Publication Information: Hong, N., Zhan, J., Wang, X., Stec, A.A., Richard Hull, T., Ge, H., Xing, W., Song, L., Hu, Y., Enhanced mechanical, thermal and flame retardant properties by combining graphene nanosheets and metal hydroxide nanorods for Acrylonitrile-Butadiene- Styrene copolymer composite. Composites Part A Appl Sci Manuf 64, 203-210, (2014).

Enhanced mechanical, thermal and flame retardant properties by combining graphene nanosheets and metal hydroxide nanorods

for Acrylonitrile-Butadiene-Styrene copolymer composite

Ningning Hong¹, Jing Zhan¹, Xin Wang¹, Anna A. Stec², T. Richard Hull², Hua Ge¹, Weiyi Xing¹, Lei Song¹*, Yuan Hu¹*

- 1. State Key Laboratory of Fire Science, University of Science and Technology of China, Hefei, Anhui 230026, PR China
- 2. Centre for Fire and Hazards Science, University of Central Lancashire, Preston, PR1 2HE, UK

Abstract

Three metal hydroxide nanorods (MHR) with uniform diameters were synthesized, and then combined with graphene nanosheets (GNS) to prepare acrylonitrile-butadiene-styrene (ABS) copolymer composites. An excellent dispersion of exfoliated two-dimensional (2-D) GNS and 1-D MHR in the ABS matrix was achieved. The effects of combined GNS and MHR on the mechanical, thermal and flame retardant properties of the ABS composites were investigated. With the addition of 2 wt% GNS and 4 wt% Co(OH)₂, the tensile strength, bending strength and storage modulus of the ABS composites were increased by 45.1%, 40.5% and 42.3% respectively. The ABS/GNS/Co(OH)₂ ternary composite shows the lowest maximum weight loss rate and highest residue yield. Noticeable reduction in the flammability was achieved with the addition of GNS and Co(OH)₂, due to the formation of more continuous and compact charred layers that retarded the mass and heat transfer between the flame and the polymer matrix.

Keywords: A. Layered structures; A. Polymer-matrix composites (PMCs); B. Mechanical properties; B. Thermal properties.

^{*} Corresponding author. Tel/Fax: +86 551 63601664.

E-mail address: leisong@ustc.edu.cn (L. Song), yuanhu@ustc.edu.cn (Y. Hu)

Introduction

Polymer nanocomposites have aroused tremendous interest in material science because of their unexpected properties including superior mechanical, thermal, and electronic properties relative to the traditional microcomposites [1]. Among all the nano-additives, layered nanofillers such as silicate clay (MMT), layered double hydroxide (LDH), and layered salt phosphate, etc., have been proven to impart effective reinforcement and flame retardancy with polymers owing to their lamellar structure and high aspect ratio [2-4].

Recently, another two-dimensional (2-D) nanostructure, graphene nanosheets (GNS), comprised of sp² hybridized carbon, has been a focal point for research, due to its unique properties and potential applications. Due to the ultrahigh mechanical strength, specific surface area, and large aspect ratio, GNS are anticipated to exhibit better enhancement in mechanical strength and flame retardancy for polymer composites than MMT or LDH [5-7]. When GNS are uniformly dispersed in polymer matrices, the load transfer from the matrices to graphene is realized, which affords high mechanical strength for the polymer/GNS nanocomposites. GNS have been widely used to reinforce poly(vinyl alcohol) and impressive 150% and 1000% increases in tensile strength and Young's modulus respectively were achieved at only 1.8 vol% GNS loading [8]. The effect of GNS on the mechanical properties of thermoset epoxy resin was also investigated and a percolation threshold value of 0.6 wt% was observed [9]. According to recent reports, GNS show great potential as a flame retardant additive to endow polymers with high flame retarding. Pioneering work by Huang's group has demonstrated the effectiveness of GNS as flame retardant in poly(vinyl alcohol). With the incorporation of 3 wt% GNS, the peak heat release rate (PHRR) of the composites was reduced by 49.1% due to the formation of a compact and dense char [10]. The inclusion of GNS led to an increase in both the thermal stability and flame retardancy of polystyrene, with the optimal properties obtained at 5 wt% GNS content [11]. The flame retardant mechanism was attributed to the layered structures of GNS that acted as barriers to slow down the heat release.

Two important issues need to be addressed when both the mechanical and flammability properties of graphene-based composites are taken into account. Pristine GNS are apt to agglomerate or even restack in matrices due to the strong van der Waals force and π - π interactions between the graphene sheets [12, 13]. In addition, the improvement especially in flame retardancy, is only achieved at high GNS concentrations, which inevitably causes a reduction in the mechanical properties [10, 14].

