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Fabrication of Ce-doped MnO₂ decorated graphene sheets for fire safety applications of epoxy composites: flame retardancy, smoke suppression and mechanism

Shu-Dong Jiang,a Zhi-Man Bai,ab Gang Tang,a Lei Song,a Anna A. Stec,c T. Richard Hull,c Jing Zhan*a and Yuan Hu*ab

Ce-doped MnO₂–graphene hybrid sheets were fabricated by utilizing an electrostatic interaction between Ce-doped MnO₂ and graphene sheets. The hybrid material was analyzed by a series of characterization methods. Subsequently, the Ce-doped MnO₂–graphene hybrid sheet was introduced into an epoxy resin, and the fire hazard behaviors of the epoxy nanocomposite were investigated. The results from thermogravimetric analysis exhibited that the incorporation of 2.0 wt% of Ce-doped MnO₂–graphene sheets clearly improved the thermal stability and char residue of the epoxy matrix. In addition, the addition of Ce–MnO₂–graphene hybrid sheets imparted excellent flame retardant properties to an epoxy matrix, as shown by the dramatically reduced peak heat release rate and total heat release value obtained from a cone calorimeter. The results of thermogravimetric analysis/infrared spectrometry, cone calorimetry and steady state tube furnace tests showed that the amount of organic volatiles and toxic CO from epoxy decomposition were significantly suppressed after incorporating Ce–MnO₂–graphene sheets, implying that this hybrid material has reduced fire hazards. A plausible flame-retardant mechanism was hypothesized on the basis of the characterization of char residues and direct pyrolysis-mass spectrometry analysis: during the combustion, Ce–MnO₂, as a solid acid, results in the formation of pyrolysis products with lower carbon numbers. Graphene sheets play the role of a physical barrier that can absorb the degraded products, thereby extend their contact time with the metal oxides catalyst, and then promote their propagate on the graphene sheets; meanwhile pyrolysis fragments with lower carbon numbers can be easily catalyzed in the presence of Ce–MnO₂. The notable reduction in the fire hazards was mainly attributed to the synergistic action between the physical barrier effect of graphene and the catalytic effect of Ce–MnO₂.

1. Introduction

Epoxy resin (EP) is one of the most widely exploited reactive polymeric resins in the laminating, adhesive, coating and casting fields due to its high tensile strength and modulus, low shrinkage when cured, high adhesion property, excellent dimensional stability and chemical and corrosion resistance.1,2 However, the high flammability and large yield of smoke and toxic gases during combustion significantly restrict their application in some areas. Recently, nanocomposite technology has brought a revolutionary advancement in the flame-retardant research community. For example, the incorporation of nano-fillers, such as layered silicates and layered double hydroxides (LDHs), can efficiently improve the fire resistance of polymer nanocomposites.1–3 However, a key problem that was realized in this system is that the large addition of fillers required (>20 wt%) resulted in the deterioration of other properties, which is undesirable for fabricating high performance materials.

Graphene, a two-dimensional (2D) form of carbon, has become a 'rising star' because of its large specific surface area, extraordinarily high electrical and superior mechanical strength.4–6 Very recently, similar to some other members of the family of carbon-based materials, such as expanded graphite, graphite oxide and carbon nanotubes, graphene has attracted interests in scientific field as a flame retardant additive.7–8 Kim et al.9 demonstrated that graphene is actually thermally stable even after exposing it to a flame, clearly
indicating the high intrinsic flame resistance of graphene. In addition, there are some other reports regarding the application of graphene as a flame retardant additive in polymer composites. However, bare graphene was used as a flame retardant in previous reports, such that a high loading is required to achieve a good flame retardant effect. Therefore, the use of graphene alone as the flame retardant still remains a challenge.

As an important functional metal oxide, manganese oxide has received considerable interest due to its distinctive physical and chemical properties, which lead to its application in catalysis, adsorption, and flame retardation. It has also been found that compounds containing cerium also have an unusually wonderful effect of flame-reduction in polymers. Recently, intensive studies have focused on combining two or more components, which can present a synergistic effect and may offer an unexpected flame resistance to polymers. Tang et al. demonstrated a novel catalytic method to endow polypropylene with obviously enhanced fire resistance in the presence of clay and a supported nickel catalyst. Yu et al. showed a synergism between multiwalled carbon nanotubes and Ni2O3 on the flame retardancy of polyethylene. Levchik et al. observed that the integration of Sb2O3 and ZnS generated a synergistic effect for the flame retardance and smoke suppression of poly(vinyl chloride). Inspired by these, this study aims to improve the fire hazards of EP by combining graphene sheets and Ce–MnO2.

