Composites Science and Technology 87 (2013) 111-117

Contents lists available at ScienceDirect





Composites Science and Technology

journal homepage: www.elsevier.com/locate/compscitech

A cyanate ester/microcapsule system with low cure temperature and self-healing capacity



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ARTICLE INFO

Article history: Received 18 October 2012 Received in revised form 29 July 2013 Accepted 2 August 2013 Available online 14 August 2013

Keywords: A. Polymer–matrix composites (PMCs) A. Smart materials B. Fracture toughness B. Thermal properties C. Crack

1. Introduction

Cyanate ester (CE) resins are important high-temperature thermosetting polymers. The cured CE polymers exhibit high strength, low dielectric constants and dissipation factors, radar transparency, low water absorption, and superior metal adhesion. Because of these characteristics, they are well-suited for use in electrical components, aerospace structures, adhesives and functional materials [1–5]. However, the CE polymer is inherently brittle and prone to microcracking under service loads. When subjected to cyclic thermo-mechanical loading, the microcracks compromise the structural integrity of CE-based composites because of fiber/matrix debonding and inter-ply delamination [6–11].

Various approaches can be employed to improve the fracture toughness of CE formulations. The approaches include addition of high-thermoplastic tougheners [12–14], rubber inclusions [15–18], inorganic or organic fillers [19–23], thermosetting resins [24,25], and flexible molecules [26–28]. However, these methods are largely ineffective in restoring strength to cracked or damaged material. In such cases, localized damage is commonly repaired by plug/patch and scarf repairs, bolted plates and injecting resins

ABSTRACT

Microcapsules filled with epoxy resin were blended into cyanate ester resin to produce systems with low cure temperature and self-healing capacity. A 4,4'-diaminodiphenylsulfone (DDS) curing agent was adopted for the systems. The mechanical properties, thermal stability and self-healing ability of the system cured at low temperature were investigated. Cyanate ester systems with 2.5 wt% and 5.0 wt% microcapsules (MCs) demonstrated an 11~43% increase in fracture toughness (K_{IC}) relative to the neat resin cured at high temperature, and exhibited slightly lower thermal stability than the neat resin. The self-healing ability of cyanate ester with MCs was influenced by MC content and healing temperature, although for a formulation with 5.0 wt% MCs, recovery of 85% of the original fracture toughness was achieved by heat treatment of fractured samples for 1 h at 220 °C.

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[29–31], although these methods often cause new problems by altering the local stiffness, introducing local stress concentrations (e.g., from mechanical fasteners), and adding weight to the structure.

An alternative approach to repair involves the addition of polymeric microcapsules (MCs) containing a healing agent released when the MCs are ruptured by growing cracks. The addition of appropriate MCs can provide self-healing functionality and increase fracture toughness when the damage is caused by service loads [32–34]. Investigations of self-healing thermosetting materials have focused primarily on the effectiveness of self-healing in different polymers or composites fabricated at relatively low temperatures, and these studies have demonstrated benefits to both fracture toughness and the self-healing ability of epoxy resins [32–35].

In previous work, we developed systems comprised of CE resin filled with MCs, and showed that judicious additions of MCs improved mechanical properties [36]. However, MCs contents above 5.0 wt% significantly decreased the thermal decomposition temperature of the cured CE, an observation was attributed to the lower thermal stability of the MCs [36]. To design formulations for specific applications, chemists must first optimize the properties of the components of these smart materials. To date, few studies have evaluated the use of poly(urea–formaldehyde) (PUF) microcapsules filled with epoxy resins (PUF/EP MCs) as self-healing agents for high-temperature CE resins.

In this paper, we report the design and characterization of a high performance self-healing CE system with MCs (designated

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CE/MCs) that can be processed at low temperature (≤ 180 °C). Small amounts of MCs were blended into the CE resin to minimize the influence on the thermal stability of the CE matrix, and 4,4'diaminodiphenylsulfone (DDS) catalyst was used in the CE formulation to decrease the polymerization reaction temperature, thus ensuring a low cure temperature. The DDS catalyst also served as a curing agent for epoxy resin released from the MCs.

