Research Article

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Materials characterization of advanced fillers for composites engineering applications

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Abstract: Four different minerals were investigated; hollow spheres of calcium carbonate, platy mica, needle like wollastonite and glassy perlite and characterized via iGC for surface energy, Freeman powder rheology for flow characterization, cyclic uniaxial die compaction for modulus of elasticity and frequency dependent sound absorption properties. Particle surface energy and particle shape strongly affected the packing density of powder beds. In the case of higher porosity and thus lower bulk density, the powders acoustic absorption was higher in comparison with higher packing density materials. Surface energy profiles and surface energy distributions revealed clear convergence with powder rheology data, where the character of the powder flow at defined consolidation stresses was mirroring either the high cohesion powders properties connected with the high surface energy or powder free flowing characteristics, as reflected in low cohesion of the powder matrix.

Keywords: mineral fillers, surface properties, sound absorption, mechanical properties, powder rheology

1 Introduction

Modern globalized automotive and aerospace production companies apply advanced composites and coating systems in their final product lines. At the present time, the goal is to enhance overall price/performance ratio and increase productivity with respect to the processing time, using a lean technology approach and energy input [1]. That is why a wide range of synthetic or natural based micro/nano fillers were being used in technical practice [2–4]. For that reason detailed knowledge of the filler particles surface interface is important for exact adjustments of the consecutive technological operations during production cycles. Micro/nano particles filled synthetic polymers are commonly used for the structural as well as non-structural components and advanced coatings applications [5]. Previously minerals served as the additives in polymer systems mainly as a cost reducing component [6]. Progressing technological improvements in minerals processing and polymer chemistry allowed their novel role as the functional additives, thus bringing specific mechanical and functional properties to the final composites structures [5, 7–9]. The packing density of the powder bed is strongly dependent on the applied consolidation stress, surface energy [10], surface roughness and particle shape of the individual particles [11–13]. That is why our focus in this study was aimed at the characterization of the physico-chemical (surface energy, surface energy distribution and surface energy profiles characterization) and material properties (powder rheology, sound absorption and die compaction...
Table 1: Studied mineral samples labelling and description

| Sample | Description                                                                                       |
|--------|---------------------------------------------------------------------------------------------------|
| 1      | Calcium carbonate hollow spheres (University of Birmingham, UK), $d_{50} = 991$ nm diameter*, density 2.40 g/cm³#. |
| 2      | Muscovite Mica (Imerys, Kings Mountains, USA), $d_{50} = 17$ µm diameter*, specific surface area 9.7 m²/g, platy with an aspect ratio 1.7, density 0.45 g/cm³#. |
| 3      | Wollastonite type VANSIL W-10 (Vanderbilt Minerals, Norwalk, USA), $d_{50} = 49$ µm diameter*, specific surface area 0.5 m²/g, needle like with an aspect ratio 13.5, density 0.65 g/cm³#. |
| 4      | Perlite, volcanic glass (Supreme Perlite Company, USA), $d_{50} = 447$ nm diameter*, density 1.10 g/cm³#. |

*As measured in Palacky University lab. #According to manufacturer’s data sheet.

behavior) both in the individual as well as in particles collective cooperation setups of free poured powder beds and die compression configurations. During die compression the deformation of the free powder bed is continuously proceeding, hence increasing the compression stress in the system. This is recorded as the stress-displacement dependency as obtained for given deformation rates and applied maximum stress. Obtained loading and unloading curves reflect the intimate inter-particle interactions and ongoing processes of deformation and displacement associated with the proceeding applied compression stress. The following sequence of individual processes during agglomeration has been recognized; namely the feed of loose packing, elastic–plastic contact deformation, pore filling by fine particles, plastic deformation of particles to create large contact areas, breakage of edges, particle breakage and plastic deformation of entire sample [14]. During the loading cycle, the deformation took place in contacts as well as the volume deformation of individual powder particles [15, 16], reflecting compaction of the sample with translation and rotation movements of particles, at relatively small contact deformations followed by the deformation of entire particles and their breakage [14]. Elastic recovery of the compacted powder bed is an important processing parameter affecting consecutive creation of defects and fractures in the compressed material due to the proceeding fast recovery or spring back, thus influencing final mechanical properties of the compressed body [14, 17].