To solve the above problems, many strategies have been adopted. The idea of using two or more components to achieve synergistic effect has already been explored before. In flame retardant fields, the combination of nanofillers with other flame retardants has been demonstrated to efficiently impact the mechanical and flame retardant performance of polymers [15-18]. Fang's group has revealed that the incorporation of 2-D clay or GNS and 1-D carbon nanotubes (CNT) gave rise to a sharp increase in the mechanical properties and obviously slowed down the whole combustion process [17, 18]. Uniform exfoliation of 2-D and 1-D nanofillers and the formation of a 3-D network structures were responsible for the

dramatic synergies. Very recently, the combination of transition metals Lewis acids and 2-D clay provided a promising method, which involved catalytically enhanced char formation and the formation of a surface-protective layer having a network structure [19, 20]. Systematic work done by Tang's group has demonstrated that dramatically improved fire retardancy of polyolefin was realized through promoting carbonization by the combination of 2-D clay with Ni₂O₃ [21, 22].

In this work, a new strategy is designed in order to simultaneously improve the mechanical properties and flame retardancy of acrylonitrile-butadiene-styrene (ABS) by combining 2-D GNS and 1-D metal hydroxide nanorods (MHR). The structures and morphologies of the GNS and MHR were characterized by X-ray diffraction (XRD) and transmission electron microscopy (TEM). The mechanical properties of the resultant ABS composites were measured by the tensile and bending testing, and the dynamic mechanical analysis (DMA). Also, the thermal degradation and the flame retardant behaviors of the resultant ABS composites were evaluated using thermogravimetric analysis (TGA) and cone calorimeter methods.

Experimental

2.1 Materials

Natural graphite (Spectrum Pure) and acetone were bought from Sinopharm Chemical Reagent Co., Ltd (Shanghai, China). Potassium permanganate (KMnO₄), sodium nitrate (NaNO₃), sulfuric acid (H₂SO₄, 98%), hydrogen peroxide (H₂O₂, 30%), hydrochloric acid (HCl, 37%) and hydrazine monohydrate (N₂H₄·H₂O) were bought from Guangfu Fine Chemial Research Institute (Tianjin, China). Cobalt chloride hexahydrate (CoCl₂·6H₂O), nickel sulfate hexahydate (NiSO₄·6H₂O), ferric nitrate nonahydrate (Fe(NO₃)₃·9H₂O), urea, sodium hydroxide (NaOH) and potassium hydroxide (KOH) were analytical grade and purchased from Sinopharm Chemical Reagent Co., Ltd (China). ABS (727) was supplied as pellets by Qimei New Material Co., Ltd. (China).

2.2 Synthesis of GNS and MHR

Graphite oxide (GO) was prepared from exfoliation of natural graphite by the modified Hummers' method [23]. GO was first subjected to ultrasonication in a 500 mL three-neck-flask to obtain a stable suspension (2 mg/mL). Then 12 mL N₂H₄·H₂O was added to the solution with magnetic stirring. The temperature was then raised to 100 °C and maintained for 2 h. During the reaction process, the yellow suspension gradually changed into a black precipitate. The mixture was filtered and rinsed with distilled water, and then dried at 80 °C.

Three different MHR were synthesized by a simple hydrothermal method. The molar ratio of starting ions, the temperature and the reaction time play a critical role in the morphologies of the resultant products. In a typical synthesis of $Co(OH)_2$, 5 mmol $CoCl_2 \cdot 6H_2O$ and 10 mmol urea were dissolved in 50 mL distilled water with magnetic stirring. After stirring for 30 min, the solution was sealed into a 60 mL Teflon-lined stainless steel autoclave, which was then maintained at 95 °C for 12 h [24]. For the synthesis of Ni(OH)_2, 16 mmol NiSO₄ \cdot 6H_2O and an

appropriate amount of NaOH were each dissolved in 20 mL deionized water. Afterwards, the latter solution was added dropwise to the former solution under vigorous stirring. The mixed solution was then transferred to an autoclave and kept in an oven maintained at 180 °C for 24 h [25]. The synthesis procedure of FeOOH was similar to that of Ni(OH)₂, using Fe(NO₃)₃·9H₂O (12 mmol) and KOH (48 mmol) as starting materials. The hydrothermal reaction was carrried out at 100 °C for 12 h. After being cooled to room temperature, the precipitates were filtered, washed with distilled water and ethanol, and then dried at 70 °C [26].

2.3 Preparation of ABS/GNS/MHR nanocomposites

The solubility of GNS in certain solvents provides a good approach to the preparation of polymer/GNS nanocomposites. GNS and/or MHR with certain proportions were dispersed to 300 mL acetone by sonication for 30 min. Then ABS pellets were added to the solutions by vigorous stirring. The resulting suspension was placed in an oil-bath at 70 °C to allow sufficient dissolution. The mixture was drop-cast into a Teflon tray and dried at 80 °C in a vacuum oven. Finally, the ABS nanocomposites were hot pressed at 205 °C under 10 MPa using a Shanghai Ximaweili press vulcanizer (China). The compositions of the sample are listed in Table 1.

