Research Article

High Strain Rate Compressive Behavior of Polyurethane Resin and Polyurethane/Al₂O₃ Hollow Sphere Syntactic Foams

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Polyurethane resins and foams are finding extensive applications. Seat cushions and covers in automobiles are examples of these materials. In the present work, hollow alumina particles are used as fillers in polyurethane resin to develop closed-cell syntactic foams. The fabricated syntactic foams are tested for compressive properties at quasistatic and high strain rates. Strain rate sensitivity is an important concern for automotive applications due to the possibility of crash at high speeds. Both the polyurethane resin and the syntactic foam show strain rate sensitivity in compressive strength. It is observed that the compressive strength increases with strain rate. The energy absorbed up to 10% strain in the quasistatic regime is 400% higher for the syntactic foam in comparison to that of neat resin at the same strain rate.

1. Introduction

A class of closed cell porous composites called syntactic foam is prepared by dispersing hollow filler particles in a matrix medium [1, 2]. The mechanical (compressive, tensile, and flexural) [3, 4], thermal (coefficient of thermal expansion (CTE) and thermal conductivity) [5–7], and electrical (dielectric constant) [8–10] properties of syntactic foams can be tailored based on the volume fraction and the wall thickness of the reinforcing hollow spherical filler particles. Epoxy, vinyl ester, polyester, and bismaleimide resins are examples of polymers that have been used previously in syntactic foams [2, 11]. Lightweight metals such as magnesium and aluminum are also extensively used in fabricating syntactic foams [12]. The density of metal matrix syntactic foams can be as low as 1 g/cm³, whereas the densities of commonly fabricated polymer matrix syntactic foams (PMSFs) are in the range of 0.4–0.8 g/cm³. Hollow glass microballoons (HGM), fly ash cenospheres, and carbon and polymer hollow spheres are widely used as fillers in syntactic foams [2, 11–13]. Interest in developing high performance syntactic foams has resulted in development of hollow particles of ceramics such as SiC [14, 15] and Al₂O₃ [16]. CTE of hollow glass microballoon (HGM)/vinyl ester syntactic foams was found to be 60.4% lower than that of the matrix resin. In comparison, vinyl ester matrix syntactic foam containing SiC hollow spheres has CTE up to 79.3% lower in comparison to the matrix resin [14]. Thus, with the development of the ceramic hollow spheres, properties of syntactic foam beyond the commonly used HGM can be obtained.

Elastomeric matrices have not been commonly used in fabricating syntactic foams and only a few studies are available on polyurethane matrix syntactic foams containing glass hollow particles [17, 18]. Polyurethanes are commonly used in automotive industries as coatings, plastic components, and seat cushion foam [19]. This makes it necessary to understand the strain rate-dependent properties as high strain rates are encountered during an automobile crash. These polymers are also used as rocket motor liners for fastening the composite pellet grains within the motor [20]. In particular, to secure hydroxyl-terminated polybutadiene based composite solid propellants in the rocket motor, butadiene and isocyanate
based polyurethanes are used as the liner materials [20]. They are also used as thermal insulation in liquid fuel tanks in spacecraft launch vehicles [21, 22]. Another application for polyurethanes and their foams is in the biomedical field where they are used as bone scaffolds and models for cancellous bones due to their biocompatibility and possessing identical mechanical properties to bones [23, 24].

The existing literature on polyurethane matrix syntactic foams with HGM fillers is summarized in Table 1. Most of these studies have utilized polyurethane foam as the matrix materials and filled HGMs for mechanical property improvement. The gas filled foams are known to have very low mechanical properties. There is lack of results on utilizing the polyurethane resin as a solid matrix medium reinforced with hollow spheres. In the current study, polyurethane matrix syntactic foam reinforced with hollow alumina spheres is fabricated and characterized for quasistatic and high strain rate compressive properties.

### 2. Materials and Methods

The polyurethane resin and syntactic foam were fabricated by Deep Springs Technologies, Toledo, OH. The polyurethane potting system was a two-part polybutadiene based polyurethane with an isocyanate (30211/40008 Urethane Potting System, Potting Solutions LLC) mixed by weight 100:18.4 ratio. The polyurethane resin density was measured to be 1.49 ± 0.02 g/cm³.

