Flame-Retardant Performance of Transparent and Tensile-Strength-Enhanced Epoxy Resins

Liang Li 1,2 and Zaisheng Cai 1,*

1 College of Chemistry, Chemical Engineering and Biotechnology, Donghua University, Shanghai 201620, China; liliang1618@126.com
2 School of Pharmacy, Yancheng Polytechnic College, Yancheng 224005, China
* Correspondence: zshcai_dhu@126.com; Tel.: +86-178-5160-0015

Received: 19 December 2019; Accepted: 29 January 2020; Published: 4 February 2020

Abstract: In this study, a flame-retardant additive with 9,10-dihydro-9-oxa-10-phosphaphenanthrene-10-oxide (DOPO) groups denoted DSD was successfully synthesized from DOPO, 4,4′-diaminodiphenyl sulfone (DDS), and salicylaldehyde. The chemical structure of DSD was characterized by FTIR–ATR, NMR, and elemental analysis. DSD was used as an amine curing agent, and the transparent, tensile strength-enhanced epoxy resins named EP–DSD were prepared via thermal curing reactions among the diglycidyl ether of bisphenol A (DGEBA), 4,4′-diaminodiphenylmethane (DDM), and DSD. The flame-retardancy of composites was studied by the limiting oxygen index (LOI) and UL-94 test. The LOI values of EP–DSD composites increased from 30.7% for a content of 3 wt % to 35.4% for a content of 9 wt %. When the content of DSD reached 6 wt %, a V-0 rating under the UL-94 vertical test was achieved. SEM photographs of char residues after the UL-94 test indicate that an intumescent and tight char layer with a porous structure inside was formed. The TGA results revealed that EP–DSD thermosets decomposed ahead of time. The graphitization degree of the residual chars was also investigated by laser Raman spectroscopy. The measurement of tensile strength at breaking point shows that the loading of DSD increases the tensile strength of epoxy thermosets. Py-GC/MS analysis shows the presence of phosphorus fragments released during EP–DSD thermal decomposition, which could act as free radical inhibitors in the gas phase. Owing to the promotion of the formation of intumescent and compact char residues in the condensed phase and nonflammable phosphorus fragments formed from the decomposition of DOPO groups, EP–DSD composites displayed obvious flame-retardancy.

Keywords: thermosets; amine curing agent; DOPO derivatives; mechanical properties; epoxy resins

1. Introduction

Epoxy resins are used in a broad variety of fields, such as electrical, electronics, and adhesive, as well as casting applications and construction because of their high adhesion, outstanding electrical and mechanical properties, and good alkali and moisture resistance [1–3]. However, the flammability and low transparency of epoxy resin composites are major disadvantages in their application, especially when used in circuit boards and coatings [4,5]. Several approaches have been utilized to enhance the thermal properties of epoxy resins, the most widely used method of which is the addition of flame-retardant additives, such as halogenated compounds, nitrogenous compounds, organophosphorus compounds, and inorganic materials [6–9].

Nanocomposites have taken a particular place in the preparation of transparent and flame-retardant coatings due to their thermal, light transmittance, and mechanical properties [10,11]. Morteza et al. prepared a new class of transparent epoxy-based nanocomposite coatings containing starch-modified nano-zinc oxide (ZnO–St), which can be applied as top-coats because of their transparency. They studied
the curing behaviors of epoxy and transparent nanocomposites containing ZnO and ZnO-St, and found that the ZnO-St prolonged curing and increased the amount of heat release because of its autocatalytic effect [12]. Hadi et al. developed two kinds of layered double hydroxides (LDHs) and prepared the transparent nanocomposite coatings with sodium dodecylbenzene sulfonate (SDBS)-modified LDHs, Mg–Al and Zn–Al LDHs. Cure kinetics and viscoelastic behavior were studied. Network formation in the presence of SDBS-modified Zn–Al LDH nanoplatelets was considered to play an important role in the enhancement of epoxy ring-opening [13]. However, the flame retardancy of these nanocomposites, such as the limiting oxygen index (LOI) and UL-94 test, have not frequently been mentioned.

Due to the demand for environmentally friendly flame retardants (FRs) [14,15], organophosphorus compounds have been considered as dependable FR alternatives because of their good flame retardancy without the generation of lots of toxic smoke and corrosive gases [16]. While heating organophosphorus flame-retardant systems, an intumescent char layer is generated, acting as a physical isolation, which blocks heat transfer from the flame to the materials [17,18]. When combined with a nitrogen element, an obvious improvement in the flame-retardant efficiency is produced due to a phosphorus–nitrogen synergistic effect [19,20].

DOPO (9,10-dihydro-9-oxa-10-phosphaphenanthrene-10-oxide) has been frequently used as starting material for the synthesis of organophosphorus compounds. As a phosphorus element supplier, it can react with various electron-deficient compounds, such as unsaturated nitrogenous compounds, by the addition reaction of P–H, to produce various DOPO-based derivatives, and can then form phosphorus–nitrogen or phosphorus–silicon synergism [8,21,22]. There are lots of works reporting on the molecular design and synthesis of flame-retardant additives based on DOPO, as well as relevant flame-retardant epoxy resin composites [23,24]. The DOPO derivatives have proved to be excellent FRs due to their versatile flame extinguishing behavior in both the gas phase and condensed phase. In particular, for an intumescent flame-retardant system containing phosphorus and nitrogen, it can not only form an intumescent residue char, which works as a barrier to inhibit heat reflux in the condensed phase, but can also release many phosphorus fragments, which restrain the radical reaction in the gas phase [25–27]. However, these phosphorus-containing epoxy resin composites have some disadvantages, such as a low transparency, decreased mechanical properties, and a higher added ratio of DOPO-based derivatives due to their low phosphorus content, leading to a low number of applications [28,29]. This is necessary, not only to enhance the flame-retardant performance, but also to maintain its mechanical properties and transparency for coating. Yang et al. employed a hydroxyl-functionalized hyperbranched polymer (H30) to improve the mechanical properties of a diglycidyl ether of bisphenol A (DGEBA) epoxy resin. They found that the tensile strength was improved by adding a proper content of H30, and the glass transition temperature ($T_g$) was not reduced by the addition of H30 in appropriate amounts [30]. Increasing the three-dimensional epoxy networks contributed to improvement of the tensile strength. As a nitrogen element supplier, 4,4′-diaminodiphenyl sulfone (DDS) is an easily available amine curing agent and compatible with DGEBA epoxy resin. When combined with phosphorus-containing compounds and reactive groups such as –OH, it can construct a phosphorus–nitrogen synergism with a multidimensional curing reaction.