Sample	Compositions					Tensile	Elongation	Bending	Bending
	ABS (wt%)	GNS (wt%)	Co(OH)2 (wt%)	Ni(OH) ₂ (wt%)	FeOOH (wt%)	Strength (MPa)	at break (%)	Strength (MPa)	Modulus (MPa)
ABS	100					28.8±0.9	28.0±2.0	39.0±1.8	1370±68
ABSG1	99	1				29.5±1.1	4.0±0.3	41.2±2.9	1579±75
ABSG2	98	2				40.7±2.0	3.1±0.2	44.0±2.1	2029±98
ABSCo4	96		4			39.1±1.6	1.1±0.1	45.5±1.8	1494±78
ABSG1Co4	95	1	4			43.2±2.3	5.9±0.4	48.4±2.1	2107±86
ABSG2Co4	94	2	4			41.8±1.7	3.7±0.3	54.8±3.3	2253±103
ABSG2Ni4	94	2		4		39.3±1.4	2.5±0.2	51.9±2.6	2193±118
ABSG2Fe4	94	2			4	40.5±1.6	5.6±0.4	42.7±2.3	2020±92

Table 1 Compositions of ABS nanocomposites and their corresponding mechanical properties.

2.4 Measurements and Characterization

X-ray diffraction (XRD) patterns were performed with a Japan Rigaku D/Max-Ra rotating anode X-ray diffractometer equipped with a Cu-K α tube and Ni filter (λ =0.1542 nm).

Transmission electron microscopy (TEM) was used to observe the morphology with an acceleration voltage of 200 kV. Specimens for TEM observation were prepared by dispersing the sample suspension onto a lacy carbon film supported by a Cu grid.

Scanning electron microscopy (SEM) was performed on a XL-30 scanning electron microscope with the acceleration voltage of 20 kV. The fracture surfaces and char residues of samples were previously coated with a conductive layer of gold before analysis.

The tensile strength of the nanocomposites was measured with an electronic universal testing instrument (MTS System Co., Ltd, China) at a crosshead speed of 50 mm/min. The elongation was in the range of error, so only the tensile strength was reported.

The bending properties of the nanocomposites were tested with an electronic universal testing instrument (MTS System Co., Ltd, China) at a crosshead speed of 2 mm/min according to GB/T8812.1-2007.

Dynamic mechanical analysis (DMA) was performed on the Perkin-Elmer Pyris Diamond DMA from 20 °C to 170 °C at a heating rate of 5 °C/min, at a frequency of 10 Hz in the tensile configuration.

The thermogravimetric analysis (TGA) was carried out on the TGA Q5000 IR thermogravimetric analyzer (TA instruments) using a heating rate of 20 °C/min in a nitrogen or air atmosphere.

The combustion property of the nanocomposites was investigated using a FTT cone calorimeter according to ISO 5660 standard. Every sample with a size of $100*100*3 \text{ mm}^3$ was heated at a heat flux of 35 kW/m².

Results and Discussion

3.1 Characterization of GNS and MHR

The morphologies of the as-synthesized GNS and MHR were observed by TEM. Fig. 1a shows the existence of thin carbon sheets of size up to 1 μ m. GNS sheets seem to associate closely with each other to form an overlapped structure. As shown in Fig. 1b, the sample consists of a large quantity of wire-like Co(OH)₂ with a uniform diameter of 40-60 nm and length of several micrometers. From Fig. 1c, the rod-like morphology is widely observed with uniform diameter and well dispersion. Based on the mutability of Ni(OH)₂, the morphology of the final product has been successfully controlled by tuning the key parameters of molar ratio, temperature and time. FeOOH exhibits almost rod-like morphology with diameter of 80-100 nm, indicating high yield and good uniformity were achieved by the hydrothermal method (Fig. 1d).

The crystalline phases of the GO, GNS and MHR were characterized by XRD as presented in Fig. 2a. After chemical reduction, a broad (002) diffraction halo at about 24.5° is observed for the GNS, corresponding to a d-spacing of 0.35 nm. This result indicates the GO is well reduced and the as-reduced GNS sheets appear to partially restack [27]. In the case of $Co(OH)_2$ nanorods, all the diffraction peaks can be well indexed, which are consistent with the result in the literature [24]. No peaks for other phases are found in the as-synthesized nanorods. Ni(OH)₂ obtained herein has very high peak intensity, suggesting that the nanomaterials are well crystallized [25]. Pure FeOOH with high crystallinity has been successfully synthesized, as evidenced by the diffraction peaks attributed to orthorhombic α -FeOOH [26]. Fig. 2b shows the TGA curves of GNS and MHR in air. There are three major weight loss steps for GO as the temperature increases. The first step before 100 °C is ascribed to the removal of absorbed water; the second one occurs at around 200 °C due to the decomposition of labile oxygen functional groups; almost all the sample is oxidized in the last step when the temperature raised to 800 °C [28]. GNS presents a gradual weight loss with an increase in temperature, which can be attributed to the removal of water molecules and labile oxygen-containing groups on the GNS sheets. A large weight loss of over 80 wt% occurs in the temperature range of 550-800 °C due to the oxidation of the graphene skeleton in air [27]. There is an abrupt weight loss between 250 and 350 °C due to removal of structural water and transformation of Co(OH)₂ to Co₃O₄. The temperature of maximum decomposition rate (T_{max}) of 296 °C is observed in the DTG curve. It is observed that the temperature for 5% weight loss ($T_{-5\%}$) and T_{max1} of Ni(OH)₂ shift to higher temperatures of 274.8 and 341.0 °C respectively. The weight loss in the first step can be attributed to the loss of water and the phase transformation of Ni(OH)₂ to NiO. The decomposition behaviour of FeOOH is similar to that of $Co(OH)_2$, exhibiting the T_{-5%} (261 °C), T_{max} (274 °C), and 87 wt% residue at 800 °C.