2. Experimental

2.1. Materials

For MC shell materials, a formulation of urea (U) and 37 wt% formaldehyde (F) was selected (Mauinckrodt Chemicals Advantor Performance Materials Inc.), and triethanolamine (TEA) was used to control the solution pH (Alfa Aesar). Analytical grade DDS was selected as catalyst (Alfa Aesar). Industrial grade CE resin (bisphenol A dicyanate (2,2'-bis(4-cyanatophenyl)isopropylidene, 98% pure, molecular weight: 278) were acquired (Zhejiang Shangyu Shengda Biochemical Co. Ltd., China), and bisphenol A epoxy resin (EP) was selected (Henkel). PUF/EP MCs were prepared following a specific protocol [37]. Fig. 1 shows the morphologies of PUF/EP MCs. The MCs are 5~150 µm, and the mean diameter is ~80 µm. The epoxy content of the MCs was ~82% (by weight), the thermal decomposition temperature (T_d : temperature at 5% weight loss) of the MCs was ~213 °C. Fig. 2 shows the chemical structures of the main materials.

2.2. Preparation of CE/MCs

To prepare the formulations, the CE resin was heated to 100 °C. After melting to a clear liquid, 1 wt% DDS was added. The temperature was raised to 120 °C and held for 30~40 min while stirring, then PUF/EP MCs (2.5 wt% and 5.0 wt%) were added and the mixture was stirred for an additional 10~20 min. Afterward, the mixture was poured into a pre-heated mold, degassed, and cured according to the following schedule: 130 °C/2 h + 150 °C/ 2 h + 180 °C/2 h. The cure schedule used for the neat CE was 150 °C/2 h + 180 °C/2 h + 200 °C/2 h, followed by a post-cure at 220 °C for 4 h. After curing, samples were cooled slowly to room temperature.

2.3. Characterization

Fourier transform infrared (FTIR) spectroscopy was performed by scanning KBr discs of the samples (Nicolet, ThermoScience),



Flexural properties were evaluated using a load frame (Instron 4045). The three-point bend fixture featured contact points with a 5 mm radius of curvature. Tests were conducted at 25 °C using a cross-head speed of 0.5 mm/min, and 10~15 specimens per composition were tested. The flexural strength, σ_f was calculated using following equation:

$$\sigma_f = \frac{3FS}{2wd^2} \tag{1}$$

where F is the applied force at fracture, S the span, and w and d are specimen width and depth.

Fracture toughness (K_{IC}) was measured using standard singleedge notched beam (SENB) specimens under mode I conditions. SENB specimens were loaded to fracture to determine the fracture toughness of the materials. The size and dimension of the specimen is shown in Fig. 3. The notch was introduced by first cutting a notch with a diamond saw, then generating a pre-crack by gently tapping a fresh razor blade at the base of the notch. The size of the original crack was 0.45W < a < 0.7W. The specimen was tested at a constant 0.5 mm/min displacement rate until fracture, and 10~15 samples for each composition were tested. Fracture toughness (K_{IC}) was calculated using:

$$K_{IC} = Y \frac{3PS\sqrt{a}}{2BW^2} \tag{2}$$

where *P* is the critical load, *B* the thickness, *W* the width, *S* the span, and *Y* is defined as:

$$Y = 1.93 - 3.07 \left(\frac{a}{W}\right) + 14.53 \left(\frac{a}{W}\right)^2 - 25.11 \left(\frac{a}{W}\right)^3 + 25.80 \left(\frac{a}{W}\right)^4$$
(3)

Fractured specimens were examined by light microscopy (Keyence VHX-600). Fractured specimens were rejoined using adhesive tape to ensure mutual contact of the crack surfaces, then heated according to the following schedule: (a) heat to 200 °C and hold for 1 h and (b) heat to 220 °C and hold for 1 h. After heat treatment, the K_{IC} of the specimen was measured using the same test procedure used for the control samples. Then the self-healing efficiency (η) was calculated using Eq. (4) [32]:

$$\eta = \frac{K_{IC \ healed}}{K_{IC \ origin}} \tag{4}$$

where $K_{IC \ origin}$ is the fracture toughness of the original specimen and $K_{IC \ healed}$ is the facture toughness of the specimen after healing.