In this work, one polyfunctional DOPO derivative defined as DSD was successfully synthesized and used as an amine curing agent to improve the flame-retardancy and tensile strength of DGEBA epoxy resins. Its chemical structure was well-characterized by FTIR–ATR, $^1$H NMR, and $^{31}$P NMR. The flame-retardant EP–DSD composites were prepared, and their flame-retardant properties were studied by the limiting oxygen index (LOI) and UL-94 test. Thermal decomposition, combustion behaviors, the structure of char residues, and the mechanical behaviors of these prepared thermosets were also investigated.
2. Materials and Methods

2.1. Materials

DOPO was supplied by Shanghai Nuotai Chemical Technology Reagent Co., Ltd. (Shanghai, China) and recrystallized in ethanol before use. DDS (4,4′-diaminodiphenyl sulfone) and salicylaldehyde were purchased from Aladdin Industrial Co., Ltd. (Shanghai, China). Tetrahydrofuran (THF) and 4,4′-diaminodiphenylmethane (DDM) were purchased from Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China). Diglycidyl ether of bisphenol A (DGEBA, commercial name: E-44) was provided by Kunshan Lvxun Chemical Co., Ltd. (Kunshan, China).

2.2. Synthesis of DSD

The synthetic routes of DSD are illustrated in Scheme 1.

![Scheme 1. Synthetic routes of the intermediate imine (DS) and a 9,10-dihydro-9-oxa-10-phosphaphenanthrene-10-oxide (DOPO)-based flame retardant (DSD).](image)

The intermediate imine coded as DS was synthesized by a typical condensation reaction; 4,4′-diaminodiphenyl sulfone (0.04 mol, 9.92 g) and THF 100 mL were fed into a 250 mL three-necked flask equipped with a magnetic stirrer and a reflux condenser, and the mixture was then heated to 40 °C. Subsequently, salicylaldehyde (0.08 mol, 9.77 g) dissolved in THF (60 mL) was added dropwise for 30 min under the protection of nitrogen. After this, the solution was heated to reflux temperature and maintained for 6 h. After cooling, the orange precipitate was filtered, washed with THF thoroughly, and dried at 60 °C for 18 h in a vacuum oven (17.62 g, yield: 96.5%).

The DSD was synthesized by an addition reaction. DS (0.02 mol, 9.13 g) and THF (80 mL) were added to a three-necked flask (250 mL) equipped with a thermometer, a reflux condenser, and a stirrer. Under the protection of nitrogen, DOPO (0.04 mol, 8.64 g) dissolved in THF (80 mL) was added for 1 h. With stirring, the reaction solution was heated and maintained at reflux temperature for 12 h. Then, the reaction solution was cooled, filtered by vacuum filtration, washed thoroughly with anhydrous alcohol and distilled water, and dried at 100 °C for 24 h until reaching a constant weight. A white solid was obtained (16.97 g, yield: 95.3%).

2.3. Preparation of Flame-Retardant Epoxy Resin

Flame-retardant epoxy resin was prepared via thermal curing reactions among DGEBA, DDM, and DSD. The formulations of modified epoxy resin are listed in Table 1, where the sum amounts of phenolic and amine protons are equal to the number of epoxy groups in DGEBA. DSD and DDM were blended at 90 °C for 30 min under a mechanical stirrer. When the mixture became transparent and uniform, the DGEBA was introduced and mixed for 5 min under a vacuum pump. Then, the mixture was poured into PTFE molds and cured at 120 °C for 2 h, followed by heat maintenance at 160 °C.
for another 2 h. Finally, the modified epoxy resin samples of a fixed size were cooled to room temperature slowly. The neat epoxy resin samples were also prepared under similar processing conditions, but without DSD.

Table 1. The formulas of EP–DSD composites.

| Samples  | Composition (g) | DSD (wt %) | P (wt %) | N (wt %) |
|----------|----------------|------------|----------|----------|
|          | DGEBA | DDM | DSD       |          |          |
| EP0      | 100   | 22  | 0         | 0        | 2.54     |
| EP–DSD3  | 100   | 21.16 | 3.75  | 3        | 0.21     | 2.49     |
| EP–DSD6  | 100   | 19.38 | 7.62  | 6        | 0.42     | 2.35     |
| EP–DSD9  | 100   | 19.37 | 11.81 | 9        | 0.63     | 2.37     |

2.4. Characterization

The chemical structure of DSD was characterized by FTIR–ATR using a PerkinElmer Spectrum 2 (PerkinElmer, Akron, OH, USA), equipped with an attenuated total reflectance (ATR) accessory. 

1H-NMR and 31P-NMR were performed on an Avance spectrometer (Bruker, Rheinstetten, Germany) at room temperature, and DMSO-d6 was used as the solvent.

