

Preparation of a novel fame retardant based on phosphorus/nitrogen modifed lignin with metal–organic framework and its application in epoxy resin

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Abstract

To reduce the fre risk of epoxy (EP) resin, a renewable lignin-based fame retardant functionalized with a metal–organic framework (MOF) HKUST-1 (Cu₃(BTC)₂, BTC = benzene-1,3,5-tricarboxylate) was prepared by a facile and green method. The chemical structure of fame retardant was characterized by Fourier transform infrared (FTIR) spectroscopy, X-ray photoelectron spectroscopy (XPS), X-ray difraction (XRD) and scanning electron microscopy (SEM). The results showed that HKUST-1 was *in-situ* grown on the surface of phosphorus/nitrogen modified lignin, which was treated as an efficient flame retardant (termed PN-lignin@HKUST-1) for subsequent EP resin. The fame-retardant properties of EP composites were evaluated by the limiting oxygen index (LOI), vertical burning (UL-94), and cone calorimetry tests. As compared with neat EP thermoset, the presence of 15 mass% PN-lignin@HKUST-1 increased the LOI value from 28.5 to 33.2% and upgraded the UL-94 rating from V-2 to V-0. Besides, the peak heat release rate (PHRR) and total heat release (THR) were reduced by 41.2 and 33.8%, respectively. Observations of char residues could indicate that addition of PN-lignin@HKUST-1 contributes to a compact and dense char layer, which blocks the release and transfer of heat and harmful gases during combustion.

Graphic abstract

Keywords HKUST-1 · Lignin · Flame retardant · Epoxy resin

Extended author information available on the last page of the article

Introduction

As a conventional thermosetting plastic, epoxy (EP) resin is widely used in daily life, such as automotive adhesives, building decoration, electronic device insulating material $[1-4]$ $[1-4]$. Epoxy resins can be divided into five categories according to molecular structure, i.e., bisphenol-A epoxy resins, bisphenol-F epoxy resins, aliphatic glycidyl ether epoxy resin, glycidyl ester type epoxy resin and polyphenol type glycidyl ether epoxy resin. Among them, the most considerable output is bisphenol-A epoxy resin, which accounts for 90% of the epoxy resin applied in the world [[5\]](#page-10-2). Due to its excellent adhesion, high strength, electrical insulation and corrosion resistance, epoxy resin plays an increasingly signifcant role in many industries including laminates, adhesives, electronic/electrical industries [\[6,](#page-10-3) [7](#page-10-4)]. However, the limiting oxygen index (LOI) of epoxy resin is only about 26%, which means it is very fammable in air and seriously limits its application in many cases [[8\]](#page-10-5). Therefore, it is of great necessity and importance to develop a more efficient system to further improve the fame-retardant properties of epoxy resin.

The most common and efective method was adding fame retardants in the preparation of epoxy resin. Traditional fame retardants were usually halogen-fame retardants. Despite its outstanding fre resistance, epoxy resins containing halogen element fame retardants would produce toxic fumes during combustion, which were harmful to the environment and human health [[9](#page-11-0)]. Hence, there was a desire need to explore eco-friendly fame retardants to replace halogen-fame retardants. Compared with traditional fame retardants, intumescent fame retardants had the characteristics of high carbon content, good fameretardant efect and environmental protection. Intumescent fame retardants were constituted of the acid, carbon and gas sources, which expanded rapidly to form protective char layers. The carbon source was mainly polyhydroxy compounds, such as pentaerythritol and ethylene glycol. With the continuous research on biomass resources in recent years, it had been discovered that lignin can also be used as a carbon source [[10](#page-11-1)].

Lignin is one of the essential components of plant cell wall and is also the second most abundant renewable resource in nature next to cellulose $[11-13]$ $[11-13]$ $[11-13]$. It is a biopolymer containing many aromatic ring structures and diferent functional groups, namely carbonyl, carboxyl, methoxyl, aliphatic and aromatic hydroxyl moieties [[14](#page-11-4)]. On account of the high carbon yield after decomposition, lignin has received extensive attention in the preparation of bio-based fame retardants. It was reported that about 35–40% of carbon can be produced after pyrolysis of lignin [[15](#page-11-5)]. At that point, lignin can be incorporated into

polymers to improve the thermodynamic properties of the material. Besides, chemical modifcation of lignin and the introduction of fame-retardant elements such as phosphorus and nitrogen could enhance the fame-retardant perfor-mance of lignin-based composites [\[16–](#page-11-6)[20](#page-11-7)].