In this work, a hybrid of Ce–MnO2 decorated graphene sheets (Ce–MnO2–GNS) were synthesized by co-assembly between positively charged Ce–MnO2 nanoparticles and negatively charged graphene sheets (GNS). GNS have a negative charge nature due to the presence of a small amount of carboxyl, epoxy and hydroxyl groups. In addition, modified Ce–MnO2 with NH2 groups exhibited the positive charge nature. The self-assembly of charged nanomaterials via electrostatic interactions is a controllable route for generating hybrid materials in an aqueous environment. Thus, the self-assembly of GNS and Ce–MnO2 could simply be carried out by utilizing the electrostatic interactions between the negative charge of GNS and the positive charge of modified Ce–MnO2. Subsequently, a Ce–MnO2–GNS hybrid was added into EP for the investigation of reduced fire hazards behavior. The synergistic effects between Ce–MnO2 and GNS for thermal stability, fire resistance and smoke suppression properties of the as-prepared nanocomposite were systematically investigated by thermogravimetric analysis (TGA), cone calorimetry, steady state tube furnace (SSTF) and thermogravimetric analysis/infrared spectrometry (TG-IR). Interestingly, it was found that a number of properties of the resultant nanocomposite could be remarkably enhanced. Meanwhile, the flame-retardant mechanism was studied by the analysis of the char residue and pyrolysis fragments. Such an aqueous synthetic strategy may be potentially applicable to the fabrication of other inorganic materials–GNS hybrids, which have potential applications in flame retardant field.

2. Experimental section

2.1. Materials

Graphite powder, concentrated sulfuric acid (98%), sodium nitrate, KMnO4, 30% H2O2 solution, hydrochloric acid, hydrazine hydrate (80%) and Ce(NO3)3·6H2O, HNO3 were purchased from Sinopharm Chemical Reagent Co. Ltd. (Shanghai, China). Aminopropyltriethoxysilane (APTES, 99%) was obtained from Aldrich (America). All the starting chemicals were of analytical grade and were used without further purification. Deionized water was used as the solvent.

2.2. Synthesis of GNS

Graphite oxide (GO) was first synthesized from graphite powder by the Hummers method, and then the aqueous GO dispersion was chemically reduced by hydrazine to generate a GNS, as described elsewhere. The obtained GNS was centrifuged, washed with absolute ethanol and then dried under vacuum. The yield of the final GNS was about 70%.

2.3. Synthesis of Ce–MnO2

Typically, 0.003 mol of KMnO4 and 0.00085 mol of Ce(NO3)3·6H2O were dissolved in 35 mL of deionized water, followed by the addition of 2 mL concentrated HNO3 (16 mol L−1) under magnetic stirring. The solution was transferred to a Teflon-lined autoclave (50 mL), maintained at 140 °C for 3 h, and then finally cooled to room temperature. After washing with 30 mL of distilled water and 30 mL of ethanol three times, brown black precipitates were obtained, which were then dried at 60 °C overnight. The yield of the final Ce–MnO2 was 85%.

2.4. Electrostatic assembly of Ce–MnO2–GNS hybrid

The Ce–MnO2 nanoparticles were first reacted with APTES to modify their surface with NH2 groups. The self-assembly of GNS and Ce–MnO2 was carried out by electrostatic interactions. Briefly, 0.09 g of modified Ce–MnO2 and 0.009 g GNS were added to 60 mL of H2O under ultrasonic agitation for 5 h. The Ce–MnO2–GNS hybrid was obtained after centrifugation and washing with 50 mL of water, and then dried in air at 60 °C. The yield of the final Ce–MnO2–GNS hybrid was about 90%.

2.5. Preparation of Ce–MnO2–GNS–EP composite

Typically, the preparation of epoxy composite with 2 wt% Ce–MnO2–GNS hybrid was as follows: Ce–MnO2–GNS (0.2 g) hybrid was dispersed into tetrahydrofuran and ultrasonically mixed until the Ce–MnO2–GNS was completely dispersed to form a black suspension. Then, epoxy resin (8.2 g) and 4,4'-diaminodiphenyl-methane (DDM, 1.6 g) were introduced into the abovementioned suspension and stirred until homogeneous mixtures were obtained. The mixtures were heated in a vacuum oven at about 60 °C to remove excess solvent. Subsequently, the samples were cured at 100 °C for 2 h and post cured at 150 °C for 2 h. After curing, the samples were cooled to room
temperature. For other samples containing neat EP, Ce–MnO$_2$–EP and GNS–EP, similar synthetic procedures were deployed except that the additives were varied.