TGA was performed on a TG 209 F1 thermal analyzer (NETZSCH, Selb, Germany), both in nitrogen and air atmosphere, with an operating temperature of 30 to 800 °C and heating rate of 10 °C/min.

Py-GC/MS was conducted on a Frontier PY-2020iD pyrolyzer connected to a Shimadzu GC–MS QP-2010 Ultra (SHIMADZU, Kyoto, Japan). The chromatographic separation was carried out in a capillary column (HP DB-5MS, 30 m length, 0.25 mm diameter, 0.25 mm thickness, Agilent, Valtbrone, Germany), and the carrier gas (helium) flow rate was 1 mL/min. The temperature of the chromatographic column was firstly held for 3 min at 40 °C, then heated to 300 °C at the rate of 15 °C/min, and finally kept at 300 °C for 10 min.

The limiting oxygen index (LOI) was measured on an oxygen index flammability gauge (ATS FAAR, Milan, Italy) based on ASTM D2863. The specimens were 130 mm in length, 6.5 mm in width, and 3 mm in thickness.

The UL-94 test was carried out according to the UL-94 standard, with specimen sizes of 130, 12.7, and 3 mm.

SEM was used to investigate the internal and external morphology of residue chars, which showed a raised appearance in the UL-94 test and was carried out on a TM-1000 scanning electron microscope (Hitachi, Tokyo, Japan), with 300 times magnification. The residue chars were stripped off by a needle and treated with gold sputtering on a copper station.

Raman spectra measurement was implemented on an SPEX-1430 laser Raman spectrometer (SPEX, Metuchen, NJ, USA), with an argon laser and a size of 633 nm.

The tensile strength was tested on an H10K-S material universal testing machine (Tinius Olsen, Philadelphia, PA, USA), according to ASTM D638-08, at a stretching speed of 5 mm/min. The specimens were 75 mm long, 10 mm wide, and 2.1 mm thick, with a gage length of 30 mm. The results were averaged from eight tests.

3. Results

3.1. Characterization of DSD

The synthesis route of DSD is illustrated in Scheme 1. The intermediate DS was synthesized through aldimine condensation between DDS and salicylaldehyde. DSD was prepared from the addition reaction between the C=N groups of DS and the P–H groups of DOPO. The structure of DSD was characterized by FTIR and NMR.
The FTIR spectra of DOPO and DSD are shown in Figure 1. In the FTIR spectra of DOPO, the absorption peak at 2436 cm\(^{-1}\) was assigned to the stretching vibration of P–H. The peaks at 1591 and 1475 cm\(^{-1}\) were attributed to the stretching vibrations of P–CAr, and stretching vibration peaks of 1278 and 1230 cm\(^{-1}\) (P=O), 1045 and 1014 cm\(^{-1}\) (P–O–C), and 1195 and 900 cm\(^{-1}\) (P–O–CAr) could be observed. As for DS FTIR spectra, the peak at 1628 cm\(^{-1}\) could be attributed to the stretching vibrations of C=N, suggesting that DS was synthesized successfully. In the FTIR spectra of DSD, the absorption peaks were detected at 3269 and 1512 cm\(^{-1}\) (N–H), 1332 cm\(^{-1}\) (C–N), 1591 and 1475 cm\(^{-1}\) (P–CAr), 1285 and 1228 cm\(^{-1}\) (P=O), 1042 and 1004 cm\(^{-1}\) (P–O–C), and 1195 and 919 cm\(^{-1}\) (P–O–CAr). In addition, the peak of P–H, which was observed at 2436 cm\(^{-1}\) in DOPO, was not detected in the spectrum of DSD. This phenomenon means that the –P–H bond in the molecule of DOPO completely participated in the additive reaction with the –C=N– bond in DS.

![Figure 1. FTIR spectra of DOPO, DS, and DSD.](image)

\(^1\)H-NMR and \(^{31}\)P-NMR were conducted on a Bruker Avance spectrometer (400 MHz) at room temperature, using DMSO-d\(_6\) as the solvent. \(^1\)H-NMR (DMSO-d\(_6\), \(\delta\), ppm): 5.22–5.32 (2 H, P–C–H); 5.37–5.48 (2 H, –NH–); 6.44–8.30 (16 H, Ar–H); 9.48–9.73 (2 H, –OH).

The \(^{31}\)P-NMR spectrum of DSD is presented in Figure 2. Obviously, there was a double-peak between 28.51 and 31.17 ppm, corresponding to the two phosphorus atoms in DSD. It was considered that the bulky sulphone and rigid DOPO group existed in the molecular structure of DSD, and their steric hindrance effects caused the unequal phosphorus peaks [2,31].

![Figure 2. \(^1\)H-NMR (a) and \(^{31}\)P-NMR(b) spectra of DSD.](image)

Elemental analysis was also executed to confirm the chemical structure of DSD. Elemental analysis values were as follows: C, 66.29%; H, 4.60%; N, 2.99%; and P, 7.04%. Calculated values were as follows: C, 67.56%; H, 4.31%; N, 3.15%; and P, 6.97%. It is clear that the elemental analysis values of
DSD were in accordance with the calculated values. All of these results indicated that the DSD was synthesized successfully.

3.2. Flame-Retardant Properties of Epoxy Thermosets

The flame-retardant properties of EP and EP–DSD thermosets were evaluated by the LOI and UL-94 test. The relevant data from these tests are listed in Table 2.