In the past several years, 9,10-dihydro-9-oxa-10-phosphaphenanthrene-10-oxide (DOPO) had been reported as an inorganic phosphorus fame-retardant intermediate since it could serve as an acid and gas source [[21,](#page-11-8) [22\]](#page-11-9). Its molecular structure contained a P–H bond, biphenyl ring and phenanthrene ring, with a higher thermal and chemical stability than the general, noncyclic organophosphate [[23–](#page-11-10)[27\]](#page-11-11). Due to these unique advantages, lignin-based fame retardants modifed with DOPO could signifcantly improve fameretardant efect, which showed great potential and broad application prospects [[28\]](#page-11-12).

Currently, metal–organic frameworks (MOFs) have attracted widespread interest for their superior capabilities and unique properties on the felds of fre-proof and fame retarding [[29](#page-11-13)[–31](#page-11-14)]. In a word, MOFs are crystalline porous materials with a periodic network structure, consisting of inorganic metal centers (metal ions or metal clusters) and bridging organic ligands connected by self-assembly [\[32](#page-11-15)]. The organic groups on the branches can also contain phosphorus, nitrogen and other fame-retardant elements [[33](#page-11-16)]. In addition, by reason of large pore volume and specifc surface area, MOFs can absorb toxic gas and improve the smoke-suppression performance of polymer materials [\[34](#page-11-17)]. Furthermore, MOFs have many unsaturated metal sites that can provide active catalytic centers for catalytic oxidation of CO, which can reduce the amount of CO released during combustion [[35\]](#page-11-18). Hence, the introduction of MOFs structure to epoxy resin is a promising attempt to achieve the solution.

In this work, research was aimed at promoting the high value-added utilization of industrial lignin and improving the fame resistance of epoxy (EP) resin. The phenolated alkali lignin was functionalized by coordinating the grafting of nitrogen and phosphorus (melamine and DOPO mixture) (referred to as PN-lignin) and further cellulose foam HKUST-1 $(Cu_3(BTC)_2, BTC=benzene-1,3,5-tricarboxy$ late) structure, which is one of the MOFs, was introduced by a green and convenient method (as shown in Scheme [1](#page-2-0)). The chemical composition and structure of prepared fame retardant (referred to as PN-lignin@HKUST-1) were characterized by Fourier transform infrared (FTIR), X-ray difraction (XRD) and X-ray photoelectron spectra (XPS). Epoxy resin was subsequently cured to fabricate EP composites by blending with as-prepared fame retardant (referred to as EP/PN-lignin@HKUST-1). The surface morphologies of PN-lignin@HKUST-1 and EP/PN-lignin@HKUST-1 were observed by scanning electron microscopy (SEM), and thermal behaviors were determined by thermogravimetric analysis (TGA). The fame-retardant properties of EP/

Scheme 1 Synthesis of PN-lignin@HKUST-1 as EP fame-retardant additive

PN-lignin@HKUST-1 composites were also measured by limiting oxygen index (LOI), vertical burning (UL-94), and cone calorimetry tests. This article would provide valuable information for the fame-retardant modifcation of lignin and reliable reference for the application of HKUST-1 based on lignin in epoxy resin.

Experimental

Materials

Alkaline lignin (Al-lignin) was purchased from TCI Shanghai Co. Ltd., China. DOPO, carboxymethyl cellulose sodium (CMC) and 4,4-diaminodiphenylmethane (DDM) were provided by Shanghai Macklin Co. Ltd., China. Diglycidyl ether of bisphenol-A type epoxy resin (commercial name: E 51) was purchased from Xingchen Epoxy Resins Factory (Nantong, China). Melamine (MEL) was obtained from Shanghai Lingfeng Co. Ltd., China. Formaldehyde, N,N-dimethylformamide (DMF), anhydrous copper acetate $(Cu(OAc₂), tri$ mesic acid (H_3BTC), glacial acetic acid (CH_3COOH), diethyl ether, ethanol and phenol were purchased from Nanjing Chemical Reagent Co. Ltd., China.

Phenolation of alkaline lignin (Ph‑lignin)

First, 20.00 g of Al-lignin was added to a flask containing 80 mL of H₂SO₄ solution (2.00 mol L⁻¹), heated up to 80 °C and stirred for 1.5 h. Then, when the temperature was raised to 95 °C, 18.00 g of phenol (0.20 mol L⁻¹) was added, and the mixture was stirred at refux for 1.5 h. After the reaction was completed, the mixture was washed three times by 500 mL of diethyl ether and dried in a vacuum oven at 70 °C overnight to a constant mass.

Preparation of MEL and DOPO mixture (MEL‑DOPO)

12.96 g of DOPO (0.06 mol) was added to 280 mL of ethanol aqueous solution (70 mass%) and stirred for 30 min, allowing the DOPO to be fully dispersed in the solution. Then, 10.00 g of MEL (0.08 mol) was added and stirred for 6 h when the temperature was raised to 70 °C. After completion, the mixture was cooled to room temperature, washed repeatedly with ethanol three times and dried in the vacuum oven at 70 °C for 12 h.