3 Methods

3.1 Thermal analysis

For powder moisture content thermogravimetry (TG) and differential thermal analysis (DTA) experiments were performed on a simultaneous DTA-TG apparatus (Shimadzu DTG 60, Japan). Measurements were performed at a heat flow rate of 10°C/min in a static air atmosphere at the temperature range from 30°C to 300°C. Samples moisture content as determined by thermal analysis (TG DTG) ranged from 0.1 to 0.9 wt.%.

3.2 Inverse gas chromatography surface energy analysis

Inverse gas chromatography (iGC) was performed by surface energy analyzer (SEA) (Surface Measurement Systems, UK). Samples were placed in 4 mm (internal diameter) columns to give a total surface area of approximately 0.5 m². Nonane, octane, hexane, heptane, dichloromethane, acetone, acetonitrile, ethyl acetate and ethanol were used as eluents for iGC. All reagents were obtained from Sigma Aldrich (USA) and were of analytical grade. The injection of vapors was controlled in order to give pre-determined fractional coverage of the sample. By gradually increasing the amount of vapors injected, it is possible to build up a surface heterogeneity plot [5, 18]. Surface energy distribution was calculated according to the standard methodology as described by Mohammadi-Jam and Waters [19].

2 Materials

As filler particles the inorganic minerals calcium carbonate hollow spheres (The University of Birmingham), muscovite mica (Imerys, Kings Mountains, USA), wollastonite type VANSIL W-10 (Vanderbilt Minerals, Norwalk, USA) and perlite (Supreme Perlite Company, USA) were used in this study. The basic description, labelling and physical parameters of the studied fillers are given in Table 1.
3.3 Powder rheology

Powder rheology measurements were conducted on a FT4 Powder rheometer (Freeman Technology, UK). All experiments were performed under the ambient laboratory temperature of 24°C and relative humidity of 45%. Consolidation stresses of 9 kPa were used for samples 1, 2 and 4, but 22 kPa for sample 3.

3.4 Uniaxial die compression testing

Uniaxial die compression testing was performed on universal testing machine Autograph AGS-X (Shimadzu, Japan) equipped with the Compact Thermostatic Chamber TCE Series. A stainless steel cylindrical die of 13 mm inner diameter and 40 mm height was used. Measurements were performed at 0.2, 1.2 and 10 mm/min deformation rates. Each measurement was repeated three times and mean average values were calculated. The compressive as well as release stress-strain dependencies were then determined. From these tests the modulus of elasticity (E) was calculated as the slope of the first linear part of unloading curve (down ramp), and the bulk modulus (K) was calculated from the same data according to Eq. (1), where at this phase of unloading only the elastic response of the studied powder material took place [14, 20]:

$$K_{ab} = \frac{\sigma^A_z - \sigma^B_z}{e^A_z - e^B_z} = K + 4K\frac{1 - 2\nu}{1 + 2\nu}$$

(1)

where \(K_{ab}\) is the slope of the down ramp from maximum axial stress (point A) to half the maximum axial stress (point B), \(\sigma_z\) is the axial stress, \(e_z\) is the axial strain and \(\nu\) is the Poisson’s ratio [20]. In this investigation, the Poisson’s ratio was assumed to have a value of 0.25 as referred earlier by [20, 21].

3.5 Powder bed sound absorption testing

Sound absorption coefficient measurements of freely poured powder beds were performed by means of the transfer function method ISO 10534-2 standard [22]. There were measured frequency dependencies of the sound absorption coefficient by means of a two-microphone impedance tube (BK 4206) in combination with three-channel signal PULSE multi-analyzer (BK 3560-B-030) and power amplifier (BK 2706) in the frequency range of 150-6400 Hz (Brüel & Kjær, Denmark). The normal incidence sound wave absorption of the tested loose powder samples of defined layer thickness (ranging from 10 to 100 mm) was determined. All experiments were performed under ambient laboratory conditions of 45% relative humidity and at 24°C. The sound absorption coefficient is given by the ratio of the dissipated power in the studied material to the incident acoustic power [5, 23]. Based on the measured frequency dependencies of the sound absorption coefficient the noise reduction coefficient (NRC) was calculated as the arithmetical average of the sound absorption coefficient at the excitation frequencies 250, 500, 1000 and 2000 Hz [5]. The longitudinal elastic coefficient \(K_1\) of the powder bed is similar to Young’s modulus of elasticity of the material and was expressed as follows [24, 25]:

$$K_1 = c^2\rho_b = (4hf_{p1})^2\rho_b$$

(2)

where \(c\) is the propagating sound speed, \(\rho_b\) is the powder bed bulk density, \(h\) is the powder bed height and \(f_{p1}\) is the primary absorption peak frequency.