2.6. Characterization

The powder X-ray diffraction (XRD) patterns of the as-synthesized samples were recorded with a Japan MapAHF X-ray diffractometer equipped with graphite-monochromatized Cu K$_\alpha$ irradiation ($\lambda = 0.154056$ nm), employing a scanning rate of 0.02° s$^{-1}$ in the 2θ range of 10–60°. Fourier transform infrared (FTIR) spectrum analyses were operated on samples pelletized with KBr powders in the range of 4000–400 cm$^{-1}$ using an IR Fourier transform spectrophotometer (Nicolet, ZOSX). Raman spectroscopy measurements were carried out at room temperature with a SPEX-1403 laser Raman spectrometer (SPEX Co, USA) with excitation provided in the backscattering geometry by a 514.5 nm argon laser line. The microstructures of Ce–MnO$_2$–GNS were observed by a JEOL JSM-2010 field-emission scanning electron microscope (SEM). Transmission electron microscopy (TEM) images were obtained on a Hitachi model H-800 transmission electron microscope with an accelerating voltage of 200 kV. Prior to observation, Ce–MnO$_2$–GNS were dispersed in deionized water followed by ultrasonication for 30 min. The homogeneous mixtures were dripped on carbon-coated copper grids. Ce–MnO$_2$–GNS–EP nanocomposite was cut into ultrathin sections using a CM1900 microtome (Leica, Wetzlar, Germany). The ultrathin sections were transferred from liquid nitrogen to carbon-coated copper grids and then observed by TEM. Energy-dispersive X-ray spectroscopy (EDS) analysis was obtained with an EDAX detector installed on the same TEM, and a copper grid was used. TGA of samples were carried out with a Q5000 thermal analyzer (TA Co., USA) from 50 °C to 700 °C at a heating rate of 20 °C min$^{-1}$ in an air atmosphere. X-ray photoelectron spectroscopy (XPS) spectra were recorded on an ESCALAB 250 X-ray photoelectron spectrometer employing a monochromatic Al K$_\alpha$ X-ray source. TG-IR was performed using the TGA Q5000 IR thermogravimetric analyzer which was coupled with the Nicolet 6700 FT-IR spectrophotometer via a stainless steel transfer pipe. Direct pyrolysis–mass spectrometry (DP-MS) analysis was performed with a Micromass GCT-MS spectrometer using the standard direct insertion probe for solid polymer materials at a heating rate of 15 °C min$^{-1}$ in the range of 30–700 °C. The cone calorimeter test was carried out following the procedures given in ISO5660. Square specimens (100 × 100 × 3 mm$^3$) were irradiated at a heat flux of 35 kW m$^{-2}$, corresponding to a mild fire scenario. The fire toxicity was assessed using a SSTF (ISO TS 19700), as shown in Scheme 1. An amount of sample (about 20 g) in the form of granules or pellets, which was spread evenly along the furnace boat was introduced into a tube furnace at a constant rate. A current of air was passed through the furnace over the specimen to support combustion. The effluent was expelled from the tube furnace into a mixing chamber, where carbon monoxide concentration and smoke density could be measured.

3. Results and discussion

3.1. Characterization of Ce–MnO$_2$–GNS hybrid

The composition and phase purity of GO, GNS, Ce–MnO$_2$ and Ce–MnO$_2$–GNS were first examined by powder XRD, as depicted in Fig. 1A. For pure Ce–MnO$_2$, four relatively strong peaks appeared at 2θ of around 22.1, 37.4, 42.7, and 56.7°, which are assigned to (120), (031), (300), and (160) reflections of the orthorhombic γ–MnO$_2$, respectively, with lattice constants of $a = 6.3600$ Å and $c = 4.0900$ Å (JCPDS Card no. 14-0644). No signal for the phases of GNS (002) or GO (001) could be detected in the Ce–MnO$_2$–GNS hybrid. The relatively high content and good crystallinity of Ce–MnO$_2$ in the hybrid gave strong diffraction peaks, covering the diffraction of the GNS. Fig. 1B shows the EDS spectrum of Ce–MnO$_2$, in which the signals for Mn and O with the atomic ratio of approximately 1 : 2 are clearly observed, indicating the chemical composition of MnO$_2$. Cu and C elements come from the carbon-coated copper grid, which is used as the support for the characterization of the Ce–MnO$_2$–GNS hybrid. In addition, very small amounts of Ce are detected, indicating that Ce exists in the MnO$_2$ structure.

The morphology and size of the Ce–MnO$_2$–GNS hybrid were characterized by TEM and SEM (Fig. 2). As can be observed from Fig. 2A–D, Ce–MnO$_2$ particles with different sizes were deposited on the GNS, and the GNS showed a representative wrinkle and ripped shape. The TEM image of Ce–MnO$_2$–GNS–EP composite is also provided (Fig. 2E), from which one can see the Ce–MnO$_2$ and GNS was still held together in the EP matrix.