The re-bonding area is calculated using an image processing code (MATLAB). The relative re-bonded area (RRBA) is taken as the fraction of the total available fracture surface that is bonded, and is defined as RRBA = A_b/A_t , where A_t is the total fracture area available for bonding and A_b is the bonded area.

Dynamic mechanical analysis (DMA) was performed using a single cantilever beam clamping setup (TA Q800 DMA) between 30 and 350 °C using a heating rate of 3 °C/min at 1 Hz. Sample dimensions were 35 mm \times 10 mm \times 2 mm. The glass transition temperature (T_g) was determined from the peak temperature in the tan δ -temperature plot.

The thermogravimetric (TGA) analyses were performed from 30 to 800 °C at a heating rate of 10 °C/min using nitrogen flowing at 50 ml/min (TA Instruments SDTQ500).



Fig. 1. Spherical PUF/EP MCs.



PUF

Fig. 2. The chemical structures of the main materials.



Fig. 3. Dimensions of SENB specimen.

3. Results and discussion

3.1. Mechanical properties of CE/MCs

The flexural strength and fracture toughness of CE/MCs systems are shown in Fig. 4. Compared to the neat CE cured at high temperature, the CE/MCs systems cured at low temperature show increased flexural strength and fracture toughness (K_{IC}). The strength and fracture toughness increase by 35% and 11% for the system with 2.5 wt% MCs, and by 50% and 43% for the system with 5.0 wt% MCs. The increases in mechanical properties can be understood from considerations of the modified chemistry, as explained below.

First, the DSC curve of the CE resin (Fig. 5a) shows that the introduction of DDS catalyst reduces the polymerization reaction peak temperature of neat CE from 300 °C to 180 °C, thus the amount of unreacted cyanate ester (—OCN) groups decreases at lower temperature, increasing the strength and toughness. Moreover, amine groups in DDS react with —OCN groups to form isoureas that contain the flexible chains [21,38,39], improving the impact strength. Secondly, because the PUF shell walls contain -OH and amine groups, the MCs catalyze the polymerization reaction of the CE resin [3,21,38–41] and reduce the unreacted -OCN groups, yielding the DSC curve of CE/PUF and CE/MCs system in Fig. 5a. The -OH and amine groups react with -OCN groups to form iminocarbonate and isoureas, respectively, which can improve the toughness.

The formation of iminocarbonate and isoureas is indicated by comparing the FTIR spectra of cured CE and the interfacial region between CE and MCs in cured CE/DDS/MCs (Fig. 5b). For all samples, the absorption peaks at 1564 cm⁻¹ and 1367 cm⁻¹ in the FTIR spectra are attributed to the triazine ring [3]. For the interfacial region between CE and MCs in cured CE/DDS/MCs, the absorption peaks between 3710 cm⁻¹ and 3090 cm⁻¹ are attributed to --NH and —OH, the absorption peak at 1690 cm^{-1} is assigned to the stretching vibration of C=NH in imidocarbonate and isourea [42], the peak at 1640 cm^{-1} is attributed to the isourea [21], and the absorption peak near at 1615 cm⁻¹ is attributed to imidocarbonate [40]. The epoxy resins released from MCs catalyze CE and react with -OCN group/triazine rings to form aliphatic oxazolidinone rings. This assertion is evidenced by comparing the DSC curves of CE, EP, CE/EP and CE/MCs (Fig. 5a), and by the absorption peak at 1752 cm⁻¹ in the FTIR spectrum of the interfacial region between CE and MCs in cured CE/DDS/MCs system (Fig. 5b) [3,42]. The oxazolidinone rings can increase fracture toughness [43,44].