Preparation of PN‑lignin

Approximately 4.00 g of phenolated lignin, 20.00 g of MEL-DOPO (mass ratio, Ph-lignin/MEL-DOPO=1:5) and 7.20 g (0.24 mol) of formaldehyde were added to a fask with 300 mL of DMF and stirred under refux condition at 75 °C for 3 h. After the reaction was fnished, 500 mL of distilled water was added and the solid was fltered, washed three times with 500 mL of distilled water and dried at 70 °C under vacuum for 24 h.

Preparation of PN‑lignin@HKUST‑1

First of all, accurately 1.47 g of CMC (6.06 mol) with 2.00 g of PN-lignin was added into 70 mL of deionized water and mechanically stirred under room temperature for 1 h. Secondly, 1.33 g of $Cu(OAc)$, (6.64 mmol) dissolved in 30 mL of deionized water and 1 mL of HAc were added dropwise and kept in stirring for 5 min. Thirdly, 1.15 g of H_3BTC (5.50 mmol) dissolved in 15 mL ethanol was sequentially added, and the mixture was continuously stirred in a closed system for 4 h. Finally, the end product was obtained by centrifugation and dried in the vacuum oven at 70 °C for 24 h.

Preparation of fame‑retardant EP composites

A traditional curing method was applied to fabricate epoxy composites with some modifcations. Typically, 3.00 g of PN-lignin@HKUST-1 and 15.00 g of EP were mechanically stirred at 1000 rpm for 1.5 h to obtain a uniform mixture system. Next, 3.00 g of curing agent DDM (mass ratio, DDM/ $EP = 1:5$) was added and kept in stirring for 1.5 h. The uniformly mixed thermoset was poured into the mold

Table 1 Information about lignin-based materials

Sample	Components
Al-lignin	Alkaline lignin
ph-lignin	Phenolation of Al-lignin
PN-lignin	ph-lignin mixed with MEL-DOPO
EP/PN-lignin	PN-lignin mixed with epoxy resin
PN-lignin@HKUST-1	PN-lignin loaded by HKUST-1
EP/PN-lignin@HKUST-1	EP/PN-lignin loaded by HKUST-1

and cured at 100 °C for 2 h after bubbles were removed in the vacuum oven. For further curing, the temperature was raised to 150 °C for 2 h. To evaluate the effect of modification, the neat EP thermoset and EP composites containing PN-lignin (referred to as EP/PN-lignin) were also prepared according to the similar procedure.

Above, lignin-based materials information was listed in Table [1](#page-3-0).

Measurements

The Fourier transform infrared (FTIR) spectra were recorded with a Nicolet 6700 FTIR spectrometer by using compressed potassium bromide (KBr) disks. The scanning range of infrared spectrometer was 4000–500 cm⁻¹ with 32 scans per spectrum. The thermogravimetric analysis (TGA) was obtained on DTG-60AH (SHIMADZU, Japan). The epoxy resin composites (about 5 mg) were heated (30–800 °C) at a rate of 20 $^{\circ}$ C min⁻¹ under the nitrogen flow of 20 mL min⁻¹. The X-ray difraction (XRD) patterns were tested with the Japanese Ultima IV combined multifunctional horizontal X-ray difraction spectrometer. The scanning speed was 10° min−1, and the scanning difraction angle 2*θ* was 5–80°. The X-ray photoelectron spectroscopy (XPS) spectra were recorded by an AXIS UltraDLD spectrometer with Al Kα (1486.6 eV) radiation. The surface morphologies of samples were observed using a JEOLJSM-7600 scanning electron microscopy (SEM) (JEOL, Japan) at the accelerating voltage of 15 kV. The limiting oxygen index (LOI) values of composites were measured using an HC-2C oxygen index meter (Jiangsu Institute of Chemical Industry, China). The size of three samples was 100 mm \times 10 mm \times 3 mm according to ASTM D2863-97 standard. The Vertical burning (UL-94) test was conducted on a CZF-2 instrument (Jiangsu Institute of Chemical Industry, China), and the size of three epoxy resin composites was $100 \text{ mm} \times 10 \text{ mm} \times 3 \text{ mm according}$ to ATSM D3801 standard. The combustion behavior of the samples was measured by a cone calorimeter (Fire Testing Technology, UK). For each formulation, the samples (100 mm \times 100 mm \times 3.2 mm dimension) were exposed to a radiant cone under a heat fux of 50 kW m−2 according to ISO 5660 standard. The samples were backed with aluminum foil in a metal frame. Laser Raman spectroscopy (LRS) was also used to test the residue by confguring a laser system with a wavelength of 532 nm.