The syntactic foams were prepared with the polyurethane resin as the matrix and hollow alumina spheres (Al₂O₃-HP) as the filler. The hollow alumina spheres are formed by a deposition technique followed by inert atmosphere sintering at 1650°C. A sacrificial polymer precursor is used to deposit the Al₂O₃ nano pellets over its surface and is followed by sintering process, where the polymer melts away and hollow Al₂O₃ spheres are formed. The average diameter and wall thickness of spheres were 3 mm and 146 μm, respectively, and the particles were treated with a silane adhesion promoter (Xiameter OFS-6020) prior to potting. The nominal Al₂O₃-HP true density was 0.55 g/cc. The density of synthesized syntactic foam was measured to be 1.13 ± 0.024 g/cm³.

The syntactic foam and the matrix resin were tested under compression at a wide range of quasistatic and high strain rates. Cylindrical specimens of 10 mm diameter and 10 mm thickness were tested at two different quasistatic strain rates of 0.001 s⁻¹ and 0.01 s⁻¹ using an Instron 4469 electromechanical test machine fitted with a 50 kN load cell. The load and displacement data were recorded using Bluehill software. At least five specimens were used for compression testing of each material. The high strain rate properties were investigated using an in-house developed split-Hopkinson pressure bar (SHPB) setup. The basic theory and the related physics of SHPB can be found in the published literature [25–27]. The SHPB consists of two long slender aluminum bars (Young’s modulus, density, and sound wave velocity of 70 GPa, 2700 kg/m³, and 5092 m/s, resp.) with the specimen sandwiched between them. The time dependent strain rate $\dot{\varepsilon}(t)$, stress $\sigma(t)$, and strain $\varepsilon(t)$ are calculated by

\[
\dot{\varepsilon}(t) = \frac{2q_0 \varepsilon_0(t)}{l_0},
\]

\[
\sigma(t) = \frac{A E \varepsilon_0(t)}{A_0},
\]

\[
\varepsilon(t) = \int_0^t \dot{\varepsilon}(t) dt,
\]

where, $q_0$ is the velocity of sound waves in the bar; $\varepsilon_0(t)$ and $\varepsilon(t)$ are the reflected and transmitted axial strain pulses as a function of time, respectively, $A$ and $E$ represent the cross-sectional area and the Young’s modulus of the bar material, respectively; and $A_0$ and $l_0$ are the cross-sectional area and the length of the test specimen, respectively. The stresses at the front and the back surface of the specimen are evaluated using the incident, reflected, and the transmitted strains using

\[
\sigma_{\text{front}} = E \left( \varepsilon_0(t) + \varepsilon_r(t) \right),
\]

\[
\sigma_{\text{back}} = E \varepsilon_0(t),
\]

### Table 1: Existing studies on polyurethane/hollow glass microballoon syntactic foams.

| Reference | Syntactic foam | Results |
|-----------|----------------|---------|
| [17]      | Polyurethane foam filled with HGM | Addition of 15% sepiolite nanofibers and 10% HGM resulted in 240% and 300% increase in the tensile and compressive strength of rigid polyurethane foam |
| [18]      | Polyurethane foam filled with HGM | 90 kg/m³ density foam showed 12% increase in compressive modulus and 10% increase in the compressive strength with addition of 1 wt.% of HGM |
| [30]      | Polyurethane foam filled with HGM (9.5 wt.%) | Isocyanate CPR-16 formulation: addition of HGM resulted in 45% increase in the compressive strength, with very little increase in the composite density. |
| [31]      | Polyurethane foam filled with HGM (0–40 vol.%) | Flexural modulus increases by 80% with addition of 10% HGM. |
| [32]      | Polyurethane foam filled with silane treated HGM (0–20 wt.%) | 50% increase in compressive modulus was observed with addition of 10 wt.% of HGM. |
| [33]      | Polyurethane resin filled with HGM (50 vol.%) | The polyurethane foam showed strain rate sensitivity and the collapse stress increased by 238% at a strain rate of 1185 s⁻¹ in comparison to quasistatic compression. |
where \( \varepsilon_i(t) \) is the incident axial strain pulse. The high strain rate result from the SHPB testing is valid only in the case of dynamic equilibrium, that is, when the stresses at the front and back surfaces of the specimen are comparable [28].

Optical photography of the initial and post test specimens was conducted using a Nikon D7000 DSLR camera equipped with an AF-S VR Micro-Nikkor 105mm f/2.8G IF-ED macro lens. A Hitachi S-3400N (Hitachi America Ltd., Tarrytown, NY) scanning electron microscope (SEM) was used to observe the specimens before and after testing. The SEM is equipped with secondary electron (SE) and back-scattered electron (BSE) detectors. The specimens were sputter coated with gold before the SEM observation using Leica EM SCD050 (Leica Microsystems Inc., Buffalo Grove, IL).