Thirdly, MCs behave in a visco-elastic manner at small deformations and in a plastic manner at larger deformations [45]. They can also act as stress concentrations under triaxial loading conditions, resulting in localized shear yielding or microcracking in the matrix [46]. As microcracks propagate, MCs arrest propagation, rupture and debond, absorbing energy and stabilizing the crack. Fourthly, due to the reaction between —OH, amine, epoxy resin, and —OCN groups at the CE/MCs interface, the interfacial adhesion is strong, which improves the mechanical properties. Finally, because



Fig. 4. Flexural strength and fracture toughness of CE/MCs systems.



Fig. 5. DSC curves of the different CE systems and FTIR spectra of cured CE and the interfacial region between CE and MCs in cured CE/DDS/MCs system.

CE/MCs systems are cured at low temperature, thermal decomposition of matrix is avoided, then the mechanical property of the systems can be improved. In this study, the CE has the same structure as BADCy reported in Ref. [36], but the DSC curves reported in this work and in Ref. [36] are different, the phenomenon is caused by the different thermal history and mixing processes employed of the sample before DSC experiment, as well as a different batch of CE.

Fig. 6 shows images of the fracture surface of CE and CE/MCs system after fracture toughness measurements, along with FTIR spectrum of core materials of MCs. Compared to the neat CE, the

CE/MCs system exhibit more irregular fracture surface, which is consistent with the higher measured toughness values. Microcracks surrounded the MCs, and ruptured MCs are evident on fracture surfaces of CE/MCs samples, indicating enhanced fracture energy absorption. In addition, the accompanying FTIR spectrum of the released core materials from the region surrounding the fractured MCs shows an absorption peak characteristic of an epoxy group at 910 cm⁻¹ (Fig. 6c). Here, the released core material was collected with injection needle under a light microscope and diluted with acetone, then the dilute core material was applied to KBr disc. After the KBr disc dried at 100 °C for 20 min, it was ready for FTIR experiment.

3.2. Thermal stability of CE/MCs

Fig. 7 shows TGA curves of CE/MCs systems. The decomposition temperature (T_d) at 5% weight loss and the temperature of maximum rate of weight loss (T_{max}) for neat CE polymer are 433 °C and 443 °C, respectively. The T_d and T_{max} of CE with 2.5 wt% MCs are 430 °C and 441 °C, and the T_d and T_{max} of CE with 5.0 wt% MCs are 412 °C and 439 °C. In contrast, the T_d for both MC loadings is slightly lower than for the neat CE, while the T_{max} for both MC contents remains roughly unchanged. The decreased T_d for the CE/MCs system is attributed to the lower thermal properties of the MCs and the creation of the molecular networks such as isoureas, iminocarbonate and oxazolidinone rings (Fig. 5b).

3.3. Thermal dynamic mechanical properties of CE/MCs

Fig. 8 shows DMA curves of CE/MCs systems. The storage moduli in the low temperature range for CE/MCs systems are slightly higher than that of neat CE. As the temperature reaches 170 °C, the storage moduli of the CE/MCs demonstrate a significant drop because of the lower stiffness of the MCs and the low molecular weight epoxy resin core materials. In addition, the drop in storage modulus can also be attributed to the flexible chains in the new networks formed by -OH, amine groups, epoxy groups and -OCN [3,21,38–44], which have lower thermal stability. The two tan delta peaks (T_{g1} and T_{g2}) of CE/MCs systems marked in Fig. 8 indicate multiphase materials. The broadness of the relaxation peak indicates the complexity of the phase morphology of the material. The T_{g1} (at 296 °C, 301 °C and 288 °C) is strong and sharp and belongs to the transition of triazine rings. The T_{g1} values for the CE/MCs systems are close to the T_{g1} value for the neat CE. The small T_{g2} at 224 °C is attributed to the MCs and the formation of new networks such as iminocarbonate, isoureas and oxazolidinone rings at the CE/MCs interfaces (Fig. 5b).





Fig. 6. Fracture surfaces of CE and CE/MCs systems along with FTIR spectrum of core materials of MCs.