Results and discussion

Characterization of prepared fame retardants

As shown in Fig. [1a](#page-3-1), DOPO is predominantly columnar in shape with a varying length and relatively smooth surface. It can be seen from Fig. [1b](#page-3-1) that foam HKUST-1 displays the shape of cubic crystal, which is consistent with previous report [[36](#page-11-19)]. The crystal surface is also covered with a thin web of fbers since HKUST-1 crystals are grown from the copper irons (Cu^{2+}) bridging of CMC chains and ligands. From Fig. [1](#page-3-1)c, d, it shows that the surface of DOPO columns becomes coarse and irregular. It is because DOPO is coated with lignin by Mannich reaction between lignin and MEL to form PN-lignin. There are many small cubes unevenly distributed on the surface of PN-lignin@HKUST-1, and the fber webs are not particularly obvious, which is due to the adhesion of CMC to the surface of lignin and then, the adsorption of metal ions to generate HKUST-1. Furthermore, the size of foam HKUST-1 slightly decreases because the growth of HKUST-1 on the lignin surface is restricted. It may be because Cu^{2+} is adsorbed to the surface under the effect of CMC [\[37](#page-11-20)]. These results preliminarily indicate that DOPO is coated with lignin and HKUST-1, and consequently fame retardant PN-lignin@HKUST-1 is prepared.

To demonstrate the chemical structure of the as-prepared samples, the FTIR spectra are performed and presented in Fig. [2.](#page-4-0) The absorption peaks at 1625/1570 and

Fig. 1 SEM images of DOPO (**a**), foam HKUST-1 (**b**) at 10,000 times magnifcation, and PN-lignin@HKUST-1 (**c**, **d**) at 5000 times magnifcation

Fig. 2 FTIR spectra of Al-lignin (**a**), HKUST-1 (**b**), PN-lignin (**c**) and PN-lignin@HKUST-1 (**d**)

Fig. 3 XRD spectra of foam HKUST-1 (**a**), PN-lignin@HKUST-1 (**b**) and PN-lignin (**c**)

1440/1370 cm⁻¹ are attributed to –COOH and –CH₂–/–CH₃ in foam HKUST-1, respectively [\[38\]](#page-12-0). The absorption peaks of 754, 1205 and 1569 cm⁻¹ are assigned to the stretching vibrations of P–O–Ph, P=O and P–Ph, which belong to DOPO structure [\[39](#page-12-1), [40\]](#page-12-2). Moreover, the presence of P–O–C vibration can be observed at 1050 cm^{-1} . The appearance of these characteristic absorption peaks in the FTIR spectrum of PN-lignin@HKUST-1 confrms that DOPO and HKUST-1 are successfully introduced into modifed lignin [\[41\]](#page-12-3).

The crystal structures of foam HKUST-1 and synthesized fame retardant are determined by XRD. As shown in Fig. [3](#page-4-1)a, the XRD spectrum suggests the intact difraction characteristic peak profles of HKUST-1 [\[42\]](#page-12-4). In peculiar, the appearance of lattice plane in (200) facet is due to the formation of cubic crystal structure, as evidenced by SEM observations [[36\]](#page-11-19). From Fig. [3](#page-4-1)b, it reveals that the characteristic peaks of PN-lignin@HKUST-1 are the same as those of HKUST-1 except that their peak intensity is weakened. It is because only part of the crystal structure of HKUST-1 is exposed on the surface, and the rest is covered by functionalized lignin or destroyed after adsorption [[43](#page-12-5)]. Besides, the peak of PN-lignin (Fig. [3](#page-4-1)c) disappears when HKUST-1 has been loaded, which is mainly because of the high intensity of HKUST-1 characteristic peaks.

XPS is an efficient equipment to analyze the elemental composition and type of chemical bonds in PN-lignin@ HKUST-1. As shown in Fig. [4a](#page-5-0), b, the P element exists in the XPS spectrum of PN-lignin@HKUST-1. Two main peaks obviously appear in the Cu2p spectrum (Fig. [4c](#page-5-0)) at 932.89 and 952.59 eV, which are attributed to $Cu2p_{3/2}$ and $Cu2p_{1/2}$, respectively. Furthermore, the spectrum shows two weak peaks at 934.31 and 954.31 eV, corresponding to the satellite peak of Cu2p_{3/2} and Cu2p_{1/2}, respectively. In addition, the O1s spectrum of PN-lignin@HKUST-1 (Fig. [4d](#page-5-0)) is divided into three peaks appearing at 532.06, 532.98 and 533.76 eV, which are ascribed to Cu–O, C=O and C–OH, respectively [[44\]](#page-12-6). The above evidences fully prove the successful preparation of fame retardant PN-lignin@HKUST-1.