![Scheme 1](Image)

Scheme 1 Illustration of the steady state tube furnace.

![Fig. 1](Image)

Fig. 1 Typical XRD patterns of GO, GNS and Ce–MnO$_2$–GNS hybrids (A) and EDS spectrum (B) of the Ce–MnO$_2$–GNS hybrid.
The FTIR spectra of GO, GNS and Ce–MnO₂–GNS in the range of 4000–400 cm⁻¹ are shown in Fig. 3. Almost all the characteristic peaks of GO disappear for GNS after chemical reduction, including the absorption peaks O–H deformation vibrations of tertiary C–OH (1410 cm⁻¹), O–H deformation vibrations of COOH groups (1620 cm⁻¹), and C=O stretching vibrations of COOH groups (1720 cm⁻¹).³⁰,³¹ In the case of Ce–MnO₂, the typical characteristic peak of lower wavenumber at 574 cm⁻¹ can be clearly observed, which is commonly attributed to the Mn–O vibrations of the [MnO₆] octahedral.³²,³³ After the electrostatic assembly of GNS and Ce–MnO₂, compared with GNS, the bands assigned to the Mn–O stretching vibration at 574 cm⁻¹ still exist,³²,³³ and several additional absorption bands at 1040, 1128, and 1620 cm⁻¹ were present, attributed to the Si–O–Si, Si–OMn, and N–H stretching or bending vibrations, respectively.³⁷ It implies that Ce–MnO₂ nanoparticles are successfully loaded on the large surface of GNS during the electrostatic assembly process.

Raman scattering spectroscopy is an effective method to examine the microstructure of carbonaceous materials because of its sensitivity toward these materials. Fig. 4 shows the Raman spectra of Ce–MnO₂, GO, GNS and Ce–MnO₂–GNS. The Raman spectrum of the Ce–MnO₂ has a distinct sharp peak located at 636 cm⁻¹, which can be attributed to the Mn–O vibrations perpendicular to the direction of the MnO₆ octahedral double chains of Ce–MnO₂.³⁴ As shown in Fig. 4, GO exhibits a Raman shift at about 1593 (G band), corresponding to the E²g mode of graphite, which is related to the vibration of the sp²-bonded carbon atoms in a two-dimensional hexagonal lattice. Another dominant peak at about 1357 cm⁻¹ (D band) is related to the defects and disorder in the hexagonal graphitic layers.³⁵ The intensity ratio (I_D/I_G) of the D to the G bands of the GO is about 1.30. Moreover, it is observed that the I_D/I_G of GNS increased to 1.69. After electrostatic interaction between Ce–MnO₂ and GNS, Ce–MnO₂–GNS shows a higher I_D/I_G ratio value than GNS due to the decrease in the ordered graphitic structure after the attachment of Ce–MnO₂. In addition, the peak at about 636 cm⁻¹ corresponding to the Mn–O stretching mode is still observed in the Raman spectrum of Ce–MnO₂–GNS. The results of XRD, EDS, FTIR and Raman spectra further indicate that the hybrid structure composed of GNS and Ce–MnO₂ was successfully synthesized.³⁶,³⁷

XPS analysis was used to investigate the surface composition and oxidation state of the Ce–MnO₂–GNS hybrid. In Fig. 5A, it can be seen that the product contains C, O, Si, N, Mn and Ce elements. The high resolution C1s peak of the Ce–MnO₂–GNS hybrid can be deconvoluted into four components corresponding to carbon atoms in different functional groups: the graphitic C–C group (284.6 eV), C–O–C group (286.3 eV), C–OH group (286.1 eV), and C–N group (288.3 eV).
group (285.3 eV) and carboxyl C=O group (288.4 eV). Two peaks corresponding to Mn2p3/2 and Mn2p1/2 are located at 642.2 eV and 654.4 eV, respectively. The Ce3d spectrum of the Ce–MnO2–GNS hybrid shown in Fig. 5 is composed of individual overlapping peaks, which are resolved into two sets of spin–orbital multiplets 3d5/2–3d3/2. The feature of v, v, v and u, u, u, corresponding to the Ce in 4+ oxidation state, while the presence of v and u demonstrates the presence of Ce in 3+ oxidation state. In this work, the peaks at 882.5, 888.4 and 898.5 eV are assigned to the v, v and v of Ce3d5/2, and the peaks at 901.2, 906.9 and 916.3 eV are observed for u, u and u of Ce3d3/2. The presence of Ce3+ in the hybrid is confirmed by the appearance of weak signals at 886.7 (v) and 904.8 eV (u).39,40