4 Results and discussion

As mentioned in the introduction, inorganic nano/micro particles are used as functional fillers in many polymer based composites, where the dominant factors are particle uniformity, shape, diameter and surface chemistry, which influence the physical and material properties of the final product. These properties include for example mechanical strength, fracture performance, thermal behavior, barrier properties, and electrical conductivity [5]. It is very well known from previous studies, that the filler particle-matrix adhesion is of paramount importance for high strength composites preparation. That is why, we have studied surface energy profiles and surface energy distribution by means of iGC testing of the studied materials with respect to obtain their surface wettability characteristics in detail. As shown in Figure 1, observed surface energy profiles have typical exponential decay dependencies, indicating two major areas: the high energy areas observed in the surface coverage regions ranging from 0 to 0.2 depending on the type of the studied powders reaching maximum of the total surface energy in the range from 60 to 85 mJ/m² and the low energy areas observed in the surface coverage regions from 0.05 to 1 exhibiting equilibrium total surface energy of about 35 to 60 mJ/m². The highest surface reactivity as represented by the observed total surface energy was found for the sample 2 (mica) characteristic with dispersive part of the surface energy dominant over the polar part. The surface energy polar part ranged from 16 to 1 mJ/m².
Table 2: Results from the shear cell flow experiments, measured at the consolidation stress of 9 kPa calcium carbonate, mica and perlite (samples 1, 2 and 4) and 22 kPa wollastonite (sample 3) and the temperature of 25°C

| Sample | Cohesion [kPa] | UYS [kPa] | MPS [kPa] | FF [-] | AIF [%] | MCS [kPa] |
|--------|----------------|-----------|-----------|--------|---------|-----------|
| 1      | 0.86           | 2.98      | 14.04     | 4.71   | 30.13   | 3.67      |
| 2      | 0.87           | 3.38      | 19.50     | 5.76   | 35.60   | 4.25      |
| 3      | 0.77           | 4.10      | 64.40     | 15.70  | 48.90   | 8.49      |
| 4      | 0.20           | 0.78      | 16.20     | 20.70  | 36.50   | 3.91      |

UYS – unconfined yield strength, MPS – major principal stress, FF – flow function, AIF – angle of internal friction, MCS – minor consolidation stress.

Figure 1: Surface energy profiles of studied fillers: A – calcium carbonate hollow spheres (sample 1), B – mica (sample 2), C – wollastonite (sample 3), D – perlite (sample 4). Square – polar part of the surface energy, triangle – dispersive part and circle – total surface energy

The surface energy distributions of studied powders are shown in Figure 2. These are of a typical peak like shapes where the highest population of 5.2% area increment at the total surface energy of 72 mJ/m² (sample 1) was found. The lowest population of about 0.55% area increment at the total surface energy of 45 mJ/m² was found for the sample 4 (perlite). Similar patterns were observed for the distributions of the polar parts of the surface energy, however, the dominating part was the dispersive energy for all tested powder materials. Obtained dispersive part distributions ranged from 45 to 85 mJ/m² (calcium carbonate, sample 1), from 44 to 84 mJ/m² (mica, sample 2), from 42 to 50 mJ/m² (wollastonite, sample 3) and from 28 to 76 mJ/m² (perlite, sample 4). The polar parts distributions ranged from 0 to 17 mJ/m². Density function theory calculations have theoretically shown that negative surface energies are possible for porous alumina in its high surface area nanocrystalline form [26, 27]. This was attributed to the particularly strongly bound state of dissociated water. However the negative energies for the perlite sample are probably either an artefact of the model or due to its glassy nature.

Results of the powder rheology measurements are shown in Figure 3. These are of a typical yield locus and Mohr’s circles dependencies as a function of the applied normal stresses. The obtained individual shear flow parameters are given in Table 2. It is evident, that the most cohesive character of the powder flow was found for the samples 1 (calcium carbonate) and 2 (mica) as indicated by the observed flow functions of 4.7 calcium carbonate (sample
Figure 2: Surface energy distribution of studied fillers: A – calcium carbonate hollow spheres (sample 1), B – mica (sample 2), C – wollastonite (sample 3), D – perlite (sample 4). Square – polar part of the surface energy, triangle – dispersive part and circle – total surface energy.

