation, CO$_2$ also serves as a plasticizer for PMMA [53–55] and it was shown that CO$_2$ has greater solubility in polymers with branched structures compared to linear polymers. In the current study, carbon dioxide has comparably high solubility in PMMA due to specific interactions it has with the carbonyl groups of PMMA. The higher affinity between CO$_2$ and carbonyl groups of PMMA implies higher plasticization capacity, which is manifested in lowered glass transition temperature and viscosity [56]. The solubility of CO$_2$ in PMMA was reported to be about 6.3% (by weight) at 20°C and 6.89 MPa, 9.3% at 40°C and 13.78 MPa, and 10.5% at 70°C and 20.67 MPa [57]. Therefore, because the solubility of CO$_2$ in PMMA changes as a function of processing parameters (pressure and temperature), it is necessary to also consider the effect of the processing parameters on the final foam morphology.

In a typical experiment, the polymer nanocomposite sample was saturated with supercritical CO$_2$ at a preselected temperature and pressure. In order to understand the effect of processing conditions on foam morphology, several experiments were conducted at 65 and 85°C and at 8.97, 13.45, and 17.93 MPa. For this study only one sample containing highly fluorinated silica nanoparticles at high concentration (FF-High with 2.9% silica content) was used, thereby, eliminating sample variations and the effect of surface modification and silica concentration. Other processing parameters such as saturation time, rate of pressure reduction, bubble fixation temperature, and foaming duration were all kept constant at 24 hrs, 1.5 MPa/s, 62°C, and 1 min, respectively. The resulting SEM micrographs of fracture surfaces are presented in Figure 7. Visual observation of the SEM micrographs indicates uniform cell size distribution across samples.

The effect of pressure on cell density and cell size at two different temperatures is presented in Figure 8. A slight decrease in cell density and a slight increase in average cell size are observed in Figures 8(a) and 8(b), although the changes are within experimental error. These results suggest that the effect of soaking pressure is low compared to the effect of surface modification for the samples investigated (FF-High with 2.9% silica content). This result can be explained by the fact that these samples already present high affinity towards CO$_2$ due to the presence of fluorinated surfactants, and therefore, nucleation of supercritical CO$_2$ at the (surface modified) silica/PMMA interface is already optimized.

The effect of temperature was found to be more complex than the effect of pressure. The effect of temperature at 8.97 MPa is presented in Figure 9. Results suggest a change of mechanism as a function of temperature. For example, at the lowest temperature, the cell density is low. Increasing the temperature leads to an increased cell density initially, but further increase in temperature leads to decreasing cell densities. These results can be explained by the change in the glass transition temperature of PMMA upon supercritical CO$_2$ saturation. It was shown before that CO$_2$ acts as a plasticizer for PMMA [55] and increasing pressure and temperature lead to more carbon dioxide absorption. The original glass transition temperature of neat PMMA is around 105°C. Previous studies reported that the plasticizer effect of carbon dioxide becomes evident at 8 MPa at which the glass transition temperature is reported to be 40°C [53, 58]. It was also shown that as the amount of absorbed CO$_2$ increases, the glass transition temperature decreases. When foaming takes place at temperatures lower than the glass transition temperature, not all nucleated bubbles can grow due to the high viscosity (resistance) of the PMMA bulk. This leads to low cell density at low temperatures. At high temperatures (above the $T_g$), although bubble growth is easier due to low polymer viscosity, growing bubbles could easily coalesce leading to decreased cell density. The highest cell density could be observed in the vicinity of the glass transition temperature as the viscosity at this state would be considerably higher than that at high temperatures preventing bubble coalescence but low enough that bubbles can grow.

Table 2 summarizes all the samples tested. The first row represents the neat PMMA sample, taken as a control. The next six samples indicate the effect of concentration and surface modification on cell density and cell size. The operating conditions (temperature and pressure) of the foaming process were varied for a single sample containing highly fluorinated silica nanoparticles (FF) with high concentration (2.9%). This
Figure 7: SEM images of the foamed highly fluorinated, high concentration (FF-High) sample at various pressures and temperatures.

Figure 8: (a) Cell density and (b) cell size of the highly fluorinated, high concentration (FF-High) sample processed at 65°C and 85°C at saturation pressures 8.97, 13.45 and 17.93 MPa. The error bars show the standard error of five samples.
Table 2: Summary of the cell density and cell size measurements for all samples used in the current study. $T$ is the foaming temperature and $P$ is the saturation pressure.