TGA measurements were used to investigate the influence of Ce–MnO2 on the thermal stability of GNS. Fig. 6 gives the TGA curves of Ce–MnO2, GNS and Ce–MnO2–GNS hybrid performed up to 800 °C under air at a heating rate of 20 °C min⁻¹. Ce–MnO2 displays a maximum weight loss of about 4% around 587 °C, corresponding to the loss of oxygen from MnO2 lattice and resulting in the phase transformation to Mn2O3. Pure GNS shows a consecutive mass loss with a weight loss of about 97.5 wt%, which should be attributed to the removal of carbon skeleton by burning of GNS. In contrast, the resulting Ce–MnO2–GNS hybrid exhibits remarkably different thermogravimetric behavior with about 52 wt% of residue at 800 °C. These results demonstrate that the combination of Ce–MnO2 with GNS significantly improves the residue yield of GNS.

3.2. Thermal and fire properties of EP and its nanocomposites

The thermal stability of the EP and its nanocomposites was evaluated by TGA, and derivative thermogravimetric (DTG) curves were generated. As shown in Fig. 7, all the nanocomposites presented degradation behaviors similar to that of pure EP, and the main difference is degradation temperatures. The thermal degradation process of pure EP mainly consists of two stages based on the TGA profile, which mainly correspond to the decomposition of the macromolecular chains in the temperature range of 350–480 °C and further thermal oxidation of char residue beyond 500 °C. The T₅% and T₅₀% are defined as the temperatures where 5% and 50% weight losses occur, while T₃₇₄% is defined as the maximum decomposition temperature. The data are listed in Table 1. In comparison with neat EP, the T₅%, T₅₀% and T₃₇₄% of Ce–MnO2–EP nanocomposite were decreased because of the catalytic degradation of Ce–MnO2. After introducing GNS into the EP matrix, the T₅%, T₅₀% and T₃₇₄% were slightly lower than that of the pure EP, which is attributed to the high heat conductivity of GNS. However, it is found that the Ce–MnO2–GNS–EP nanocomposite demonstrates improved thermal stability with 4.6 °C, 8.8 °C and 8.9 °C increments for T₅%, T₅₀% and T₃₇₄%, respectively, as compared to those of pure EP. As far as the char yield is concerned, the char residue at 700 °C is significantly increased.

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Table 1  TGA data for EP and its nanocomposites in air

<table>
<thead>
<tr>
<th>Sample</th>
<th>T₅% (°C)</th>
<th>T₅₀% (°C)</th>
<th>T₃₇₄% (°C)</th>
<th>Char residue at 700°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>EP</td>
<td>369.6</td>
<td>422.7</td>
<td>380.3</td>
<td>0.6</td>
</tr>
<tr>
<td>Ce–MnO2–EP</td>
<td>335.8</td>
<td>384.9</td>
<td>362.2</td>
<td>3.0</td>
</tr>
<tr>
<td>GNS–EP</td>
<td>356.6</td>
<td>414.7</td>
<td>374.6</td>
<td>0.4</td>
</tr>
<tr>
<td>Ce–MnO2–GNS–EP</td>
<td>374.2</td>
<td>431.5</td>
<td>389.2</td>
<td>3.4</td>
</tr>
</tbody>
</table>

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Fig. 5 XPS spectra of the Ce–MnO2–GNS hybrid (A), C1s (B), Mn2p (C) and Ce3d (D).

Fig. 6 TGA thermograms of GNS, Ce–MnO2 and Ce–MnO2–GNS in an air atmosphere.

Fig. 7 TGA and DTA curves of pure EP, GNS–EP, Ce–MnO2–EP, Ce–MnO2–GNS–EP.
when Ce–MnO$_2$ or Ce–MnO$_2$–GNS was added due to the catalytic carbonization of Ce–MnO$_2$. The improved char yield provides a protective shield to mass and heat transfer between air and polymeric materials, and reduces the heat release rate during combustion. From the DTG curves (Fig. 7), it can be clearly seen that the addition of Ce–MnO$_2$–GNS or GNS notably decreases the maximum mass loss rate (the peak of DTG curves) compared to that of pure EP, implying the mass barrier effect of GNS.