Thermal behavior of prepared fame retardants

To investigate the mass-loss behavior and thermal decomposition process of lignin and fame retardants under nitrogen atmosphere, the TGA and DTG curves are presented in Fig. [5](#page-5-1) with the detailed data listed in Table [2](#page-5-2). The degradation route is traced by detecting the initial degradation temperature (T_i) , where a mass loss of 5 mass% takes place), the maximum temperature (T_{max}) , where the massloss rate reaches a maximum) and the mass of residue left. As can be seen from Table [2](#page-5-2), the thermal decomposition of Al-lignin starts from 105.54 °C and shows a sharp decrease at 333.36 °C, with 41.57 mass% residue left at 800 °C. The T_i of PN-lignin and PN-lignin@HKUST-1 are both lower than that of Al-lignin. The onset degradation temperature decreases due to the incorporation of DOPO, which has a high gas-phase activity and condensed-phase activity (through char formation) $[25, 45]$ $[25, 45]$ $[25, 45]$. In addition, the O=P–O bond in DOPO is less stable than the common C–C bond, which is also responsible for the advanced decomposition. Compared with original Al-lignin, the char residues of PN-lignin and PN-lignin@HKUST-1 are reduced because of the relatively low lignin content (Fig. [5](#page-5-1)a), which are 18.24 mass% and 31.13 mass%, respectively. However, the char residue of PN-lignin@HKUST-1 is much higher than that of PN-lignin. The underlying reason is the catalytic ability of metal copper ions in HKUST-1 on the char formation process, which can contribute to protecting the

Fig. 5 TGA (**a**) and DTG (**b**) curves of Al-lignin, PN-lignin

polymer when their composites are exposed to the fre [\[46\]](#page-12-8). As shown in Fig. [5b](#page-5-1), the decomposition of PN-lignin exhibits one rapid and violent degradation stage with T_{max} at about 354.73 °C, which is primarily attributed to the degradation of DOPO and MEL, along with the formation of polyphosphate and release of ammonia $(NH₃)$, nitrogen (N_2) and water vapor (H_2O) [[47,](#page-12-9) [48\]](#page-12-10). The followed stage after 450 °C is potentially assigned to the degradation of polyphosphate for the generation of phosphoric acids and metaphosphoric acids [[47](#page-12-9)]. The degradation process of PN-lignin@HKUST-1 is more complicated, and the T_{max} (309.22 °C) is much lower than that of PN-lignin, on account of the introduction of foam HKUST-1, which

Table 2 TGA data of Al-lignin, PN-lignin and PN-lignin@HKUST-1 under N_2 atmosphere

Sample	T_i /°C	T_{max} /°C	Residue/ mass%
Al-lignin	105.54	333.36	41.57
PN-lignin	87.10	354.73	18.24
PN -lignin $@$ HKUST-1	83.57	309.22	31.13

has a relatively weaker thermal stability and decomposes gradually at about 320 °C [\[36\]](#page-11-19).

Fig. 6 SEM images of EP thermoset (**a**, **b**) and EP/PN-lignin@ HKUST-1 composite (**c**, **d**) at 10,000 times magnifcation

Dispersibility of fame retardant in prepared EP composites

To verify the compatibility and interfacial interaction between fame retardant and polymeric matrix, the SEM images of fractured sections for EP thermoset and EP/PNlignin@HKUST-1 are shown in Fig. [6.](#page-6-0) It can be clearly seen that there are few folds distributed on the fat and smooth fractured surface of neat EP thermoset. Almost no embedded particles are visible under this circumstance. By contrast, a series of irregular folds and particles of diferent sizes on the fractured surface of EP/PN-lignin@HKUST-1 composite can be observed. This obvious diference is attributed to the inclusion of PN-lignin@HKUST-1, which implies the strong interfacial interaction between additives with EP thermoset matrix. The result shows that PN-lignin@HKUST-1 has been absolutely mixed with EP thermoset and possessed a great dispersibility in the composite.