Table 2. Flame-retardancy of EP and EP–DSD thermosets.

| Samples    | LOI (%) | UL-94, 3.0 mm bar | Ignition of Cotton | Dripping | Rating |
|------------|---------|-----------------|-------------------|----------|--------|
|            |         | $t_1 + t_2$ a (s) | Yes               |          |        |
| EP         | 24.8    | NR b            | Yes               | Yes      | No rating |
| EP–DSD3    | 30.7    | 11.5 + 5.8      | NO                | NO       | V-1    |
| EP–DSD6    | 34.2    | 5.5 + 4.2       | NO                | NO       | V-0    |
| EP–DSD9    | 35.4    | 3.1 + 2.3       | NO                | NO       | V-0    |

a $t_1$ and $t_2$: average after-flame times after the first and the second flame application. b No record due to burn out.

As shown in Table 2, the LOI value of neat EP was only 24.8%, which continued burning with fire drippings after the first ignition, and the drippings could ignite the cotton. In contrast, the EP/DSD composites showed an improved flame-retardancy, and the LOI values of the EP–DSD thermosets obviously increased from 30.7% to 35.4% with the increasing of the DSD content from 0 to 9 wt %. When the loadings of DSD were more than 6 wt %, the composites passed V-0 ratings in UL-94 tests. These results demonstrate that DSD significantly improved the flame-retardancy of epoxy thermosets with a high efficiency.

The combustion of neat EP and EP–DSD thermosets during the UL-94 test was recorded by a digital camera. The relevant screenshots are shown in Figure 3. Obviously, the neat EP combusted vigorously with flaming drips after the first ignition, which showed that it was highly flammable. In regard to EP–DSD thermosets, the combustion behaviors differed from those of neat EP, and they all extinguished spontaneously. With the increasing of the DSD content, the burning times after the first and second ignition decreased gradually. Moreover, the pyrolytic gases were found to be continuously ejected from the interiors of samples during combustion, and this ejection of gas could even blow the flame out at times. Some studies have also reported this phenomenon, called “blowing-out”, which helps to reduce the combustion time and improve the flame-retardancy of thermosets [32].

3.3. Morphology and Structure of Residual Char after the UL-94 Test

To know more about the combustion behavior of neat EP and EP–DSD thermosets in the UL-94 test, the photographs recorded by a digital camera and SEM micrographs of char residues are shown in Figure 4.
Figure 3. Digital photos of EP thermosets during the UL-94 test.

3.3. Morphology and Structure of Residual Char after the UL-94 Test

To know more about the combustion behavior of neat EP and EP–DSD thermosets in the UL-94 test, the photographs recorded by a digital camera and SEM micrographs of char residues are shown in Figure 4.
As presented in Figure 4a, neat EP possessed a loose and cotton-like char layer with distinct cracks after the UL-94 test. Unlike neat EP, the residual char of EP–DSD thermosets maintained a cuboid morphology, which the UL-94 test required for samples, and the char layer of EP–DSD thermosets was relatively tight and smooth. Furthermore, for EP–DSD thermosets, it should be noted that there were several raised stomates marked with the blue box on the surface of residual char, where pyrolytic gases were found to be ejected during the UL-94 test. It is possible that the loading of DSD enhanced the strength of the char layer, and the strength was strong enough to maintain the shape of raised stomates due to the high internal pressure caused by the accumulation of pyrolytic gases during combustion.

The SEM micrographs of char residues corresponding to the red box in Figure 4a are shown in Figure 4b,c. It is clear that the char residue of neat EP displayed a discontinuous morphology with some cracks. For EP–DSD thermosets, the external char layer showed a tight and intumescent morphology with small holes, and the internal char layer possessed a smooth morphology with some honeycombed cavities. It is presumable that this tight and intumescent char layer with a honeycombed cavity inside acted as an insulation layer, and could block the fluxion of air and heat transfer during combustion [33].

In addition, this insulation layer could also prevent the pyrolytic gases generated inside the samples from escaping. As the amount of pyrolytic gases increased, the inside pressure increased. Under sufficient pressure, the pyrolytic gases broke through the char layer and blew out the flame due to the quenching effect of phosphorus radicals and the diluting effect of nonflammable gases. All of these effects mean that the EP–DSD thermosets achieved a short after-flame time.

3.4. Morphology of Residual Char after Thermal Degradation under Nitrogen Atmosphere

To learn more about the morphology of the char layer, neat EP and EP–DSD thermosets (12.7 mm × 12.7 mm × 3.0 mm) were heated to 600 °C at a heating rate of 10 °C/min under nitrogen atmosphere in a muffle furnace, and naturally cooled to room temperature. A digital photograph of epoxy thermosets before and after thermal degradation is presented in Figure 5.
The intensity of D and G bands was represented by the relevant peak area, and the ratio of the intensity of D and G bands (ID/IG) of neat EP and EP–DSD9 thermosets was calculated by the ratio of (D band) and 1582 cm\(^{-1}\) (G band). Generally, the D band corresponds to carbons in a disorder arrangement, while the G band relates to carbons in graphite layers [9,36,37].

For further investigation, the residual chars for neat EP and EP–DSD9 thermosets after thermal degradation under nitrogen atmosphere were examined by laser Raman spectroscopy. The corresponding Raman spectra of residue char are presented in Figure 6. Both of the two spectra displayed two peaks at 1355 cm\(^{-1}\) (D band) and 1582 cm\(^{-1}\) (G band). Generally, the D band corresponds to carbons in a disorder arrangement, while the G band relates to carbons in graphite layers [9,36,37]. The intensity of D and G bands was represented by the relevant peak area, and the ratio of the intensity of D and G bands (ID/IG) of neat EP and EP–DSD9 thermosets was calculated by the ratio of peat areas. As shown in Figure 6, the ID/IG values of neat EP and EP–DSD9 thermosets were...
1.33, respectively. This means that the residual char of EP–DSD9 thermosets possessed a higher graphitization degree, and thus had a higher heat stability.