Cone calorimetry is a widely used method for measuring the flammability of various materials in real-world fire. The heat release rate (HRR) and peak heat release rate (THR) curves for EP and its nanocomposites are shown in Fig. 8A and B, and some important parameters obtained from the cone calorimeter tests, such as the peak heat release rate (pHRR), THR, total smoke release (TSR) and the average mass loss rate (AMLR), are also tabulated in Table 2. It is shown that neat EP burns extremely rapidly after ignition and the pHRR value reaches 1653 kW m$^{-2}$. GNS is usually used to impart flame retardant properties to polymers due to its unique 2D nanosheet structure. As expected, incorporating GNS into EP decreases pHRR to 1156 kW m$^{-2}$. For Ce–MnO$_2$–EP, the pHRR decreases to 920 kW m$^{-2}$. In addition, the pHRR of Ce–MnO$_2$–GNS–EP shows further reduction compared with that of both GNS–EP and Ce–MnO$_2$–EP. In addition, the THR values of GNS–EP, Ce–MnO$_2$–EP and Ce–MnO$_2$–GNS–EP are significantly reduced by 17%, 26% and 35%, respectively, compared to that of pristine EP. Fig. 8C shows the digital photographs of the residual chars after cone calorimeter tests of EP and its nanocomposites. As can be seen, neat EP almost does not form a char while the GNS–EP and Ce–MnO$_2$–EP nanocomposites form char with many holes and cracks. The addition of Ce–MnO$_2$–GNS leads to the increase in char yield and the formation of a continuous and compact char layer, which are good in agreement with the TGA results. The continuous and compact char surfaces are good barriers to protect the underlying polymers and inhibit the exchange of degradation products, combustible gases and oxygen. The results of flammability exhibit that Ce–MnO$_2$–GNS–EP nanocomposite show the best flame retardancy among EP nanocomposites. By combining TGA and cone calorimetry results, it is reasonably believed that some epoxy chains participated in the carbonization process due to the barrier effect of GNS and the catalysis of Ce–MnO$_2$, and more and better chars are formed, which decrease the flammability of EP nanocomposites.

Table 2  Cone calorimetry data of EP, GNS–EP, Ce–MnO$_2$–EP and Ce–MnO$_2$–GNS–EP composites

<table>
<thead>
<tr>
<th>Sample</th>
<th>pHRR (kW m$^{-2}$)</th>
<th>THR (MJ m$^{-2}$)</th>
<th>TSR (m$^2$ m$^{-2}$)</th>
<th>AMLR (g s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>EP</td>
<td>1653 ± 58</td>
<td>129.9 ± 3.4</td>
<td>3075 ± 55</td>
<td>0.249 ± 0.01</td>
</tr>
<tr>
<td>GNS–EP</td>
<td>1156 ± 30</td>
<td>107.8 ± 2.2</td>
<td>2666 ± 28</td>
<td>0.175 ± 0.007</td>
</tr>
<tr>
<td>Ce–MnO$_2$–EP</td>
<td>920 ± 27</td>
<td>96.7 ± 1.8</td>
<td>2222 ± 19</td>
<td>0.124 ± 0.005</td>
</tr>
<tr>
<td>Ce–MnO$_2$–GNS–EP</td>
<td>765 ± 25</td>
<td>83.8 ± 1.5</td>
<td>1809 ± 17</td>
<td>0.110 ± 0.003</td>
</tr>
</tbody>
</table>
3.3. Smoke suppression properties

EP with aromatic and aliphatic chains, which decompose to yield predominantly aromatic hydrocarbons, produces very large amounts of smoke and toxic gases. It is well-known that in most cases the real killer in fires is not the heat of the fire itself but the smoke and toxic gases. Therefore, reduction of smoke and toxic gases (especially carbon monoxide) will be beneficial to fire rescue when a fire accident occurs. Fig. 9A presents CO rate during the thermal decomposition of EP and its nanocomposite are shown in Fig. 10. Some small molecular gaseous decomposition products evolved from EP and its nanocomposite are identified unambiguously by characteristic strong FTIR signals, such as -C=H groups for allyl alcohol, acetone, ethers and various hydrocarbons (3100–2800 cm⁻¹), CO (2360 cm⁻¹), CO (2180 cm⁻¹) and aromatic compounds (1605, 1510 and 1460 cm⁻¹). Typical thermal degradation products of EP nanocomposites that are similar to those of pure EP are observed. To provide a clear comparison, the intensities of typical gaseous volatiles for pure EP and its nanocomposite are presented in Fig. 11. With the incorporation of inorganic additives, the intensity of gas emission is shifted to lower values. Moreover, it can be obviously observed that the addition of Ce–MnO₂–GNS significantly decreased the evolution of all the evolved compounds and showed the best smoke suppression performance among all the nanocomposites.