Thermal behavior of prepared EP composites

The TGA, DTG curves of EP thermoset and its composites under N_2 atmosphere are shown in Fig. [7,](#page-6-1) and the corresponding data are listed in Table [3](#page-7-0). The thermal decomposition process of neat EP thermoset undergoes only one main degradation step, which starts from 333.74 °C and shows a sharp decrease at 405.35 °C, leaving a char residue of 1[7](#page-6-1).21 mass% at 800 °C (Fig. 7a). The T_i of EP/PN-lignin and EP/PN-lignin@HKUST-1 composites are shifted to a lower range, which can be attributed to the early degradation of the fame retardants and the catalytic efect of the metals [[49\]](#page-12-11). The T_{max} of all of the additive modified composites is also brought forward to diferent extents compared with neat EP thermoset. It can be seen from Fig. [7b](#page-6-1) that T_{max} shows a decreasing trend as $EP > EP/10\%$ PN-lignin > $EP/15\%$ PN-lignin>EP/10% PN-lignin@HKUST-1>EP/15% PNlignin@HKUST-1. For intumescent fame-retardant system, the faster decomposition of fame retardants is conducive to accelerating the charring of polymeric matrix. This tendency suggests that the fame retardants can come into play at a lower temperature range, especially for EP/PN-lignin@ HKUST-1, the T_{max} of which is about 20 $^{\circ}$ C lower than that of EP/PN-lignin at the same content. The early decomposition of the composite can be explained by the presence of copper irons (Cu^{2+}) , which can promote the decomposition of molecular chains of polymer due to the catalytic interaction of open metal sites and unsaturated covalent bonds [[49](#page-12-11)]. As shown in Table [3,](#page-7-0) the char residue of EP/PN-lignin and EP/PN-lignin@HKUST-1 is much higher than that of the EP thermoset, which increases greatly with increasing the additive amount. Therefore, the rapid degradation and catalysis of PN-lignin@HKUST-1 are beneficial for the quick formation and signifcant enhancement of char residue [\[50](#page-12-12)].

Flame‑retardant properties of prepared EP composites

Table [4](#page-7-1) shows the fame retardancy of EP thermoset and its composites evaluated by the limited oxygen index (LOI) and UL-94 vertical test. The LOI value of pure EP is only 26.1%, indicating that EP thermoset tends to burn easily in air. The LOI values of EP composites are improved to different degrees by the addition of fame retardants. When the amount of PN-lignin added is 10 mass%, the LOI value of EP composite is 28.5%, while has no fame-retardant rating. The LOI value increases from 28.5 to 29.8% as the amount of PN-lignin content increases from 10 to 15 mass%, and the latter just achieves V-2 rating in UL-94 vertical test. When 10 and 15 mass% of PN-lignin@HKUST-1 are applied in the EP, the LOI value remarkably increases to 31.7% and 33.2%, respectively, which suggests that it is difficult for the composites to burn under such conditions. Meanwhile, EP/10% PN-lignin@HKUST-1 can reach V-1 rating and EP/15% PN-lignin@HKUST-1 can even reach V-0 rating in UL-94 vertical test, revealing an important role of PNlignin@HKUST-1 in restraining the infammation. These results show that the lignin-based fame retardant modifed by nitrogen and phosphorus can improve the fame-retardant properties of EP thermoset, but they may not meet the application requirements in practice. However, the LOI value and UL-94 fame retardant level of the composites can be dramatically improved when the HKUST-1 structure is introduced, which is mainly attributed to the synergistic fame-retardant efect of DOPO and HKUST-1.

Table 3 TGA data of EP thermoset, EP/PN-lignin and EP/PNlignin@HKUST-1 composites under N_2 atmosphere

Sample	T_i /°C	$T_{\rm max}$ /°C	Residues/ mass%
EP	333.74	405.35	17.21
EP/10% PN-lignin	324.11	387.83	19.74
EP/15% PN-lignin	309.26	384.97	23.72
EP/10% PN-lignin@HKUST-1	332.62	364.96	18.05
EP/15% PN-lignin@HKUST-1	330.14	360.73	20.14

Cone calorimetry test is a comprehensive evaluation method for the fre risk of polymers by monitoring many important real-time and statistical parameters throughout combustion process. The heat release rate (HRR) and total heat release (THR) curves for EP thermoset, EP/15% PNlignin and EP/15% PN-lignin@HKUST-1 composites are given in Fig. [8,](#page-8-0) and the key combustion parameters collected from the test are listed in Table [5.](#page-8-1) The ignition time (time to ignition, TTI) of EP/15% PN-lignin and EP/15% PNlignin@HKUST-1 composites is shorter than that of neat EP thermoset, indicating that addition of PN-lignin and PN-lignin@HKUST-1 can reduce the initial pyrolysis temperature, which is in exact accordance with the TGA results. For catalytic char formation system, the short ignition time means that flame retardants can take effect in the early combustion process, decomposing inorganic acids and metal ions to promote the formation of residue char [[51,](#page-12-13) [52\]](#page-12-14). The high peak heat release rate (PHRR) and THR values of the EP thermoset are as high as 1089.6 kW m⁻² and 155.2 MJ m⁻², respectively. The PHRR value is decreased to 906.5 kW m⁻², and the THR value is also decreased to 129.4 MJ m⁻² as 15 mass% PN-lignin is added in the composite, which exhibits relatively good fame retardancy. While the incorporation of 15 mass% PN-lignin@HKUST-1 is capable of further reducing the values of PHRR and THR to 640.1 kW m^{-2} and 102.7 MJ m−2, respectively, corresponding to 41.2 and 33.8% reductions compared to neat EP thermoset. On the one hand, PN-lignin@HKUST-1 can generate non-fammable gases such as ammonia, carbon monoxide and water vapor during combustion, which reduce the concentration of fammable gases in the polymer and absorb large amounts of heat to reduce the surface temperature [[48\]](#page-12-10). On the other hand, the char residue of EP/15% PN-lignin@HKUST-1 (11.80 mass%) is far more than that of EP thermoset (0.50 mass%) and EP/15% PN-lignin (5.10 mass%). As mentioned, it mainly owes to the catalytical char formation capacity of copper ions in HKUST-1, which can facilitate the generation of protective char layer, thereby blocking the oxygen and thermal, and inhibiting the volatilization of fammable gas. Based on these results, it can be inferred that EP/PNlignin@HKUST-1 presents the extremely superior fame retardancy.

