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Surface free energy and mechanical performance of LDPE/CBF composites containing toxic-metal free filler



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ABSTRACT

Heavy-metal contamination in children's toys is a widespread problem, and the international community has issued a series of safety standards to restrict and control the use of toxic metals in toys. In this work, a colored filler (CBF) was prepared using pearl oyster shell (POS) as the green raw material and azo dye as the colorant. Its surface properties were subsequently studied in comparison to those of POS powder using the inverse gas chromatography method. The dispersion surface free energy profiles for both CBF and POS showed that this component contributed the major part (> 70%) to the total surface free energy. The CBF possessed lower polar surface free energy and was relatively more hydrophobic. It also showed a lower thermodynamic work of cohesion, allowing its better dispersion in a low density polyethylene (LDPE) matrix. Mechanical performance studies showed that adding CBF could significantly increase the tensile strength, elastic modulus, flexural strength and flexural modulus of LDPE composites. The absence of toxic metals coupled with excellent mechanical performance makes the CBF an ideal candidate as a filler for children's toys fabrication.

1. Introduction

Coloring adds aesthetic appeal and enhances values to toys and other children's products. The most widely used colorants include dyes and pigments, which are distinguishable by their mode of application. Among them, inorganic pigments have been applied since ancient times [1], and are considered to have advantages over their organic counterparts in terms of heat and light stability. However, most inorganic pigments are derived from toxic or transition metals, such as Cd, Pb, Cr, Co, Hg and Ce, which adversely affect the environment and represent a threat to human health [2]. These metals may be released and absorbed via saliva during mouthing, sweat during dermal contact or gastric fluids after ingestion [3,4]. Although direct release of these metals from plastics may be expected to be low due to the complex and strong polymer structure of toys, scenarios of repeated exposure pose significant concerns for children's health.

Mollusk shells (e.g. clam, oyster, mussel and pearl oyster shells), with their predominantly calcium carbonate (CaCO₃) content plus a small amount of biomacromolecules, can be used as a potential

substitute for commercial CaCO3 filler [5]. Fombuena et al. [6] prepared epoxy resin composites incorporating seashells. Li et al. [7] reported excellent mechanical performance of polypropylene composites filled with Mytilus edulis shell. Mustata et al. [8] also investigated the thermal properties of composites based on epoxy resin and conch shells. Considering the increasing concern of toxic heavy metal contamination in plastics and the inherent mechanical properties of mollusk shell material, we attempted to prepare a colored filler (CBF) using pearl oyster shell (POS) as the green raw material and azo dye as the colorant. However, it has been well recognized that the nature of the filler e.g. its surface free energy, particle size, surface area, and structure—or degree of irregularity, influences its reinforcing ability. A better understanding of the filler's surface properties is critical to determining the most effective polymer reinforcement fillers. Therefore, in the present work, the surface properties of CBF were determined in comparison to those of POS powder using the inverse gas chromatography (IGC) method. In order to provide further evidence for its suitability in larger scale practical applications, Vicat softening temperature and the mechanical performance of LDPE/CBF composites

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were also investigated. Ultimately the work is aimed at probing the promising use of CBF in the masterbatch industry and, therefore, solving the significant problem of high levels of toxic metals in toys.

2. Theoretical basis

The basic theory of IGC is presented here and more details can be found elsewhere [9–16]. The surface free energy is defined as the average free energy per unit area surface of a material. The total free surface energy (γ_s^T) is often the combination of dispersion (γ_s^D) and polar (γ_s^P) components. Dispersive (apolar) interactions, also known as Lifshitz-van der Waals interactions, consist of London, Keesom, and Debye interactions. Specific (polar) interactions explain other types of interactions, such as acid-base interactions, hydrogen bonding and π bonding [17]. A standard method of surface characterization is that the γ_s^D is first determined using a series of *n*-alkanes as probes (in this case, octane, nonane and decane); then the acid-base parameters can be calculated from polar probes (in this case, toluene, acetone, acetonitrile and dichloromethane[DCM]). For the calculation of γ_s^D , the Dorris-Gray method [18] is commonly used and thus was applied in this work. The contribution of acid-base properties of a solid is often obtained by first measuring the free energies of adsorption (ΔG^p) for different polar probes. From the ΔG^p , we can calculate the acid-base numbers related to the polar surface free energy.