The possible reasons for the reduced smoke and gases toxicity of Ce–MnO₂–GNS–EP are illustrated as follows. From the TGA results (Fig. 7), the synergistic effect between GNS and Ce–MnO₂ can promote char formation in the EP nanocomposite, which implies that there is more compact char residue formed on the surface of the sample with Ce–MnO₂–GNS. In addition, the digital photographs of the residual chars (Fig. 8C) further confirm this point. The compact char residue and the physical barrier effect of GNS can restrain the release of combustible gases such that the released flammable gases can completely burn, which leads to the slow CO production rate. Metal catalysis has also been found to be an effective way to eliminate volatile organic compounds and toxic
gases during combustion reactions. As an important functional metal oxide, MnO₂ has exciting catalytic effects due to its distinctive physical and chemical properties. Zhang et al. reported that manganese oxides are promising catalyst in CO oxidation. Li et al. demonstrated that the MnO₂ microspheres showed a higher catalytic ability for the catalytic elimination of benzene. Moreover, the presence of cerium can improve the oxygen storage capacity of the catalyst through the transformation between Ce⁴⁺ and Ce³⁺ under controlled conditions, which promotes the catalytic ability of the catalyst. Therefore, it is reasonable to believe that the catalytic effect of Ce–MnO₂ plays an important role in the elimination of volatile organic compounds and toxic gases. The reduction of the volatile organic compounds and toxic gases in the smoke and gases will be beneficial to fire rescue when an accident occurs.

3.4. Flame retardant mechanism

It is well-known that layered nanofillers, including layered silicate and LDHs, play a role in the condensed phase by acting as "char reinforcers". When a condensed phase action is the main mechanism for flame retardant additives, its efficiency strongly depends on the structure and composition of the char during combustion. Therefore, investigating the properties and the microstructure of the resultant char layers provides an insight into understanding how the flame retardant additives act in the condensed phase. As indicated in Fig. 12A, the residual char of pure EP presents a rough and loose char layer with a mass of cracks and holes dispersed on the surface. Moreover, the char displays a porous and incompact surface with high-magnification SEM imaging (Fig. 12B). However, when 2 wt% of Ce–MnO₂–GNS is added into EP, the cracks and holes are reduced (Fig. 12C and D) and the porous and incompact surface changes into a more compact surface. More detailed information regarding the char surface is obtained from the inset of Fig. 12D; it can be observed that the final char contains a mass of nano-sized spheres, and the char nanospheres were connected to each other to form a "macromolecular-chain" structure. The dense char layer can lower the efficiency of heat and volatiles transference because of the strong hindering effect, and can provide better flame shielding for the underlying material during combustion. The char residue of Ce–MnO₂–GNS–EP nanocomposite after cone calorimetry tests. As can be observed, the Raman spectra of the four samples exhibit a similar shape, with two peaks at 1593 cm⁻¹ and 1358 cm⁻¹. The characteristic peak at 1593 cm⁻¹ is called G band, corresponding to the first order scattering of the E₂g mode of hexagonal graphite, while the other is called D band, arising from the vibration of carbon atoms with dangling bonds in the plane terminations of disordered graphite. The graphitization degree of the char can be estimated by I_D/I_G, where I_D and I_G are the integrated intensities of the D and G bands, respectively. Basically, the lower the ratio of I_D/I_G, the better is the structure of the char. The peak intensity ratio I_D/I_G of GNS–EP and Ce–MnO₂–EP is slightly lower than that of EP, indicating the improvement of the graphitized carbons in the

![Fig. 12 SEM images (A–D) of the surface morphology of the residue of EP and Ce–MnO₂–GNS–EP and XRD pattern (E) of the char residue of Ce–MnO₂–GNS–EP nanocomposite after cone calorimetry tests.](image)

![Fig. 13 Raman curves of the residue for EP and its nanocomposites.](image)
residual char. Furthermore, incorporating the Ce–MnO₂–GNS hybrid into EP remarkably decreases the intensity ratio \(I_D/I_G\) compared to that of EP, suggesting a synergistic effect for char formation between Ce–MnO₂ and GNS. The high graphitization degree of the char is known to be compact and efficient in terms of thermal insulation, which provides a protective shield that results in a decrease in heat and mass transfers between the flame and the material.¹⁹