Fig. 1. TEM images of the as-prepared GNS (a), Co(OH)₂ (b), Ni(OH)₂ (c) and FeOOH (d).



Fig. 2. XRD patterns (a) and TGA curves (b) of the GNS, Co(OH)₂, Ni(OH)₂ and FeOOH.

3.2 Dispersion of GNS and MHR

The dispersion state of nanofillers in a matrix is critical to the mechanical, thermal, and other properties of the composite. To investigate the morphological features of GNS and/or MHR in the ABS matrix, SEM and TEM were used. Pure ABS represents a smooth fracture surface, indicative of a typical brittle failure feature (not shown). With the introduction of 2 wt% GNS, the surface of ABS composite becomes rough (Fig. 3a). In addition, graphene aggregates are sometimes seen on the surface due to the strong van der Waals forces between GNS sheets (marked by black circles). As revealed in Fig. 3b, the fractured surface contains randomly dispersed nanorods without visible agglomerates, suggesting good dispersion of Co(OH)₂ in the matrix (marked by black rectangles). The ABSG2Co4 composite shows good dispersion and homogeneity on the fractured surface (Fig. 3c). 2-D GNS is embedded in and 1-D Co(OH)₂ is held tightly by the matrix, suggesting the formation of 3-D network structure. In the TEM image of ABSG2Co4 (Fig. 3d), the gray background represents the ABS matrix, while the black lines and flakes correspond to different orientations of GNS sheets (marked by black circles and rectangles). As shown in Fig. 3e and f, ABS composites with $Ni(OH)_2$ and FeOOH exhibit almost the similar features except for rougher surfaces. No GNS aggregates could be observed in all the ternary composite, indicating that the presence of MHR could prevent the GNS from re-stacking and thus have a good dispersion in the matrix.



Fig. 3. SEM images of the fractured surfaces of ABSG2 (a), ABSCo4 (b), ABSG2Co4 (c), ABSG2Ni4 (e) and ABSG2Fe4 (f); TEM image of ABSG2Co4 (d).

3.3 Mechanical properties of ABS nanocomposites

The effects of GNS and MHR on the mechanical properties of the ABS composites are studied, with the results summarized in Table 1. Because of the high mechanical modulus of GNS, the tensile strength (TS) increases from 28.84 MPa for pure ABS to 40.71 MPa for ABSG2, but its elongation at break largely reduces. The incorporation of 4 wt% Co(OH)₂ results in a 35.6% increase in TS, compared to pure ABS owing to the homogeneous dispersion. When the combination of GNS with $Co(OH)_2$ is applied, further enhancement in the TS of the ternary composite are observed. The TS of ABS composites reaches a maximum value of 43.16 MPa when 1 wt% GNS and 4 wt% Co(OH)₂ are added, corresponding to an increase of 49.7%. Such enhancement is attributed to the 3-D network structure formed by homogeneous dispersion of GNS and Co(OH)₂, which facilitates the successful load transfer from the matrix to the nanofillers. The enhancement in the TS is more significant than the values reported by the previous literatures [29, 30]. According to Tang's work, the incorporation of carbon nanotubes slightly improves the TS of linear low density polyethylene/Ni₂O₃ compared with those of binary composites [29]. An increase in mechanical strength was reported for epoxy/graphene nanoplate/carbon nanotube composites, where only 14.5% increase in the TS was obtained compared to epoxy/graphene nanoplate composites [30]. Compared to the ABSG2Co4, ABSG2Ni4 and ABSG2Fe4 show slightly reduced TS of 39.29 and 40.46 MPa, respectively, due to the decreased aspect ratio of Ni(OH)2 and FeOOH nanorods. The formation of a network structure due to the homogeneous dispersion of two types of nanofillers is responsible for the improvement in the mechanical properties of the composites.