Knowledge of surface free energy differences between filler particles would allow an objective decision about the extent of the interaction between filler and polymer matrix and provide a hint on the dispersion ability of the filler within the polymer matrix or about the likely extent of filler agglomeration. The extent of filler-filler interaction can be quantified by the thermodynamic work of cohesion (W_C), which in turn can be determined by IGC [19] according to the geometric mean equation [20–22]:

$$W_C = 2\sqrt{\gamma_S^D \cdot \gamma_S^D} + 2\sqrt{\gamma_S^P \cdot \gamma_S^P} = 2\gamma_S^T \tag{1}$$

Knowledge of the surface free energy and its components for CBF and POS, can be used to calculate the $W_{\rm C}$ between filler particles.

3. Experimental materials and methods

3.1. Materials

Commercial LDPE (2426H) with a melt flow rate of 1.9 g/10 min at 190 °C was supplied by Sinopec Maoming Company, China. Direct Red 28 dye (DR 28) was provided by Yiwu Yu Fang Pigment Co., Ltd., China. The polar and nonpolar probes (HPLC purity, 99.0%) were purchased from Sigma-Aldrich (St-Louis, MO, USA). The raw pearl oyster shell was collected from a pearl-processing factory in Zhuji city, China. It was first washed to remove attached impurities, and then calcined at 350 °C to remove the stratum corneum. The dried powder was subjected to fine grinding to obtain the pearl oyster shell (POS) powder. XRD analysis indicated that the major crystalline phase of POS was calcite (CaCO₃, JCPDS card no. 86-2334). The chemical compositions were determined by X-ray fluorescence (XRF, Shimadzu XRF-1800) and the results were as follows (wt%): C 14.2%, O 52.6%, Ca 33.2%.

3.2. CBF preparation

POS powder was mixed with DR 28 and water at a weight ratio of 200:1:300. After vigorous stirring for 0.5 h, the mixture was left standing for 24 h, then filtered, and the filter cake dried. The dried cake was ground to obtain the CBF. Photographs of the POS and CBF powders are shown in Fig. 1. The volume-weighted distribution analysis showed that the mean and median particle sizes of the POS were 8.5 and 7.6 μ m, respectively; of the CBF, 9.1 and 7.6 μ m, respectively. The

BET specific surface areas of POS and CBF were determined as 0.9 and $1.5 \text{ m}^2/\text{g}$, respectively, using the iGC Surface Energy Analyzer (iGC-SEA, Surface Measurement Systems, Alperton, UK) [23].

3.3. LDPE composites preparation

Before mixing, the LDPE matrix and CBF powder were oven-dried at 80 °C overnight. A series of LDPE and CBF powders with different weight ratios (100/0, 98/2, 95/5, 90/10, 85/15, 80/20, 70/30 and 60/40) were mixed with 5 wt% compatilizer PE-g-MAH, 0.1 wt% antioxidant 168, 1.5 wt% lubricant TAF and 0.1 wt% mineral oil, using a SHJ-35 parallel co-rotating twin screw extruder (Nanjing, China). The extruder has eleven independent temperature zones and temperatures were set at 200 °C (zones 1–3), 205 °C (zone 4–7) and 215 °C (zone 8–11). The extrudates were pelletized, and a plastic injection-molding machine (MA900/260, Ningbo, China) was then used to prepare test specimens. The injection molding machine has five temperature zones and temperatures were set at 180 °C (zone 1–2), 200 °C (zone 3–4) and 185 °C (zone 5). Photographs of the test specimens are displayed in Fig. 2.