Table 4 LOI values and UL-94 vertical test results of EP thermoset, EP/PN-lignin and EP/PN-lignin@HKUST-1 composites

Sample	EP/mass%	DDM/mass%	PN -lignin/mass%	PN -lignin $@$ HKUST-1/mass%	$LOI\%$	UL-94 rating
EP	83.3	16.7			26.1	$V-2$
$EP/10\%$ PN-lignin	75.0	15.0	10.0		28.5	$V-2$
EP/15% PN-lignin	70.8	14.2	15.0		29.8	$V-2$
EP/10% PN-lignin@HKUST-1	75.0	15.0		10.0	31.7	$V-1$
EP/15% PN-lignin@HKUST-1	70.8	14.2		15.0	33.2	$V-0$

Fig. 8 HRR (**a**) and THR (**b**) curves of EP thermoset, EP/15% PN-lignin and EP/15% PNlignin@HKUST-1 composites

Table 5 Detailed data obtained from cone calorimeter tests of EP thermoset, EP/15% PN-lignin and EP/15% PN-lignin@HKUST-1 composites

Apart from the heat release, the release of smoke and toxic gases is another factor to consider for fame-retardant additives, especially carbon monoxide, which constitutes a fatal disadvantage of traditional phosphorous-containing fame retardants in practical applications [[49](#page-12-11)]. Compared with the smoke release of the neat EP thermoset, the incorporation of PN-lignin and PN-lignin@HKUST-1 mildly increases the total smoke production (TSP) of composite materials. However, the average carbon monoxide yield

Fig. 9 Digital photographs of char residues of EP thermoset (**a**), EP/15% PN-lignin (**b**) and EP/15% PN-lignin@HKUST-1 (**c**) composites after cone calorimeter test

Fig. 10 SEM images of chars for EP thermoset (**a**, **b**), EP/15% PNlignin (**c**, **d**) and EP/15% PN-lignin@HKUST-1 (**e**, **f**) composites at 10,000 times magnifcation

Fig. 11 LRS spectra of the residues of EP thermoset (**a**), EP/15% PN-lignin (**b**), EP/15% PN-lignin@HKUST-1 (**c**) composites

(av-COY) of EP/15% PN-lignin@HKUST-1 composite is 0.237 kg kg⁻¹, about half that– of the EP thermoset $(0.439 \text{ kg kg}^{-1})$. It is mainly attributed to HKUST-1, which can provide active catalytic centers for the oxidation of CO [[53\]](#page-12-15). Moreover, HKUST-1 also can absorb gases released because of its high porosity and large specifc surface area. Therefore, the average carbon dioxide yield (av- $CO₂Y$) of EP/15% PN-lignin@HKUST-1 composite (2.016 kg kg−1) is much lower than that of the EP thermoset (2.821 kg kg^{-1}) [[54](#page-12-16), [55\]](#page-12-17). It is reasonable to conclude that PNlignin@HKUST-1 possesses outstanding fame-retardant ability, for the reason that it can reduce the exothermic rate, increase the amount of char residue, and inhibit the production of harmful gases.