![Raman spectra of neat EP and EP–DSD9.](image)

**Figure 6.** Raman spectra of neat EP and EP–DSD9.

### 3.5. Thermal Stability of EP and EP/DSD Thermosets

The thermal stability of neat EP and EP/DSD thermosets was investigated by TGA under air and nitrogen atmosphere, and the TG and DTG curves of epoxy thermosets are shown in Figures 7 and 8, respectively. The values of the initial degradation temperature for 5% weight loss ($T_{5\%}$), the maximum weightlessness rate ($V_{\text{max}}$), the temperature at $V_{\text{max}}$ ($T_{\text{max}}$), and the char yield at $T_{\text{max}}$ and 700 °C ($C_{700}$) are listed in Tables 3 and 4.

![TG (a) and DTG (b) curves of epoxy thermosets under air atmosphere.](image)

**Figure 7.** TG (a) and DTG (b) curves of epoxy thermosets under air atmosphere.

### Table 3. TGA data of epoxy thermosets under air atmosphere.

| Sample   | $T_{5\%}$ (°C) | $T_{\text{max}}$ (°C) | $C_{700}$ (%) | $V_{\text{max}}$ (%/°C) | $C_{700}$ (%) |
|----------|----------------|-----------------------|--------------|-------------------------|--------------|
| EP       | 369.7          | 384.9                 | 79.84        | −1.47                   | 1.21         |
| EP–DSD3  | 361.3          | 375.9                 | 79.92        | −1.25                   | 1.54         |
| EP–DSD6  | 351.1          | 369.7                 | 80.28        | −1.14                   | 2.49         |
| EP–DSD9  | 346.4          | 364.5                 | 82.97        | −1.14                   | 3.98         |

As Figure 7 shows, there were two rapid descents in the TG curve, which corresponded to the two peaks in the DTG curve. These indicated that there were two weight-loss processes in the thermal degradation of all EP thermosets under air atmosphere. However, it was clear that the TG and DTG curves of EP–DSD thermosets were different from those of neat EP. Although the loading of DSD did not change the two-stage thermal degradation of EP thermosets under air atmosphere, it advanced the decomposition time. As shown in Table 3, compared with neat EP, the loading of DSD resulted in a lower $T_{5\%}$ for EP/DSD thermosets, and it decreased from 369.7 °C for neat EP to 346.4 °C for EP–DSD9.
Besides, with the increasing of DSD content, higher char yields at \( T_{\text{max}} \) and 700 °C could be observed when compared with neat EP. This suggests that the introduction of DSD could reduce the weight loss and promote the formation of char residues, and hence improve the thermal stability.

![Figure 8. TG (a) and DTG (b) curves of neat EP and EP–DSD thermosets under nitrogen atmosphere.](image)

**Table 4.** Data of neat EP and EP–DSD thermosets under nitrogen atmosphere.

| Sample    | \( T_{5\%} \) (°C) | \( T_{\text{max}} \) (°C) | \( \text{CY}_{T_{\text{max}}} \) (%) | \( V_{\text{max}} \) (%/°C) | \( \text{CY}_{700} \) (%) |
|-----------|---------------------|-----------------------------|---------------------------------|---------------------------|-----------------------------|
| EP        | 367.2               | 387.2                       | 75.78                           | -1.76                     | 16.58                       |
| EP–DSD3   | 362.2               | 377.2                       | 79.49                           | -1.39                     | 20.48                       |
| EP–DSD6   | 354.6               | 372.1                       | 81.73                           | -1.25                     | 22.74                       |
| EP–DSD9   | 349.7               | 364.7                       | 82.13                           | -1.21                     | 22.87                       |

Under nitrogen atmosphere, the TG curves of all EP thermosets displayed a single decomposition process, as shown in Figure 8, which corresponded to the only peak in the DTG curve. With the increase of DSD content from 0 wt % to 9 wt %, the char residue at 700 °C was gradually increased from 16.58% to 23.45%. Similar to the results in air atmosphere, with the dosage of DSD, the char residual weights were much larger than those of neat EP, proving that an enhanced performance was exhibited by char forming under nitrogen atmosphere. In addition, the \( T_{5\%} \), \( T_{\text{max}} \), and \( V_{\text{max}} \) of EP–DSD thermosets were lower than those of neat EP with the increasing content of DSD.

In summary, EP–DSD thermosets possessed a higher char yield than neat EP in both air and nitrogen atmosphere. The loading of DSD lowered the \( T_{5\%} \), \( T_{\text{max}} \), and \( V_{\text{max}} \) of thermosets. These changes indicated that the loading of DSD contributed to char formation in combustion, which means a better enhancement of the flame-retardancy of epoxy thermosets.

### 3.6. Py-GC/MS Analysis of DSD, EP, and EP–DSD Thermosets

As shown in Figure 9, there were 14 kinds of pyrolysis products for DSD. However, the main pyrolytic products of DSD were benzenamine (c, relative area: 21.98%), diphenyl (e, relative area: 8.29%), dibenzofuran (f, relative area: 11.18%), o-phenylphenol (g, relative area: 36.07%), and fluorene (h, relative area: 1.17%). In consideration of the molecular structure and their relative area, it is reasonable that the breaking of C–N and C–P in DSD resulted in the main pyrolytic products above, and the diphenyl (e), dibenzofuran (f), and o-phenylphenol (g) were decomposition products of DOPO-groups. This suggest that phosphorus fragments existed in the gas phase [38].