Figure 3: Yield locus and the Mohr’s circles of the tested materials as obtained by shear cell experiments at applied 9 kPa consolidation stress (samples 1, 2 and 4) and at 22 kPa consolidation stress (sample 3): A – calcium carbonate hollow spheres (sample 1), B – mica (sample 2), C – wollastonite (sample 3), D – perlite (sample 4).
Table 3: Results of the acoustic and mechanical testing of the studied powder fillers

| Sample | Quantity | 10 | 15 | 20 | 50 | 100 |
|--------|----------|----|----|----|----|-----|
| 1      | NRC [-]  | 0.149 | 0.180 | 0.167 | 0.167 | 0.163 |
|        | $f_{p1}$ [Hz] | 808 | 544 | 520 | 312 | 304 |
|        | $c$ [m·s⁻¹]   | 32.2 | 32.6 | 41.6 | 62.4 | 121.6 |
|        | $K_l$ [MPa] | 2.5 | 2.6 | 4.2 | 9.4 | 35.5 |
| 2      | NRC [-]  | 0.233 | 0.235 | 0.222 | 0.370 | 0.261 |
|        | $f_{p1}$ [Hz] | 744 | 464 | 344 | 240 | 216 |
|        | $c$ [m·s⁻¹]   | 29.8 | 27.8 | 27.5 | 48.0 | 86.4 |
|        | $K_l$ [MPa] | 0.4 | 0.4 | 0.3 | 1.0 | 3.4 |
| 3      | NRC [-]  | 0.216 | 0.191 | 0.248 | 0.196 | 0.216 |
|        | $f_{p1}$ [Hz] | 736 | 382 | 280 | 168 | 144 |
|        | $c$ [m·s⁻¹]   | 29.4 | 22.9 | 22.4 | 33.6 | 57.6 |
|        | $K_l$ [MPa] | 0.4 | 0.3 | 0.3 | 1.0 | 2.2 |
| 4      | NRC [-]  | 0.106 | 0.095 | 0.116 | 0.136 | 0.124 |
|        | $f_{p1}$ [Hz] | 848 | 600 | 472 | 304 | 296 |
|        | $c$ [m·s⁻¹]   | 33.9 | 36.0 | 37.8 | 60.8 | 118.4 |
|        | $K_l$ [MPa] | 1.3 | 1.4 | 1.6 | 4.0 | 15.3 |

1) and 5.7 mica (sample 2) typical for cohesive powders. In contrary to this, the observed flow curves for the samples 3 (wollastonite) and 4 (perlite) indicate typical free flow powders behavior, as confirmed also by flow functions exceeding values of 15. The highest magnitude of the major principal stress of 64.4 kPa was found for sample 3 (wollastonite) at a 22 kPa consolidation stress. Other samples exhibited lower values of the major principal stress ranging from 14.04 kPa (sample 1) to 19.5 kPa (sample 2) at 9 kPa consolidation stress. The observed data correspond also with the obtained magnitudes of the unconfined yield strengths of the studied materials, which were ranging from 0.78 kPa (sample 4) to 3.38 kPa (sample 2) at the 9 kPa consolidation stress, and of 4.1 kPa (sample 3) at the 22 kPa consolidation stress.

In general, in the matrix of the sound attenuating material dissipation of the incident acoustic wave is converted to mechanical energy and heat [28]. This proceeds by combination of the following processes. Firstly by friction of the vibrating air molecules on the walls during their penetration into the pores of the sound absorbing material (by lowering kinetic energy of the incident sound field, effectiveness of this process increases with growing porosity of the absorption material). Secondly by decreasing the potential energy of the acoustic wave penetrating into the material (by lowering acoustic pressure due to the heat exchange between air and the skeleton of the absorbing material during periodic pressure changes), and thirdly by non–elastic deformation of the absorbing material body [29, 30]. Results of sound absorption experiments are shown in Figures 4 and 5 and Table 3.