3. Results and Discussion

The structure of the polyurethane/\( \text{Al}_2\text{O}_3 \)-HP syntactic foam is shown in Figure 1(a). Alumina spheres are distributed uniformly in the polyurethane matrix material. The particle-matrix interface is continuous as observed in Figures 1(b) and 1(c). The micrographs of an alumina particle are shown in Figure 2. It is observed that the particle surface has a high degree of roughness and also some porosity. These features are helpful in promoting bonding with the matrix resin. In previous studies on syntactic foams, ceramic hollow spheres of SiC and \( \text{Al}_2\text{O}_3 \) were also observed to have rough surface and porous structure [14, 29]. It is further observed in Figure 1(d) that some cracks exist in the matrix very close to the particle-matrix interface. A large difference in the stiffness of the matrix and particle material causes these cracks to form during sample preparation for microscopy. These cracks are not a microstructure feature in the bulk syntactic foam slab.

A set of representative compressive stress-strain curves of the polyurethane resin at the quasistatic strain rates of 0.001 s\(^{-1}\) and 0.01 s\(^{-1}\) are plotted in Figure 3. The stress-strain curves of the polyurethane resin under quasistatic compression show a nonlinear behavior. Such resins have a high degree of viscoelasticity. The specimen fractures when the stress reaches the maximum value at the peak of the stress-strain curve. The polyurethane resin shows strain rate sensitivity under quasistatic compression testing conditions, as observed in Figure 3. Although the initial trend of the stress-strain curve is the same, the specimens show considerably higher strength and failure strain at the higher strain rate. The compressive strength and failure strain at 0.001 s\(^{-1}\) strain rate are measured to be 9.3 \( \pm \) 1.5 MPa and 58 \( \pm \) 2.0%, respectively. In comparison, the strength and failure strain at 0.01 s\(^{-1}\) strain rate are found to be 15.2 \( \pm \) 3.6 MPa and 68 \( \pm \) 1.8%, respectively. The compressive strength increased by 63.4% and the failure strain increased by 17.2% as the strain rate was increased by an order of magnitude.

The strain signals obtained from the SHPB for the polyurethane resin tested at a strain rate of 5800 s\(^{-1}\) are shown in Figure 4(a). The strain rate, stress, and strain are
Figure 2: (a) A single alumina particle of the type used in syntactic foams and (b) close up of the particle surface showing a high degree of surface roughness.

Figure 3: The quasistatic compressive response of the polyurethane resin. Data for one specimen is presented at two different strain rates for comparison.

evaluated from the strain signals using (1) and are shown in Figure 4(b). To obtain valid results from the high strain rate testing, the stress equilibrium in the specimen needs to be attained; that is, the stress on the front and the back surfaces of the specimen should be in close agreement. Figure 4(c) shows close matching in the stress at the two faces of the specimen and validates the test. Similar calculation is conducted for each specimen. The stress-strain curves evaluated from the high strain rate testing of the resin are shown in Figure 5. The general behavior of these curves is similar to the quasistatic compression curves. Note that the y-axis scale is different in Figures 3 and 5 because the specimens demonstrate considerably high strength at high strain rates. It can also be noted in Figure 5 that higher strain rate leads to higher strength in the specimens. All specimens are observed to fracture under the applied loading conditions. The peak stress for the quasistatic and the high strain rates are plotted as a function of the strain rates (on the log scale) and are shown in Figure 6. The peak stress under high strain rate testing for the resin is considerably higher (2–6 times higher for strain rates of 4600–6600 s$^{-1}$) when compared to quasistatic compression. From Figure 6, it could be observed that two trends emerge from the variation of peak stress with respect to strain rates and that the two trends intersect at a strain rate of 5600 s$^{-1}$. These observations about the neat resin will be useful in understanding the behavior of syntactic foams.

The stress-strain response of the polyurethane/Al$_2$O$_3$-HP syntactic foam is shown in Figure 7. The general trend of the stress-strain curves of the polyurethane syntactic foam is different than that of the polyurethane resin. The stress increases linearly with the strain in the initial elastic region. After the yield point, the specimen deforms plastically before it fractures. The transformation of stress-strain curve from the polyurethane resin to its syntactic foam composite can be attributed to the addition of rigid Al$_2$O$_3$ spheres into the viscoelastic polyurethane matrix. Rigid Al$_2$O$_3$ particles are load-bearing elements in the syntactic foam microstructure. These particles reduce the overall elasticity of the material. Peaks observed in the stress-strain curve can be attributed to successive failure of the Al$_2$O$_3$ spheres as the compressive strain increases. In comparison to the quasistatic testing, the high strain rate results show higher stress values at comparable strain. The yield strength of the polyurethane syntactic foam is specified by the 0.2% offset method and is plotted as a function of the strain rate (on a logarithmic scale) in Figure 8. The yield strength of the polyurethane syntactic foam shows a trend in the strain rate sensitivity that is similar to the matrix resin. It can be expressed as a bilinear relation and the intersection point is observed at a strain rate of 800 s$^{-1}$.