From Table 1, it can be also seen that the incorporation of GNS significantly increases bending strength (BS) and bending modulus (BM) of the ABS composites, due to the high intrinsic strength and modulus of GNS [31]. The inclusion of 4 wt% Co(OH)₂ also slightly

improves the ultimate BS and BM of the ABS composite owing to the reinforced effect derived from the large aspect ratio of the nanorod. With the simultaneous addition of GNS and Co(OH)₂, it is found that both BS and BM are further increased to 54.8 MPa and 2253 MPa, respectively. In comparison with Ni(OH)₂ and FeOOH, Co(OH)₂ shows more obvious synergistic effect with GNS on improving the BS and BM of the composites.

The mechanical reinforcement is also reflected on the dynamic mechanical property. The storage modulus (E') as a function of temperature curves were plotted in Fig. 4 for ABS and its composites. The incorporation of GNS into ABS gives rise to a noticeable increase in E' of the composites due to the rigid nature of GNS. For instance, E' value at 30 °C in the glassy region is 1796 MPa, yielding 28.7% increment compared to pure ABS (1396 MPa). The incorporation of 4 wt% Co(OH)₂ gives rise to a reduced E' value of the composite. A combination effect of GNS and Co(OH)₂ on the E' value of the composites in the glassy region is clearly observed. The E' values of the ABSG1Co4 and ABSG2Co4 composites at 30 °C are 2110 and 1987 MPa, respectively, which are 51.1% and 42.3% higher than that of pure ABS. This phenomenon can be attributed to the better dispersion and improved interfacial interaction between GNS and ABS matrix due to the existence of Co(OH)₂ nanorods. By comparison, the reinforcement effects are not observed in ABSG2Ni4 and ABSG2Fe4 composites, because Co(OH)₂ nanorod with higher aspect ratio results in stronger reinforcement.



Fig. 4. Storage modulus (E0) versus temperature plots of ABS and its nanocomposites.

3.4 Thermal stability of ABS nanocomposites

The thermal stability of ABS and its nanocomposites was characterized by TGA under both nitrogen and air. The TGA (a, c) and DTG (b, d) curves are plotted in Fig. 5, and according results are summarized in Table 2. ABS degrades with a large single peak at 382.6 °C with little residue left, implying that pure ABS itself has weak charring ability. The addition of GNS does not visibly affect the thermal stability of the resin except for improving the residue to 3.5 wt%. However, the addition of Co(OH)₂ plays an important role on the

thermal stability of the ABS composite. The T_{-5%} and T_{max} of the ABSCo4 composite are found to be lower than that of pure ABS, which could be attributed to the dehydration reaction of Co(OH)₂ at low temperature. It is surprising that the residue is significantly increased to 13.7 wt%. Moreover, the rate of weight loss is greatly reduced with the incorporation of Co(OH)₂. The overall increased amount of residue may be due to the carbonization effect of Co(OH)₂ that promotes the pyrolytic species of ABS into solid char [19]. The simultaneous addition of GNS and Co(OH)₂ further increases the amount of residue (15.7 wt%) and decreases the mass loss rate (33.9%). The combination effect could be observed in the systems containing graphene and metal hydroxides, where the T_{max} and residue amount are greatly increased due to the physical barrier of graphene and the catalytic carbonization of Co(OH)₂. As far as the residue yield is concerned, the ABSG2Ni4 and ABSG2Fe4 composites are not so efficient as ABSG2Co4, which is probably assigned to that the addition of Ni(OH)₂ or FeOOH can not form an effective barrier due to lower aspect ratio.



Fig. 5. TG and DTG curves of ABS and its nanocomposites in nitrogen (a and b) and air (c and d) atmosphere. Insets in (a) and (c) are the enlarged areas of weight loss.

Sample-	T₋ _{5%} (ºC)		T _{max1} (≌C)		T _{max2} (≌C)		Residue at 700 ºC (wt%)	
	N ₂	Air	N ₂	Air	N ₂	Air	N ₂	Air
ABS	382.6	368.8	429.1	422.6	-	586.3	2.3	1.1
ABSG1	382.0	367.2	427.2	422.0	-	563.4	2.8	0.8
ABSG2	382.9	367.5	429.9	420.9	-	548.7	3.5	1.1
ABSCo4	352.9	349.0	409.0	395.9	-	514.1	13.7	3.9
ABSG1Co4	353.7	344.2	406.6	394.2	-	509.1	14.5	3.3
ABSG2Co4	357.4	352.7	407.2	397.3	-	517.0	15.7	5.6
ABSG2Ni4	368.2	352.0	425.9	415.9	-	545.9	9.8	3.6
ABSG2Fe4	378.7	349.5	421.9	408.0	-	491.3	12.5	3.1

Table 2 TGA data of ABS and its nanocomposites in both nitrogen and air atmosphere.