<table>
<thead>
<tr>
<th>Sample label</th>
<th>Surface modification (%)</th>
<th>Concentration (wt%)</th>
<th>$T$ ($^\circ$C)</th>
<th>$P$ (MPa)</th>
<th>Cell Density† ($10^9$ cm$^{-3}$)</th>
<th>Cell Diameter† ($\mu$m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PMMA</td>
<td>—</td>
<td>—</td>
<td>40</td>
<td>8.97</td>
<td>0.75 ± 0.05</td>
<td>9.62 ± 5.22</td>
</tr>
<tr>
<td>B</td>
<td>—</td>
<td>0.85</td>
<td>40</td>
<td>8.97</td>
<td>0.57 ± 0.09</td>
<td>5.36 ± 4.33</td>
</tr>
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<td>B</td>
<td>—</td>
<td>2.50</td>
<td>40</td>
<td>8.97</td>
<td>0.67 ± 0.29</td>
<td>6.90 ± 4.14</td>
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<td>F</td>
<td>~73</td>
<td>1.29</td>
<td>40</td>
<td>8.97</td>
<td>57.1 ± 10.4</td>
<td>1.62 ± 0.40</td>
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<tr>
<td>F</td>
<td>~73</td>
<td>3.16</td>
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<td>8.97</td>
<td>248 ± 12</td>
<td>1.29 ± 0.29</td>
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<tr>
<td>FF</td>
<td>&gt;100</td>
<td>1.38</td>
<td>40</td>
<td>8.97</td>
<td>132 ± 16</td>
<td>2.10 ± 0.80</td>
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<tr>
<td>FF</td>
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<td>248 ± 23</td>
<td>1.45 ± 0.54</td>
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<tr>
<td>FF*</td>
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<td>1.38 ± 0.44</td>
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<td>478 ± 32</td>
<td>1.19 ± 0.37</td>
</tr>
<tr>
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<td>13.45</td>
<td>478 ± 43</td>
<td>1.23 ± 0.34</td>
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<tr>
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<td>1.06 ± 0.32</td>
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<td>8.97</td>
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</tr>
<tr>
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<td>13.45</td>
<td>398 ± 44</td>
<td>1.52 ± 0.37</td>
</tr>
<tr>
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<td>85</td>
<td>17.93</td>
<td>400 ± 14</td>
<td>1.52 ± 0.43</td>
</tr>
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*These samples were all processed from the same silica synthesis batch.
†Standard deviation is given.

Figure 9: Cell densities of FF-High samples processed at 8.97 MPa as a function of saturation temperature. Error bars are the standard error for five samples.

strategy not only eliminates the concentration and surface modification effect but also ensures heterogeneous bubble nucleation mechanism is the dominant mechanism.

The cell densities and average cell sizes obtained in the current study are compared to those obtained from other studies performed with PMMA in Figure 10. Although the processing conditions in various studies differ from each other, considering that the goal of all of these studies is to obtain the greatest cell densities and the smallest average cell sizes, comparison of various results might prove to be useful. However, in order to achieve a direct comparison, results from various studies were normalized by the results of the neat PMMA in each study. In general, it is well known that cell density and cell size are inversely correlated with each other. This phenomenon can easily be observed in Figure 10.

4. Conclusions

Silica/poly(methyl methacrylate), PMMA, nanocomposites containing bare and fluoroalkane modified silica nanoparticles with an average size of 100 nm were studied under
varying supercritical carbon dioxide conditions. The effect of silica nanoparticle concentration and the extent of fluorination was studied systematically. The fracture surfaces of foamed samples were imaged with field emission scanning electron microscope and image processing tools were utilized to obtain cell density and average cell size. Our findings led to the following conclusions.

Surface modification of silica nanoparticles with CO\textsubscript{2}-philic fluoroalkane molecules significantly improved foam morphology. The average cell size decreased from 9.62\,µm in neat PMMA to 1.06\,µm with the addition of 2.9\% highly surface modified silica nanoparticles at 75\,°C and 8.97 MPa. The effect of surface fluorination of silica nanoparticles on cell size was observed for all nanocomposite samples regardless of the nanofiller concentration. Even the low fluorinated samples showed 6-7 times decrease in average cell size.

However, the extent of surface coverage (tethering density) of silica nanoparticles had a minor effect on foam morphology. The average cell sizes increased approximately 20\% upon going from ~73\% surface coverage to over 100\% surface coverage.

Increasing silica concentration had a remarkable effect on the average cell size and cell density. Increasing silica concentration from ~1.3\% to ~3.2\% led to a ~35\% decrease in average cell size.

Foam morphology strongly depends on the foaming temperature in reference to the glass transition temperature of the polymer. Although the glass transition of the neat PMMA used in the current study is around 105\,°C, soaking with supercritical CO\textsubscript{2}, which acts as a plasticizer, decreased it to approximately 65\,°C at 8.97 MPa soaking pressure. As a result, the foam morphology below and above the new glass transition temperature was influenced by different viscoelastic properties of the PMMA matrix. Our results suggest that the maximum cell density is reached in the vicinity of the glass transition temperature because the bubble growth was impeded either by high viscosity below the $T_g$ or by bubble coalescence above the $T_g$.

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References


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