3.4. Characterization and tests

Surface free energy characterization and specific surface area determination were both carried out using the iGC-SEA. The data were analyzed using the advanced Cirrus Plus Analysis Software of iGC-SEA. For all the experiments, approximately 300 mg of powder were packed into individual dimethyldichlorosilane-treated glass columns. The samples were run at a series of surface coverages (2–16%) with polar and nonpolar molecular probes to determine γ_s^D and γ_s^P as well as ΔG^P . The sample column was preconditioned for 1 h at 343.15 K and 0% RH with 10 ml/min helium carrier gas, under the same conditions as in the experiment. The retention times were determined with a flame ionization detector (FID) and methane gas was used as a noninteracting molecule to determine the dead volume.

Prior to any surface-energy-related experiments, the BET specific surface areas of the two samples were first determined via physical adsorption of ethanol molecules by Elution Method using the retention time at the maximum FID signal. The particle-size distributions of POS and CBF were determined using a Beckman LS13320 laser particle-size analyzer. Vicat softening temperature tests were conducted with a ZWK1302-B Thermal Deformation/VICAT Temperature tester (MTS Systems (China) Co., Ltd.), 10 N loaded at a heating rate of 50 °C/h. Uniaxial tensile tests on LDPE/CBF composites were conducted according to the ASTM D638 standard. Izod impact tests were carried out on unnotched specimens according to the ASTM D256 standard. Particle size distribution was determined using a Beckman LS13320 laser particle size analyzer.

4. Results and discussion

4.1. Surface free energies

The γ_S^T , γ_S^D and γ_S^P profiles of CBF and POS obtained from iGC-SEA are illustrated in Fig. 3. It can be observed that the γ_S^D component of the samples contributed the major part (> 70%) to the γ_S^T . In addition, it displayed a decreasing trend with increasing surface coverage and the highest-energetic sites occupying approximately 2% of the fillers. However, this decreasing trend became insignificant when the surface coverage was larger than 4%. The difference in the measured absolute γ_S^D values at low and high coverage indicated significant heterogeneity among the surface free energy sites—those at highest-energetic sites had approximately 20% higher absolute values of the surface free energy compared to those of the lowest-energetic sites. For POS, the calculated γ_S^D fell into the range of 48.0–59.3 mJ/m² across the surface coverages measured; however, the CBF showed a lower range of

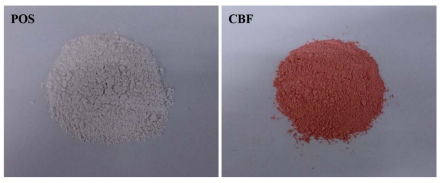


Fig. 1. Photographs of the POS and CBF powders.



Fig. 2. Specimens for the mechanical performance studies of LDPE with increasing amounts of CBF.

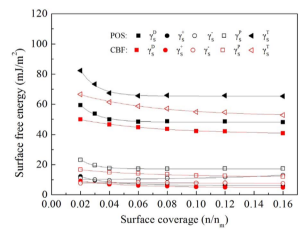


Fig. 3. Surface energy profiles for POS and CBF.

40.8–49.9 mJ/m². It is worth noting that the γ_S^D depends on the surface composition of the solid and the test temperature [24–26]. Schmitt et al. [27] reported that the modification of precipitated CaCO₃ with chemicals such as hydroxy acids or silanes could decrease the γ_S^D . The work of Papirer et al. [28] revealed a drastic decrease in surface free energy for CaCO₃ coated with stearic acid (44.4 mJ/m² for untreated and 29 mJ/m² for stearic acid-treated CaCO₃ at 70 °C). Jeong et al. [29] also reported a lower γ_S^D value for stearic acid-treated ground CaCO₃ (93.3 mJ/m² for untreatred and 34.8 mJ/m² for 1.5 wt% stearic acid-treated CaCO₃). In this work, a decrement of γ_S^D for CBF was also observed, which might be ascribed to its more uniform surface after loading with DR 28.