Thermal degradation of the samples in air is significantly different from that in nitrogen. ABS exhibits a two-step weight loss: the first step (T_{max1}=422.6 °C) is attributed to the molecular chain scission and the second one (T_{max2}=586.3 °C) is due to the oxidation of residual species. The $T_{-5\%}$ and T_{max1} of the ABS/GNS composites are seldom changed, while the T_{max2} is visibly reduced, inferring the poor thermal oxidation resistance of GNS in air. Compared to pure ABS, the T_{-5%} and T_{max} of ABSCo4 composite are largely shifted to lower temperatures, especially for the decrease of T_{max2} (about 70 °C). Interestingly, the rate of weight loss is significantly reduced by 40.7% and the final residue is increased to 3.9 wt%. In this sense, cobalt oxide promotes char formation at an earlier temperature that decreases the evolution of pyrolytic gases through enhancing the char formation. When GNS are further added, though the $T_{-5\%}$ and T_{max} of the ternary composite are largely decreased, the rate of mass loss is reduced most. It is worth noting that the combination of GNS and Co(OH)₂ leads to the highest amount of residue (5.6 wt%). In the enlarged area, the ternary composite displays a higher weight of residue than the individually filled binary composites with GNS or Co(OH)₂. Therefore there is a combined effect in terms of thermal oxidative degradation. Similarly, the thermal stability of ABSG2N4 and ABSG2Fe4 is inferior to that of ABSG2Co4, but much superior to that of pure ABS.

3.5 Flame retardancy of ABS nanocomposites

The flammability of polymeric materials is commonly characterized by cone calorimeter, which is very effective to evaluate their fire properties under real world fire conditions. Fig. 6 shows the HRR versus time curves for ABS and its nanocomposites. Important parameters including time to ignition (t_{ign}) , time to PHRR (t_{PHRR}) , total heat release (THR), mean specific extinction area (mSEA), mean mass loss rate (mMLR) and fire growth index (FGI), are

recorded in Table 3. Pristine ABS is easy to be ignited (t_{ign} =49 s) and a rather sharp PHRR value (843 KW/m²) yields after 173 s. In comparison with ABS, the PHRR of the composite is slightly increased when GNS is added. The t_{ign} is shorten due to the high thermal conductivity of GNS [11]. Incorporating GNS shows little effect on the THR, mSEA and mMLR values. The inclusion of 4 wt% $Co(OH)_2$ gives rise to a 9.8% reduction in PHRR as compared to pure ABS. However, it exhibits weak effect on reducing THR and mSEA, but the mMLR is suppressed to an extent. For the trinary ABS nanocomposites, the PHRR displays a 30.5 % reduction when 2 wt% GNS and 4 wt% Co(OH)₂ are combined. Moreover, there is an obvious synergistic effect between GNS and Co(OH)₂ on reducing fire hazards of ABS composite, including reduced mSEA and mMLR, as compared to the binary ABSG2 and ABSCo4 composites. It could be explained by the combination effect of the physical barrier of GNS that prevents the escape of volatile gases and the cartalytic carbonization of $Co(OH)_2$ that favors the char formation [29]. The flame retardancy of ABSG2Ni4 and ABSG2Fe4 is also improved with PHRR to be 638 and 658 KW/m^2 , respectively. The FGI, defined as the ratio of PHRR and t_{PHRR}, could be used to measure the overall fire behaviors of ABS composites. The smallest value of 3.18 is observed in the ABSG2Co4 composites, demonstrating highest flame retardancy.



Fig. 6. Heat release rate versus time curves for ABS and its nanocomposites.

							-
Sample	t _{ign} (s)	t _{PHRR} (s)	PHRR (KW/m²)	THR (MJ/m²)	mSEA (m²/Kg)	mMLR (Kg/Kg)	FGI (KW/s·m²)
ABS	49	173	843	84.08	1175	0.116	4.87
ABSG1	35	164	878	85.63	1151	0.167	5.35
ABSG2	34	171	855	89.00	1096	0.114	5.00
ABSCo4	46	175	760	93.99	1235	0.082	4.34
ABSG1Co4	43	173	634	79.30	1300	0.076	3.66
ABSG2Co4	45	184	586	88.49	709	0.067	3.18
ABSG2Ni4	35	183	638	90.35	976	0.083	3.49
ABSG2Fe4	39	188	658	92.29	1107	0.111	3.50

Table 3 Cone calorimeter data of ABS and its nanocomposites at a heat flux of 35 KW/m².