Fig. 7. TGA curves of CE/MCs systems.



Fig. 8. DMA curves of CE/MCs systems.

3.4. Self-healing ability of CE/MCs

Fig. 9 shows the dependence of self-healing efficiency (η) of healed CE/MCs systems on heating schedule. After heat treatment at 200 °C for 1 h, the CE with 2.5 wt% and 5.0 wt% MCs recover ~36% and ~43% of the original fracture toughness. When CE/MCs are healed at 220 °C for 1 h, CE with 2.5 wt% and 5.0 wt% MCs recover even more of the corresponding original fracture toughness, ~56% and ~85%, respectively.

The self-healing capacity of CE/MCs systems arises from the release of uncured epoxy from ruptured MCs, and the subsequent reaction with residual amine or amine derives within matrix to polymerize and re-bond the crack surfaces [47]. In particular, the released epoxy resin reacts with triazine rings in the matrix to form oxazolidinone rings [43,44]. The new network formation



Fig. 9. Dependence of self-healing efficiency (η) of healed CE/MCs systems on heating schedule.



(c) CE with 2.5wt% MCs

Fig. 10. Fractured surfaces of CE/MCs systems previously fractured and healed at 220 °C for 1 h, and FTIR spectrum of polymerized core materials.

significantly increases the bond strength and restores the CE mechanical properties. For a fixed MC loading, higher healing temperatures increase the self-healing efficiency η because of the more complete reaction of epoxy. Also, for a fixed healing temperature, higher MC contents lead to higher η , because the increased amount of epoxy resins leads to more extensive coverage and bonding of crack surfaces.

Fig. 10 shows the fractured surfaces of CE/MCs systems previously fractured and healed at 220 °C for 1 h and FTIR spectrum of polymerized core materials. Adhesion traces near ruptured MCs are visible on the fractured surface in Fig. 10a~c. Fig. 10a shows a microcrack that has intersected and ruptured a MC. The MC is surrounded by a roughened adhesion region caused by the spreading of epoxy resin that subsequently re-bonded to the opposing crack surface. Increasing the MC content provides more healing agent to rebond the crack surfaces (Fig. 10b). The measured values of the relative rebonded areas for CE with 2.5 wt% and 5.0 wt% MCs are 0.20 \pm 0.05 and 0.54 \pm 0.08, respectively. The latter value is slightly greater than twice the former value, an observation can be attributed to capillarity effects of interacting pads of resin in close proximity, resulting in coalescence and expansion.

The roughened peripheral regions around the MCs on the rebonded surface contain epoxy polymers, as confirmed by the disappearance of the epoxy group absorption peak at 910 cm^{-1} in the FTIR spectrum (Fig. 10c). The disappearance of the epoxy group results from the polymerization reaction of the released epoxy resins initiated by the —OH, amine group and triazine rings when heated.

4. Conclusions

Self-healing CE/MCs systems with low cure temperature were formulated and mechanical properties were measured before and after healing. The self-healing CE/MCs systems exhibited enhanced mechanical properties and self-healing capacity. The flexural strength of CE loaded with 5.0 wt% MCs was 50% greater than that of neat CE, and the fracture toughness was 43% greater. The CE/MCs systems showed slightly reduced thermal decomposition temperatures due to the MCs and theformation of new networks containing iminocarbonate, isoureas and oxazolidinone rings. When MCs ruptured and released epoxy resin core material to fill cracks, subsequent heating polymerized the epoxy and re-bonded the crack surfaces, thereby restoring as much as 85% of the original fracture toughness. The findings demonstrate the effectiveness of the MC approach for development of self-healing polymer materials, and suggest the potential for extension of a similar approach to fiber-reinforced CE composites.

Acknowledgements

The authors thank the National Natural Science Foundation of China (No. 51273135), the support of the Priority Academic Program Development of Jiangsu Higher Education Institutions (PAPD), and "Jiangsu Government Scholarship for Overseas Studies" (2011–2012). SN acknowledges support from the M.C. Gill Composites Center.

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