It should be noted that the polyurethane syntactic foams show quasistatic compressive failure strain in the range of 60–70%. Since there is a considerable difference in the shape of the stress-strain graphs of neat resin and syntactic foams, the energy absorbed up to 10% strain is calculated for comparison and plotted as a function of the strain rate in Figure 9, which
mainly includes elastic energy. The energy absorption of the foam specimens is considerably higher in comparison to the neat resin. In the quasistatic range, the energy absorbed by the foam is 3-4 times the neat resin values. The higher energy absorption in the foam specimens can be attributed to the Al₂O₃-HP. The failure of Al₂O₃-HP results in increase in cumulative energy absorption at 10% strain. This type of behavior can be utilized in energy absorbing crash structures in automotive and marine industry, where understanding the energy absorption characteristics with respect to strain rate is important.

Two points are marked as A and B in the quasistatic compressive stress-strain graph of syntactic foam Figure 7(a). The failure of a representative syntactic foam specimen corresponding to these points is shown in Figure 10. It is observed in Figure 10(a) that the initial specimen failure starts as a shear crack across the entire specimen. This point is marked as point "A" in Figure 7(a). Al₂O₃-HP also starts to fracture at point A. Continued compression results in complete crushing of particles and fragmentation of the specimen as observed in Figure 10(b), which represents the final failure of the specimen at the end of the densification region, marked as...
“B” in Figure 7(a). These features can be compared to that of a specimen that was tested at 1430 s\(^{-1}\) strain rate, shown in Figure II. This specimen shows failure due to shear cracking. The specimen has not been completely crushed because the strain encountered in the high strain rate testing is smaller compared to the strain to which the quasistatic specimens were subjected. Al\(_2\)O\(_3\)-HP present in this specimen shows signs of crushing. The high strain rate compressive failure is further observed through SEM on a specimen that was subjected to strain rate of 1670 s\(^{-1}\). The arrows marked in Figure 12(a) indicate the compression direction. Micrographs show failure of the Al\(_2\)O\(_3\)-HP in Figures 12(b), 12(c), and 12(d). It can be observed in these micrographs that the failed hollow spheres are adhering to the matrix resin, indicating a strong particle-matrix interface and effective load transfer from the matrix to the particle.

4. Conclusions

Polyurethane resin and polyurethane/Al\(_2\)O\(_3\)-HP syntactic foams are studied for compressive properties under quasistatic and high strain rates. The high strain rate compression testing is conducted using a split-Hopkinson pressure bar. The results from the present study can be summarized as follows.
The polyurethane resin shows a viscoelastic quasistatic stress-strain response. Addition of Al₃O₅-HP changes the stress-strain trend of the material. Syntactic foams are observed to have an initial linear elastic region followed by a plastic region that corresponds to particle crushing.

(ii) Both the polyurethane resin and its syntactic foam show strain rate sensitivity in compressive strength.

Two different trends were observed in the plot of compressive strength as a function of the strain rate (on the logarithmic scale) for the quasistatic and the high strain rate regimes.

(iii) The energy absorbed up to 10% strain in the quasistatic regime is 400% higher for the syntactic foam in comparison to the neat resin.
Figure 10: Fracture features of polyurethane/Al$_2$O$_3$-HP syntactic foam specimens tested under compression at strain rates of 0.01 s$^{-1}$. Parts (a) and (b) correspond to the points A and B, respectively, marked in Figure 7(a).

Figure 11: Fracture features of polyurethane/Al$_2$O$_3$-HP syntactic foam specimens tested under compression at strain rates of 1430 s$^{-1}$.

Figure 12: SEM images of a polyurethane/Al$_2$O$_3$ syntactic foam specimen tested at a strain rate of 1670 s$^{-1}$ showing the (a) specimen shear failure and (b), (c), and (d) failure of a hollow particle.
(iv) The failure of the syntactic foam occurs through shearing of matrix resin, followed by failure of the hollow particles. The interface between the Al$_2$O$_3$-HP and the matrix resin on the failed specimens was intact, indicating effective load transfer between the matrix and the hollow particle.

Conflict of Interests

The authors declare no conflict of interests.

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