The flame retardant efficiency has a direct relationship with the structure and yield of the char formed. Fig. 7 shows the digital photographs of the residual chars from the combustion of ABS composites. Pure ABS is almost completely burnt, and little char is obtained (Fig. 7a). The residue of ABSG2 is very loose as revealed in Fig. 7b. Layered GNS could somewhat prevent the evolution of gaseous products, but its efficiency is not high. The color of the char of ABSCo4 composite turns into black, indicating that cobalt compound promotes char formation (Fig. 7c). In the case of ABSG2Co4, the char looks more continuous and compact, which is likely to reduce the HRR (Fig.7d). Promoting char formation of ABS itself during combustion will be beneficial for reducing its flammability [32]. The combination effect of the physical barriers of GNS and the catalytic carbonization of Co(OH)₂ is responsible for the improved char residue. In Fig.7e, the morphology of the final residues of ABSG2Ni4 seems to be similar with that of ABSG2Co4. For ABSG2Fe4 composite, the rufous and island-like residue indicates that the main component is Fe₂O₃ from the decomposition of FeOOH rather than amorphous carbon (Fig.7f).



Fig. 7. Digital photographs of the char residues of ABS and its nanocomposites.

Conclusions

The Co(OH)₂, Ni(OH)₂ and FeOOH nanorods were successfully synthesized by a facile hydrothermal process, and their structures were characterized by TEM, XRD, FTIR and TGA. The 1-D MHR and their combination with 2-D GNS were used to prepare ABS composites. Stacking of individual 2-D GNS sheets is somewhat hindered by introducing 1-D MHR nanorods, resulting in the formation of 3-D nanohybrids network in the ABS matrix. A combined effect exists between GNS and Co(OH)₂ in the mechanical properties, thermal stability and flame retardancy of ABS composites. When 2 wt% GNS and 4 wt% Co(OH)₂ were combined, the resulting composite exhibits significant mechanical improvements. The thermal stability of ABS is also obviously enhanced with the addition of GNS and Co(OH)₂. The combination of GNS and Co(OH)₂ could confer better flame retardancy on ABS relative to all other components. A more continual, compact charred layers were formed in the ABS/GNS/Co(OH)₂ composite. This improvement is partially attributed to the formed 3-D network due to the good dispersion of GNS and Co(OH)₂, and partially to the carbonization of degradation products of ABS catalyzed by Co(OH)₂.

Acknowledgements

The work was financially supported by the National Key Technology R&D Program (2013BAJ01B05), the National Natural Science Foundation of China (No.21374111), the China Postdoctoral Science Special Foundation (2013T60621) and the Institute of Nanotechnology and Bioengineering of the University of Central Lancashire in the UK.

References

[1] Kumar AP, Depan D, Tomer NS, Singh RP. Nanoscale particles for polymer degradation and stabilization-Trends and future perspectives. Prog Polym Sci 2009;34(6): 479-515.

[2] Bitinis N, Hernandez M, Verdejo R, Kenny JM, Lopez-Manchado MA. Recent Advances in Clay/Polymer Nanocomposites. Adv Mater 2011;23(44):5229-36.

[3] Leroux F, Besse JP. Polymer interleaved layered double hydroxide: A new emerging class of nanocomposites. Chem Mater 2001;13(10):3507-15.

[4] Alongi J, Frache A. Flame retardancy properties of alpha-zirconium phosphate based composites. Polym Degrad Stabil 2010;95(9):1928-33.

[5] Kotov NA. Materials science: Carbon sheet solutions. Nature 2006;442(7100):254-5.

[6] Zaman I, Kuan HC, Dai JF, Kawashima N, Michelmore A, Sovi A, Dong SY, Luong L, Ma J. From carbon nanotubes and silicate layers to graphene platelets for polymer nanocomposites. Nanoscale 2012;4(15):4578-86.

[7] Putz KW, Compton OC, Palmeri MJ, Nguyen ST, Brinson LC. High-Nanofiller-Content

Graphene Oxide-Polymer Nanocomposites via Vacuum-Assisted Self-Assembly. Adv Funct Mater 2010;20(19):3322-9.

[8] Zhao X, Zhang QH, Chen DJ, Lu P. Enhanced Mechanical Properties of Graphene-Based Poly(vinyl alcohol) Composites. Macromolecules 2010;43(5):2357-63.

[9] Fang M, Zhang Z, Li JF, Zhang HD, Lu HB, Yang YL. Constructing hierarchically structured interphases for strong and tough epoxy nanocomposites by amine-rich graphene surfaces. J Mater Chem 2010;20(43):9635-43.

[10] Huang GB, Gao JR, Wang X, Liang HD, Ge CH. How can graphene reduce the flammability of polymer nanocomposites? Mater Lett 2012;66(1):187-9.

[11] Han YQ, Wu Y, Shen MX, Huang XL, Zhu JJ, Zhang XG. Preparation and properties of polystyrene nanocomposites with graphite oxide and graphene as flame retardants. J Mater Sci 2013;48(12):4214-22.

[12] Lin Y, Jin J, Song M. Preparation and characterisation of covalent polymer functionalized graphene oxide. J Mater Chem 2011;21(10):3455-61.

[13] Cao YW, Lai ZL, Feng JC, Wu PY. Graphene oxide sheets covalently functionalized with block copolymers via click chemistry as reinforcing fillers. J Mater Chem 2011;21(48):9271-8.