Compared to γ_S^D , the γ_S^P component evidently contributed less to γ_S^T : approximately 26.6% and 23.7% for POS and CBF, respectively,

implying a lower polarity and more hydrophobicity for both samples. γ_S^T can be determined by summing γ_S^D and γ_S^P according to the Fowkes theory [30], so that the higher γ_S^D and γ_S^P components for POS add up to a higher γ_S^T value. γ_S^T was also found to be significantly decreased at lower surface coverages for both samples. According to Eq. (1), W_C is equivalent to $2\gamma_S^T$. Thus, the W_C between filler particles was calculated as $130.60-164.82 \, \text{mJ/m}^2$ for POS and $105.57-133.11 \, \text{mJ/m}^2$ for CBF at various surface coverages. POS showed higher W_C values than CBF, by 12.8-19.2% across all coverages, indicating a higher tendency to aggregate. CBF showed comparatively lower W_C values,

which could reduce the filler particle-particle interactions, allowing its

4.2. Specific Gibbs free energy profiles

better dispersion in a polymer matrix.

The surface properties of fillers also depend on the ability to participate in specific interactions resulting from the presence of polar functional groups on the surface of the material. The ΔG^p profiles as a result of the interactions with four polar probes (toluene, acetone, acetonitrile and DCM) for CBF and POS are displayed in Fig. 4. ΔG^p changed as a function of surface coverage, further confirming the heterogeneous nature of the two samples. In Fig. 4, similar curves were generated for CBF and POS, although POS showed higher polar surface energies. The decreasing rank order for ΔG^p interactions was: acetonitrile > acetone > DCM > toluene, which was consistent with the order of solvent polarity [31,32]. Both samples showed a strong degree of interaction with all the polar probes, but predominantly with acetonitrile, and to a lesser extent with toluene. It is worth noting that CBF possessed a relatively lower polar surface free energy, indicating a more hydrophobic nature.

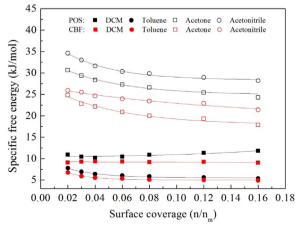


Fig. 4. Specific Gibbs free energy profiles of polar probes for POS and CBF.

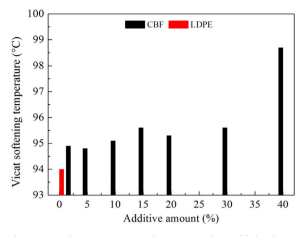


Fig. 5. Vicat softening temperature of neat LDPE and LDPE filled with CBF.

4.3. Vicat softening temperature of LDPE/CBF composites

Vicat softening temperature (VST), an important indicator for heat resistance, is the determination of the softening point for materials, such as plastics. As can be seen in Fig. 5, the VST exhibited an overall increasing trend with increasing CBF content. As compared with that of neat LDPE (94.0 °C), the VST of the composite materials increased by 0.9–4.7 °C, indicating an improvement in heat resistance. This may be attributed to that the fact that the combination of LDPE and CBF restricted the movement of LDPE molecular chains.

4.4. Mechanical properties of LDPE/CBF composites

The mechanical properties of LDPE/CBF composites are displayed in Fig. 6. For comparison, the neat LDPE is also included. It can be seen that the tensile strength of the specimens increased significantly as CBF increased. It increased from 12.65 to 14.13 MPa, with a filler loading increase of 5–40 wt%. The tensile strength is more dependent on the strength of filler-matrix adhesion [33]. Its increase might result from a

good adhesion between the CBF and the LDPE matrix, which created strong interface regions and benefited load transfer from the ductile matrix to the strong filler, and thus reinforced the composite [33].