The highest magnitude of the sound absorption coefficient was 0.54 (at the frequency of 460 Hz) simultaneously with the highest value of the noise reduction coefficient (NRC = 0.235, at the 15 mm powder bed height) were found for the powder bed of the sample 2 (mica) due to its high porosity in comparison to the other tested samples. As mentioned earlier, these results correspond very well with the observed highest cohesion of 0.87 kPa (see Table 2) and the highest surface energy of 72 mJ/m² (see Figure 2), as obtained for the sample 2. The lowest poros-
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Figure 5: Frequency dependencies of the sound absorption coefficient of studied particular fillers: A – wollastonite (sample 3), B – perlite (sample 4). Inset: Powder bed height in mm

Figure 6: Stress vs relative displacement dependencies observed during chamber uniaxial compression testing (at 10 mm/min deformation rate and 80 kN compression force): A - hollow spheres calcium carbonate filler (sample 1), B - wollastonite filler (sample 3)

ity was found for the sample 4 as indicated both by powder rheology (see the lowest value of the parameter of cohesion of 0.2 kPa in Table 2) and the lowest NRC coefficient of 0.095 at the same bed height as in the case of the mica data. These data correspond very well in the same fashion as in the previous mica (sample 2) with the surface energy distribution, which was the most populated for the perlite (sample 4) at the 45 mJ/m$^2$, which was found to be the lowest surface energy in comparison with other tested powders.

In Figure 5 results of the effect of the powder bed height on frequency dependencies of the sound absorption coefficient are shown. Here, a typical shift of the primary absorption peak frequency ($f_{p_1}$) position to lower frequencies was found (see Figure 4 and Table 3) with increasing powder bed height [5]. By the application of Eq. (2) the longitudinal elastic modulus ($K_l$) and propagated sound velocity ($c$) were calculated. Results indicate, that the highest velocity of the propagated sound wave was found in the calcium carbonate and perlite powders (samples 1 and 4) of 32.6 m/s and 36.0 m/s respectively (for the powder bed height of 15 mm). These results were confirmed also with the observed highest packing density of the latter calcium carbonate and perlite powders (samples 1 and 4) hence resulting in the obtained highest structural stiffness of the system as reflected in observed values of the $K_l$ being 2.6 MPa calcium carbonate (sample 1) and 1.4 MPa perlite (sample 4).

Results of the uniaxial die compression testing of the calcium carbonate and wollastonite (samples 1 and 3) are shown in Figures 6 and 7. Typical patterns of the loading and unloading curves were obtained with a characteristic hysteresis loop. According to the Eq. (1) the calculated bulk modulus ($K$) of 643 MPa (calcium carbonate) and 852 MPa (wollastonite) as well as elastic modulus ($E$) of 1647 MPa (calcium carbonate) and 2191 MPa (wollastonite) were obtained for 10 mm/min deformation rates and 600 MPa maximum compression stress. Results of the effect of the deformation rate on obtained bulk modulus are illustrated for the calcium carbonate (sample 1) in Figure 7. Here a typical increase of the bulk modulus with increasing deforma-
Figure 7: Stress vs relative displacement dependencies observed for hollow spheres calcium carbonate filler (sample 1) during chamber uniaxial compression testing (at 80 kN compression force): A - 10 mm/min deformation rate, B - 1.2 mm/min deformation rate and C - 0.2 mm/min deformation rate.

Figure 8: Results of the cyclic closed shell compression loading experiments of the calcium carbonate (sample 1) (deformation rate 10 mm/min, maximum force 80 kN): full circle – input energy (J), triangle up – stored energy (J), square – dissipated energy (J).

Figure 9: Results of the cyclic closed shell compression loading experiments of the calcium carbonate (sample 1) (deformation rate 10 mm/min, maximum force 80 kN): full circle – dissipated energy (J), triangle up – elastic modulus (MPa).

The equilibrium energy dissipation of 2.8 J (cycle 10). The complexity of the observed sequence of the individual deformation events proceeding in the course of the die compression of the tested hollow sphere calcium carbonate (sample 1) is evident from the Figure 9 (elastic modulus vs. cycle no. dependency). At the beginning of the compression process a clear increase in the elastic modulus between cycles 1 and 2 indicates progressing feed of loose packing followed by elastic-plastic contact deformation as found by Berg et al. [20]. The latter step was followed by an observed decrease of the elastic modulus from 1.93 MPa (cycle 2) to 1.33 MPa (cycle 3) due to the progressive pore filling by fine particles followed by plastic deformation of particles to create large contact areas as indicated by a linear increase of the $E$ in the consecutive cycles from cycle 3 to cycle 6, reaching a maximum value of elastic modulus 6.18 MPa. This reflects the state of the compressed matrix still preserving the hollow spherical structure of the filler material. This stage was then followed by proceed-
of the entire compressed body were clearly identified from the cyclic stress vs. relative displacement measurements.

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