[14] Hong NN, Pan Y, Zhan J, Wang BB, Zhou KQ, Song L, et al. Fabrication of graphene/Ni-Ce mixed oxide with excellent performance for reducing fire hazard of polypropylene. Rsc Adv 2013;3(37):16440-8.

[15] Yang F, Nelson GL. Combination effect of nanoparticles with flame retardants on the flammability of nanocomposites. Polym Degrad Stabil 2011;96(3):270-6.

[16] Yu AP, Ramesh P, Sun XB, Bekyarova E, Itkis ME, Haddon RC. Enhanced Thermal Conductivity in a Hybrid Graphite Nanoplatelet-Carbon Nanotube Filler for Epoxy Composites. Adv Mater 2008;20(24):4740-4.

[17] Ma HY, Tong LF, Xu ZB, Fang ZP. Synergistic effect of carbon nanotube and clay for improving the flame retardancy of ABS resin. Nanotechnology 2007;18(37): 375602.

[18] Song PA, Liu LN, Fu SY, Yu YM, Jin CD, Wu Q, Zhang Y, Li Q. Striking multiple synergies created by combining reduced graphene oxides and carbon nanotubes for polymer nanocomposites. Nanotechnology 2013;24(12):125704.

[19] Jang J, Kim J, Bae JY. Effects of Lewis acid-type transition metal chloride additives on the thermal degradation of ABS. Polym Degrad Stabil 2005;88(2):324-32.

[20] Kashiwagi T, Danyus R, Liu MF, Zammarano M, Shields JR. Enhancement of char formation of polymer nanocomposites using a catalyst. Polym Degrad Stabil 2009;94(11):2028-35.

[21] Tang T, Chen XC, Chen H, Meng XY, Jiang ZW, Bi WG. Catalyzing carbonization of polypropylene itself by supported nickel catalyst during combustion of polypropylene/clay nanocomposite for improving fire retardancy. Chem Mater 2005;17(11):2799-802.

[22] Song RJ, Jiang ZW, Yu HO, Liu J, Zhang ZJ, Wang QW, Tang T. Strengthening carbon

deposition of polyolefin using combined catalyst as a general method for improving fire retardancy. Macromol Rapid Comm 2008;29(10):789-93.

[23] Park S, An JH, Piner RD, Jung I, Yang DX, Velamakanni A, Nguyen ST, Ruoff RS. Aqueous Suspension and Characterization of Chemically Modified Graphene Sheets. Chem Mater 2008;20(21):6592-4.

[24] Wang ZH, Chen XY, Zhang M, Qian YT. Synthesis of Co_3O_4 nanorod bunches from a single precursor $CO(CO_3)(_{0.35})CI_{0.20}(OH)_{1.10}$. Solid State Sci 2005;7(1):13-5.

[25] Dong LH, Chu Y, Sun WD. Controllable synthesis of nickel hydroxide and porous nickel oxide nanostructures with different morphologies. Chem-Eur J 2008;14(16):5064-72.

[26] Ou P, Xu G, Ren ZH, Hou XH, Han GR. Hydrothermal synthesis and characterization of uniform alpha-FeOOH nanowires in high yield. Mater Lett 2008;62(6-7):914-7.

[27] He GY, Chen HQ, Zhu JW, Bei FL, Sun XQ, Wang X. Synthesis and characterization of graphene paper with controllable properties via chemical reduction. J Mater Chem 2011;21(38):14631-8.

[28] Dubin S, Gilje S, Wang K, Tung VC, Cha K, Hall AS, Farrar J, Varshneya R, Yang Y, Kaner RB. A One-Step, Solvothermal Reduction Method for Producing Reduced Graphene Oxide Dispersions in Organic Solvents. Acs Nano 2010;4(7):3845-52.

 [29] Yu H, Liu J, Wang Z, Jiang ZW, Tang T. Combination of Carbon Nanotubes with Ni₂O₃ for Simultaneously Improving the Flame Retardancy and Mechanical Properties of Polyethylene.
J Phys Chem C 2009;113(30):13092-7.

[30] Yang SY, Lin WN, Huang YL, Tien HW, Wang JY, Ma CCM, Li SM, Wang YS. Synergetic effects of graphene platelets and carbon nanotubes on the mechanical and thermal properties of epoxy composites. Carbon 2011;49(3):793-803.

[31] Farahani RD, Dalir H, Aissa B, Khakani MAE, Lévesque M, Therriault D. Micro-infiltration of three-dimensional porous networks with carbon nanotube-based nanocomposite for material design. Compos Part A-Appl S 2011;(42):1910-9.

[32] Song PA, Xu LH, Guo ZH, Zhang Y, Fang ZP. Flame-retardant-wrapped carbon nanotubes for simultaneously improving the flame retardancy and mechanical properties of polypropylene. J Mater Chem 2008;(18):5083-91.