The elongation at break of the LDPE composites decreased almost linearly, with a CBF loading greater than 5 wt%, because the composites became stiffer with increasing CBF content. As compared with the neat LDPE (171.6%), this value decreased from 166.4 to 75.2% as the CBF incorporation increased from 2 to 40 wt%. This substantial reduction was ascribed to the low elongation of the filler and the strong interaction with the polymer, restricting thus flow of polymer molecules [34].

In Fig. 6, a continuous increase in tensile elastic modulus is demonstrated with the incorporation of CBF. As compared with that for the neat LDPE (110.1 MPa), it increased from 137.2 to 313.3 MPa with filler loading increasing from 2 to 40 wt%. This apparent increase in Young's modulus can be attributed to increased filler-matrix interfacial interaction and enhanced load transmission from the polymer matrix to the reinforcement.

There was an overall increase in flexural strength and flexural modulus for LDPE composites filled with CBF with increases of 55.1% and 149.5% for flexural strength and flexural modulus, respectively, when the filler content increased from 2 to 40 wt%. This apparent increment could be attributed to the enhanced brittleness and stiffness of the composites.

The impact strength of a composite is influenced by many factors, including the toughness of the reinforcement, matrix fracture, the nature of the filler-matrix interfacial region, and frictional work involved in pulling out the filler from the matrix [35,36]. Among these, the nature of the interfacial region is significant and directly related to the toughness of the composite. For the CBF-filled composites, impact strength showed a small plateau followed by a significant deterioration. It increased from $65.3 \, \text{kJ/m}^2$ for neat LDPE to $69.5 \, \text{kJ/m}^2$ with a filler loading of 5 wt%; however, further increasing the incorporation content to 40 wt% led to a decrement of impact strength to $15.8 \, \text{kJ/m}^2$.

The evaluated mechanical properties showed that adding CBF could significantly increase the tensile strength, elastic modulus, flexural strength and flexural modulus of the LDPE composites. However, it

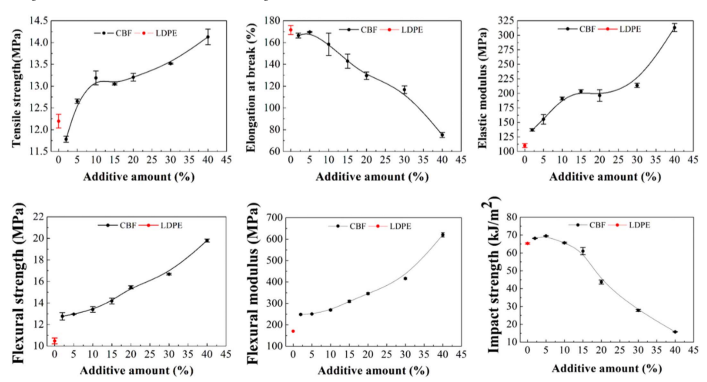


Fig. 6. Mechanical properties of neat LDPE and LDPE filled with CBF.

resulted in a decrease in impact strength and elongation at break. Therefore, the inclusion of CBF mainly played a reinforcing role. The optimal amount of CBF was determined to be 5–15 wt% with a good balance between toughness and stiffness in the LDPE composites. It is worth noting that lubricants [37] and compatilizers [38] can also affect the mechanical performance of LDPE composites. These factors will be considered in future work.

5. Conclusions

The γ_s^D profile for both CBF and POS showed that this component contributed the major part to the γ_s^T . The γ_s^D changed as a function of surface coverage, indicating that they were energetically fairly heterogeneous. The CBF also showed lower W_c , which could reduce the filler particle-particle interactions, allowing its better dispersion in the LDPE matrix. Adding CBF could significantly improve the heat resistance of LDPE and the VST increased by 0.9–4.7 °C. In addition, it could significantly increase the tensile strength, elastic modulus, flexural strength and flexural modulus of the LDPE composites. The absence of toxic metal coupled with excellent mechanical performance makes CBF an ideal candidate as a biofiller for the masterbatch industry, potentially solving the significant problem of high levels of toxic metals in toys.

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