As is well-known, the pyrolysis products during combustion play a very important role in the flame-retardant performance of materials. To explore the flame-retardant mechanism of the EP nanocomposites, the pyrolysis products during combustion were studied by the DP-MS technique. Fig. 14 depicts the total ion current (TIC) chromatogram of EP and its nanocomposite, and EI-MS spectra corresponding to the TIC peaks with the maximum intensity. The pyrolysis fragment ions with a molecular weight of \(x\) are labelled as \(x_M\) and are identified in Fig. 15. For pure EP, a strong peak at 255 \(m/z\) corresponds to \(C_{17}H_{19}O_2\). In addition, the peaks at 269 \(m/z\), 213 \(m/z\), 197 \(m/z\) and 135 \(m/z\) can be assigned to the pyrolysis products of \(C_{17}H_{21}N_2O\), \(C_{14}H_{13}O_2\), \(C_{15}H_{18}\), and \(C_{9}H_{11}O\), respectively. Moreover, some fragment ions recombine to form some new products under high temperature environments; for instance, the recombination between benzene and aniline generates carbazole (166 \(m/z\)), and the carbazole and benzene reconstitute to produce a few polycyclic aromatic hydrocarbons (497 \(m/z\)). The spectrum of the GNS–EP composite shows similar pyrolysis products as that of EP. In contrast, the main fractions in the degradation products of the Ce–MnO₂–GNS–EP or Ce–MnO₂ includes \(C_{12}H_{13}O_2\) (213 \(m/z\)), which is with lower carbon numbers than that of neat EP or GNS–EP. In fact, it has been previously reported in the literature that solid acids can catalyze the degradation of polymers, which results in the formation of pyrolysis products with lower carbon numbers, and the degradation products with lower carbon numbers could easily catalyze carbonization in the presence of metal oxides.⁴²,⁶⁵ Manganese oxides have been reported to be the most efficient solid acid for suppressing the recombination of macromolecular hydrocarbon radicals due to the presence of several acidic sites.⁴²,⁶⁶ In this work, carbon numbers of pyrolysis products from the Ce–MnO₂–GNS–EP composite also decreased compared with that of EP. Thus, this undoubtedly results from the catalytic degradation effect for EP by Ce–MnO₂.⁴²,⁶⁶ Moreover, degradation products with lower carbon numbers could extend the contacting time of with metal oxides catalyst under the physical barrier effect of GNS. With MnO₂ catalyst, Hong et al. used a simple method to synthesize multi-layer graphene flakes from pyrolyzing biodegradable poly(butylene succinate) composites.⁴⁴ Thus, it is reasonably believed that the improved fire resistant properties of the Ce–MnO₂–GNS–EP nanocomposite are attributed to the synergism of the catalytic effect of Ce–MnO₂ and the physical barrier effect of GNS. In view of the results of volatile pyrolysis fragment ions, the mechanism for the improved fire resistant properties of the Ce–MnO₂–GNS–EP nanocomposite is illustrated in Scheme 2. During the combustion process, Ce–MnO₂ with catalytic activity leads to the formation of pyrolysis products with lower carbon numbers, which can be easily catalyzed for carbonization in the presence of metal oxides. Meanwhile, Ce–MnO₂ is reduced in situ to Ce–MnO by reductive hydrocarbons through redox reactions, which can catalyze the carbonization of degradation products.⁴⁴ Moreover, GNS acts as a physical barrier, which can absorb degraded products to extend contact time with metal oxides catalyst, and continually propagate on the GNS, which serves as a template of micro-char. Furthermore, the degraded products are dehydrogenated and catalytically converted into char by the combination of the GNS physical barrier effect and Ce–MnO₂ catalysis effect.

![Scheme 2 Illustration for the flame-retardant mechanism for the combination effect of the Ce–MnO₂–GNS hybrid.](image_url)

![Fig. 14 Total ion current (TIC) curves of the decomposition process of EP and its composites, and EI-MS spectra of compounds evolved from EP and its composites at the peak maximum in the TIC curves.](image_url)

![Fig. 15 Simplified mass fragmentations of the Ce–MnO₂–GNS–EP nanocomposite.](image_url)
4. Conclusions

In conclusion, Ce–MnO₂–GNS hybrid was successfully prepared through electrostatic interactions between Ce–MnO₂ and GNS, and the composition and structure were confirmed by XRD, EDS and XPS. TGA investigation revealed that the modification of GNS with Ce–MnO₂ could enhance the residue yield of GNS compared with pure GNS. The incorporation of 2 wt% Ce–MnO₂–GNS hybrid into EP led to the improvement of degradation temperatures, char residue and DTG peak value compared to that of pure EP. Meanwhile, the pHRR and THR values of Ce–MnO₂–GNS–EP were significantly reduced compared to those of neat EP. Moreover, the amount of organic volatiles released during the combustion of EP was significantly reduced and toxic CO was suppressed after incorporating Ce–MnO₂–GNS. The notable reduction of fire hazards was mainly attributed to the synergistic action between physical barrier effect of GNS and the catalytic effect of Ce–MnO₂. Ce–MnO₂ with catalytic activity resulted in the formation of pyrolysis products with lower carbon numbers, which can easily catalyze carbonization in the presence of metal oxides. GNS acted as physical barriers, which can absorb degraded products to extend the contact time with metal oxides catalyst, and continually propagate on the GNS, which serves as a template of micro-char, while pyrolysis fragments with lower carbon numbers could easily catalyze carbonization in the presence of Ce–MnO₂.

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Notes and references