Figure 10 shows the pyrograms of neat EP and EP–DSD thermosets; they were similar, but the relative area of EP–DSD thermosets was large. It should be noted that diphenyl (e), dibenzofuran (f), and o-phenylphenol (g) were also observed in EP–DSD pyrograms, which related to decomposition of the DOPO-groups, indicating that phosphorus fragments released in the decomposition of EP–DSD thermosets could inhibit the free radical reaction in the flame in the gas phase.
DSD thermosets decreased with the increasing dosage of DSD. The rigid DOPO group constrained the molecular chain motion [39], and the elongation of EP–DSD and –NH– groups in DSD with DGEBA and DDM. However, the cross-linked network structure attributed to the larger cross-linked network structure, which was formed by the cross-linking of OH and –NH– groups in DSD with DGEBA and DDM. However, the cross-linked network structure raised the $T_g$ from 171.2 to 176.5 °C, complying with the increasing of tensile strength. This could be attributed to the larger cross-linked network structure, which was formed by the cross-linking of OH and –NH– groups in DSD with DGEBA and DDM. However, the cross-linked network structure and the rigid DOPO group constrained the molecular chain motion [39], and the elongation of EP–DSD thermosets decreased with the increasing dosage of DSD.

Table 5. Glass transition temperature ($T_g$) and cross-linking degree values of neat EP and EP–DSD thermosets.

| Sample     | $T_g$ (°C) | $\nu_e$ (mol/m$^3$) |
|------------|------------|---------------------|
| EP         | 171.2      | 1958                |
| EP–DSD3    | 173.6      | 2015                |
| EP–DSD6    | 174.9      | 2067                |
| EP–DSD9    | 176.5      | 2105                |

3.7. Mechanical Properties of Neat EP and EP–DSD Thermosets

When used as a curing agent, the influence of DSD on tensile properties was investigated by measurement of the tensile strength and elongation. The results are exhibited in Figure 10. Neat EP displayed outstanding tensile properties, with a tensile strength of 77.29 MPa and elongation of 8.39% at breaking point. The addition of DSD improved the tensile strength of EP–DSD composites from 84.37 MPa for EP–DSD3 to 88.83 MPa for EP–DSD9. Cross-linking degree values were calculated and are listed in Table 5. They increased from 1958 mol/m$^3$ for neat EP to 2105 mol/m$^3$ for EP–DSD9, and thus raised the $T_g$ from 171.2 to 176.5 °C, complying with the increasing of tensile strength. This could be attributed to the larger cross-linked network structure, which was formed by the cross-linking of OH and –NH– groups in DSD with DGEBA and DDM. However, the cross-linked network structure and the rigid DOPO group constrained the molecular chain motion [39], and the elongation of EP–DSD thermosets decreased with the increasing dosage of DSD.
4. Conclusions

In this study, a novel DOPO-based flame retardant (DSD) was successfully synthesized, and its chemical structure was confirmed by FTIR, $^1$H-NMR, and $^{31}$P-NMR. TGA analysis showed that the loading of DSD improved the thermal stability of EP composites, lowered the $T_{5\%}$ and $T_{\text{max}}$ values of EP thermosets, and increased the char residue to 3.98% and 23.45% under air and nitrogen atmosphere, respectively. The results from the LOI and UL-94 test revealed that the addition of DSD improved the flame retardancy of EP composites. Digital photos and SEM micrographs illustrated that a compact and stable intumescent char layer with a honeycombed cavity inside was formed after combustion, which was responsible for the enhanced flame-retardant properties. Furthermore, the $I_D/I_G$ values of neat EP and EP–DSD9 thermosets, which were calculated according to Raman spectra, indicate that the residual char of EP–DSD9 thermosets possesses a higher graphitization degree, and thus has a higher heat stability. Py-GC/MS analysis showed that phosphorus fragments were released in the decomposition of EP–DSD thermosets, which could inhibit the free radical reaction in the gas phase. The formation of a larger cross-linked network structure due to the cross-linking of the OH and –NH– group in DSD with DGEBA and DDM resulted in an enhancement of the tensile strength for EP-DSD composites.

Author Contributions: Conceptualization, Z.C. and L.L.; methodology, L.L.; software, L.L.; validation, L.L., Z.C.; formal analysis, L.L.; investigation, Z.C.; resources, L.L.; writing—original draft preparation, L.L.; writing—review and editing, Z.C.; visualization, L.L.; supervision, Z.C.; project administration, Z.C.; funding acquisition, L.L. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by Natural Science Foundation of the Jiangsu Higher Education Institutions of China, grant number 19KJD430009, and the Scientific Research Projects of Yancheng Polytechnic College, grant no. ygy1702.

Conflicts of Interest: The authors declare no conflict of interest.

References

1. Jang, J.B.; Kim, T.H.; Kim, T.; Kim, H.J.; Seo, B.; Lim, C.-S.; Lee, W. Modified epoxy resin synthesis from phosphorus-containing polyl and physical changes studies in the synthesized products. *Polymers* 2019, 11, 2116. [CrossRef] [PubMed]

2. Hamciuc, C.; Vlad-Bubulac, T.; Serbezeanu, D.; Carja, I.-D.; Hamciuc, E.; Lisa, G.; Forrat Perez, V. Environmentally friendly fire-resistant epoxy resins based on a new oligophosphonate with high flame retardant efficiency. *RSC Adv.* 2016, 6, 22764–22776. [CrossRef]

3. Iddrissu, I.; Zheng, H.; Rowland, S.M. DC electrical tree growth in epoxy resin and the influence of the size of inceptive AC trees. *IEEE Trans. Dielectr. Electr. Insul.* 2017, 24, 1965–1972. [CrossRef]

4. Wang, N.; Gao, H.; Zhang, J.; Qin, Y.; Wang, D. Phytic acid intercalated graphene oxide for anticorrosive reinforcement of waterborne epoxy resin coating. *Polymers* 2019, 11, 1950. [CrossRef] [PubMed]