Char residue analysis of prepared EP composites

Figure [9](#page-8-2) shows the digital photographs of EP thermoset, EP/15% PN-lignin and EP/15% PN-lignin@HKUST-1 composites after cone calorimetry test. The neat EP sample is almost burnt out with nearly no char residue left, while the other two composites have much more char yields in favor of serving as a isolated layer. It can be seen that a certain degree of swelling occurs after the combustion of EP/15% PN-lignin, but the surface of char layer appears to be loose and irregular, which cannot prevent the material

well from burning. In comparison, the addition of 15 mass% PN-lignin@HKUST-1 can form a dense char layer on the surface, accompanied with a regular appearance and signifcant expansion. It is due to the combined efects of DOPO and HKUST-1: the polyacids, metal ions and metal oxides generated by decomposition at high temperatures catalyze the formation and dehydration of char layer, resulting in a more compact and intact structure. Furthermore, the nonfammable gas produced by thermal decomposition greatly expands the char layer, thus isolating the oxygen and achieving the purpose of fame retardant.

As shown in Fig. [10](#page-8-3), SEM images are investigated to further measure the morphology of char residue of EP thermoset and its composites after the cone calorimetry test. The char layer of neat EP is cracked and fractured, whereas EP/15% PN-lignin tends to form an unbroken and compact char layer that can act as a shield against the transfer of gases and heat. However, many broken pores can still be clearly observed on the surface, which is probably due to the gases fushing out from the weak points of char layer. By contrast, the pores on the char layer of EP/15% PN-lignin@ HKUST-1 nearly disappeared after the introduction of HKUST-1, as evidenced by the high char residue and gas absorption capacity. Such a dense and complete protective layer cannot only isolate the oxygen and heat, but also efectively prevents the escape of toxic gases and smoke [\[56\]](#page-12-18). The conclusions obtained by the SEM images are in agreement with the digital graphs shown above.

Figure [11](#page-9-0) shows the Raman spectra of the char layer of EP thermoset, EP/15% PN-lignin and EP/15% PN-lignin@ HKUST-1 composites. There are two strong absorption peaks in the Raman spectra of the three samples, near 1356 and 1593 cm−1, respectively, which are the representative peaks of graphite (G and D peaks). Generally speaking, the ratio of the integrated intensities of D to G band (I_D/I_G) is usually used to assess the degree of char layer. The lower ratio of I_D/I_C means the higher graphitization degree of car-bon layer, indicating that the layer is denser [[57\]](#page-12-19). The $I_D/$ I_G value of the neat EP thermoset and EP/15% PN-lignin is 2.81 and 2.77, respectively, indicating that the graphitization degree of char layer is very low. However, the I_D/I_G value of the EP composite is further decreased to 2.61 after adding PN-lignin@HKUST-1. The metal irons in the HKUST-1 can act as catalysts to generate more graphitized carbon during combustion, which contributes to the formation of char structure with improved thermal stability.

Conclusions

In this work, fame retardant PN-lignin@HKUST-1 is developed through the dual modifcation of phenolated Al-lignin by MEL-DOPO and HKUST-1. The acquired PN-lignin@ HKUST-1 is further added into epoxy resin (EP) to prepared flame-retardant (EP/PN-lignin@HKUST-1) composites using physical blending method. The FTIR, XRD and XPS spectra demonstrate that DOPO and HKUST-1 are successfully introduced into modified lignin. The SEM morphological observation proves that PN-lignin@ HKUST-1 in EP composite possesses great compatibility and dispersibility. Compared with PN-lignin, PN-lignin@ HKUST-1 displays excellent flame-retardant properties and pretty good catalytical char formation capacity in the EP composites. It is concluded that EP/15% PN-lignin@ HKUST-1 can reach up to V-0 rating in UL-94 vertical test and the limited oxygen index (LOI) value is 33.2%. The peak heat release rate (PHRR) and total heat release (THR) of EP composite with 15 mass% PN-lignin@HKUST-1 are 640.1 kW m⁻² and 102.7 MJ m⁻², respectively. The average carbon monoxide yield (av-COY) of EP/15% PN-lignin@ HKUST-1 is 0.237 kg kg−1 and the average carbon dioxide yield (av-CO₂Y) is $(2.016 \text{ kg kg}^{-1})$. These results indicate that PN-lignin@HKUST-1 can reduce the exothermic rate, increase the amount of char residue and inhibit the production of harmful gases. In addition, EP/15% PN-lignin@ HKUST-1 forms a dense and compact char layer, which can block the internal material from exposure to heat and oxygen in combustion process.

Author contributions HY contributed to the conceptualization, data curation, formal analysis, investigation, methodology, validation, visualization, writing-original draft, writing-review and editing and funding acquisition. YQ was involved in the conceptualization and resources. DL was involved in the data curation and resources. XL and XG were involved in the project administration and funding acquisition. All authors have read and agreed to the published version of the manuscript.

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Data availability All data supporting the fndings of this research are authentic and available in this article.

Declarations

Conflict of interest The authors state that they have no known competing fnancial interests or personal relationships that could have appeared to infuence the work reported in this paper.

Ethical approval Not applicable.

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