5. Xin, D.; Qiang, H. Study on thermomechanical properties of cross-linked epoxy resin. *Mol. Simulat.* 2015, 41, 1081–1085. [CrossRef]

6. Naderi, M.; Hoseinabadi, M.; Najafi, M.; Motahari, S.; Shokri, M. Investigation of the mechanical, thermal, and anticorrosion properties of epoxy nanocomposite coatings: Effect of synthetic hardener and nanoporous graphene. *J. Appl. Polym. Sci.* 2018, 135, 46201. [CrossRef]

7. Kireev, V.V.; Bilichenko, Y.V.; Borisov, R.S.; Mu, J.; Kuznetsov, D.A.; Eroshenko, A.V.; Filatov, S.N.; Sirotin, I.S. Synthesis of bisphenol A based phosphazene-containing epoxy resin with reduced viscosity. *Polymers* 2019, 11, 1914. [CrossRef] [PubMed]

8. Movahedifar, E.; Vahabi, H.; Saeb, M.R.; Thomas, S. Flame retardant epoxy composites on the road of innovation: An analysis with flame retardancy index for future development. *Molecules* 2019, 24, 3964. [CrossRef]

9. Vahabi, H.; Laoutid, F.; Movahedifar, E.; Khalili, R.; Rahmati, N.; Vagner, C.; Cochez, M.; Brison, L.; Ducos, F.; Ganjali, M.R.; et al. Description of complementary actions of mineral and organic additives in thermoplastic polymer composites by flame retardancy index. *Polym. Adv. Technol.* 2019, 30, 2056–2066. [CrossRef]
10. Vahabi, H.; Saeb, M.R.; Formela, K.; Cuesta, J.-M.L. Flame retardant epoxy/halloysite nanotubes nanocomposite coatings: Exploring low-concentration threshold for flammability compared to expandable graphite as superior fire retardant. *Prog. Org. Coat.* 2018, 119, 8–14. [CrossRef]

11. Vahabi, H.; Jouyandeh, M.; Cochez, M.; Khalili, R.; Vagner, C.; Ferriol, M.; Movahedifar, E.; Ramezanzadeh, B.; Rostami, M.; Ranjar, Z.; et al. Short-lasting fire in partially and completely cured epoxy coatings containing expandable graphite and halloysite nanotubes additives. *Prog. Org. Coat.* 2018, 123, 160–167. [CrossRef]

12. Sari, M.G.; Saeb, M.R.; Shabanian, M.; Khaleghi, M.; Vahabi, H.; Vagner, C.; Zarrintaj, P.; Khalili, R.; Paran, S.M.R.; Ramezanzadeh, B. Epoxy/starch-modified nano-zinc oxide transparent nanocomposite coatings: A showcase of superior curing behavior. *Prog. Org. Coat.* 2018, 115, 143–150. [CrossRef]

13. Rastin, H.; Saeb, M.R.; Nonahal, M.; Shabanian, M.; Vahabi, H.; Formela, K.; Gabrion, X.; Seidi, F.; Zarrintaj, P.; Sari, M.G. Transparent nanocomposite coatings based on epoxy and layered double hydroxide: Nonisothermal cure kinetics and viscoelastic behavior assessments. *Prog. Org. Coat.* 2017, 113, 126–135. [CrossRef]

14. Chen, L.; Wang, Y.Z. A review on flame retardant technology in China. Part I: Development of flame retardants. *Polym. Adv. Technol.* 2009, 21, 1–26. [CrossRef]

15. Levchik, S.V.; Weil, E.D. A review of recent progress in phosphorus-based flame retardants. *J. Fire Sci.* 2006, 24, 345–364. [CrossRef]

16. Rakotomalala, M.; Wagner, S.; Doering, M. Recent developments in halogen free flame retardants for epoxy resins for electronic and electronic applications. *Materials* 2010, 3, 4300–4327. [CrossRef] [PubMed]

17. Wang, L.; Jiang, J.; Jiang, P.; Yu, J. Synthesis, characteristic of a novel flame retardant containing phosphorus, silicon and its application in ethylene vinyl-acetate copolymer (EVM) rubber. *J. Polym. Res.* 2010, 17, 891–902. [CrossRef]

18. Wang, X.; Feng, N.; Liu, L.; Wang, Z.; Wen, Y. Preparation of Lig-based intumescent flame retardant and its effect on flame resistance of TPO. *China Synth. Resin Plast.* 2017, 34, 6–10.

19. Benelli, T.; Mazzocchetti, L.; D’Angelo, E.; Lanzi, M.; Saraga, F.; Sambri, L.; Franchini, M.C.; Giorgini, L. New nitrogen-rich heterocycles for organo-modified bentonites as flame retardant fillers in epoxy resin nanocomposites. *Polym. Eng. Sci.* 2017, 57, 621–630. [CrossRef]

20. Duan, H.; Ji, S.; Yin, T.; Tao, X.; Chen, Y.; Ma, H. Phosphorus–nitrogen-type fire-retardant vinyl ester resin with good comprehensive properties. *J. Appl. Polym. Sci.* 2019, 136, 47997. [CrossRef]

21. Yang, H.; Song, L.; Tai, Q.; Wang, X.; Yu, B.; Yuan, Y.; Hu, Y.; Yuen, R.K.K. Comparative study on the flame retarded efficiency of melamine phosphate, melamine phosphite and melamine hypophosphite on poly(butylene succinate) composites. *Polym. Degrad. Stab.* 2014, 105, 248–256. [CrossRef]

22. Jian, R.; Wang, P.; Xia, L.; Yu, X.; Zheng, X.; Shao, Z. Low-flammability epoxy resins with improved mechanical properties using a Lewis base based on phosphaphenanthrene and 2-aminothiazole. *J. Mater. Sci.* 2017, 52, 1–15. [CrossRef]

23. Xie, M.; Zhang, S.; Ding, Y.; Wang, F.; Liu, P.; Tang, H.; Wang, Y.; Yang, M. Synthesis of a heat-resistant DOPO derivative and its application as flame-retardant in engineering plastics. *J. Appl. Polym. Sci.* 2017, 134, 44892. [CrossRef]

24. Wang, Z.; Liang, B. Synthesis of a new phosphorus-nitrogen flame retardant GAP-DOPO and its flame retardant performance for epoxy resin. *Fine Chem.* 2017, 34, 382–388.

25. Shi, Y.; Yu, B.; Zheng, Y.; Yang, J.; Hu, Y. Design of reduced graphene oxide decorated with DOPO-phosphanomidate for enhanced fire safety of epoxy resin. *J. Colloid Interf. Sci.* 2018, 521, 160–171. [CrossRef] [PubMed]

26. Fang, Y.; Zhou, X.; Xing, Z.; Wu, Y. Flame retardant performance of a carbon source containing DOPO derivative in PET and epoxy. *J. Appl. Polym. Sci.* 2017, 134. [CrossRef]

27. Xu, M.-J.; Xu, G.-R.; Leng, Y.; Li, B. Synthesis of a novel flame retardant based on cyclotriphenazene and DOPO groups and its application in epoxy resins. *Polym. Degrad. Stabil.* 2016, 123, 105–114. [CrossRef]

28. Ping, C.; Shengping, L.; Dezhong, W. *Epoxy Resin and Its Application*; Chemical Industry Press: Beijing, China, 2011.

29. Tan, Y.; Shao, Z.-B.; Chen, X.-F.; Long, J.-W.; Chen, L.; Wang, Y.-Z. A novel multifunctional organic-inorganic hybrid curing agent with high flame-retardant efficiency for epoxy resin. *ACS Appl. Mater. Interfaces* 2015, 7, 17919–17928. [CrossRef]
30. Yang, J.P.; Chen, Z.-K.; Yang, G.; Fu, S.-Y.; Ye, L. Simultaneous improvements in the cryogenic tensile strength, ductility and impact strength of epoxy resins by a hyperbranched polymer. *Polymer* 2008, 49, 3168–3175. [CrossRef]

31. Liu, C.; Chen, T.; Yuan, C.H.; Song, C.F.; Chang, Y.; Chen, G.R.; Xu, Y.T.; Dai, L.Z. Modification of epoxy resin through the self-assembly of a surfactant-like multi-element flame retardant. *J. Mater. Chem. A* 2016, 4, 3462–3470. [CrossRef]

32. Zhang, W.; Li, X.; Yang, R. Pyrolysis and fire behaviour of epoxy resin composites based on a phosphorus-containing polyhedral oligomeric silsesquioxane (DOPO-POSS). *Polym. Degrad. Stabil.* 2011, 96, 1821–1832. [CrossRef]

33. He, X.; Zhang, W.; Yang, R. The characterization of DOPO/MMT nanocompound and its effect on flame retardancy of epoxy resin. *Compos. Part A Appl. Sci. Manuf.* 2017, 98, 124–135. [CrossRef]

34. Wan, J.; Gan, B.; Li, C.; Molina-Aldareguia, J.; Kalali, E.N.; Wang, X.; Wang, D.-Y. A sustainable, eugenol-derived epoxy resin with high biobased content, modulus, hardness and low flammability: Synthesis, curing kinetics and structure-property relationship. *Chem. Eng. J.* 2016, 284, 1080–1093. [CrossRef]

35. Lin, C.H.; Chou, Y.C.; Shiao, W.F.; Wang, M.W. High temperature, flame-retardant, and transparent epoxy thermosets prepared from an acetonilavolone-based hydroxyl poly(ether sulfone) and commercial epoxy resins. *Polymer* 2016, 97, 300–308. [CrossRef]

36. Wang, Z.; Wu, W.; Zhong, Y.; Ruan, M.; Hui, L.L. Flame-retardant materials based on phosphorus-containing polyhedral oligomeric silsesquioxane and bismaleimide/diallylbisphenol a with improved thermal resistance and dielectric properties. *J. Appl. Polym. Sci.* 2015, 132, 41545. [CrossRef]

37. Jiang, S.-D.; Bai, Z.-M.; Tang, G.; Song, L.; Stec, A.A.; Hull, T.R.; Hu, Y.; Hu, W.-Z. Synthesis of mesoporous silica@Co−Al layered double hydroxide spheres: Layer-by-layer method and their effects on the flame retardancy of epoxy resins. *ACS Appl. Mater. Interfaces* 2014, 6, 14076–14086. [CrossRef]

38. Schartel, B.; Braun, U.; Balabanovich, A.I.; Artner, J.; Ciesielski, M.; Doering, M.; Perez, R.M.; Sandler, J.K.W.; Altstaedt, V. Pyrolysis and fire behaviour of epoxy systems containing a novel 9,10-dihydro-9-oxa-10-phosphaphenanthrene-10-oxide-(DOPO)-based diamino hardener. *Eur. Polym. J.* 2008, 44, 704–715. [CrossRef]

39. Kopwyhoba, A.H.; Korshunova, A.N. The peculiarities of polaron motion in the molecular polynucleotide chains of finite length in the presence of localized excitations in the chain. *Mat. Biolog. Bioinform.* 2017, 11, 141–158. [CrossRef]

© 2020 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (http://creativecommons.org/licenses/by/4.0/).