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# **A new reactive iso-eugenol based phosphate flame retardant: toward green epoxy resins**

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**Keywords:** iso-eugenol, phosphate flame retardant, bio-based resins, epoxy thermosets, high performance.

## **ABSTRACT:**

A bio-based reactive phosphate flame retardant derived from iso-eugenol was synthesized and fully characterized (<sup>1</sup>H, <sup>13</sup>C, <sup>31</sup>P NMR, FTIR, MS) with the aim of improving flame retardancy

behavior of bio-based epoxy thermosets. This new green flame retardant, diepoxy-iso-eugenol phenylphosphate (DEpiEPP) was then copolymerized either with conventional diglycidylether of bisphenol A resin (DGEBA) or with a totally bio-based glycidylether epoxy iso-eugenol (GEEpiE) resin using a bio-based camphoric anhydride (CA) as hardener. Resulting thermosets with varying rates of flame retardant (1.0 to 4.3 w% phosphorus) were then characterized (FTIR, DSC, nanoindentation, 3-point bending test, TGA, PCFC, cone calorimeter). By this way, a new solution was proposed allowing (i) to increase the bio-based content of thermosets (ii) to improve the flame retardancy properties by a reactive way and (iii) to provide epoxy thermosets with high T<sub>g</sub>, excellent mechanical properties and a curing temperature compatible with the use of vegetable fibers. For instance, with a weight content of 2.0 w% P, both GEEpiE-CA-DEpiEPP (95% bio-based content) and partially bio-based DGEBA-CA-DEpiEPP (57% bio-based content) epoxy thermosets possess high T<sub>g</sub> (respectively 129 and 105°C), bending elastic modulus and strength of respectively 4.08 GPa and 90 MPa and demonstrate good flame-retardant properties due to char promotion showing high promise for application.

## **INTRODUCTION**

Epoxy thermosets are widely applied and constitute a large part of the thermosetting materials due to their high performances.<sup>1</sup> Their chemical backbone often contains aromatic rings providing good mechanical and thermal behaviors. They offer useful adhesion and mechanical properties exhibiting also interesting chemical, electrical and heat resistances. Thus, these materials are also widely used in cutting-edge technological fields, such as aeronautics, automotive, electronic components and sports. However, most of the commonly used epoxy thermosetting polymers are petroleum-based. Furthermore, it is noteworthy that 90% of epoxy cross-linked polymers are synthesized from DGEBA (diglycidylether of bisphenol A). This resin is produced from BPA (bisphenol A) reactant,<sup>2</sup> an endocrine disruptor and nephrotoxic

compound that has received much attention due to its negative health effects<sup>3</sup>. Thus, in the view of decreasing the environmental impact, saving petroleum resources and replacing toxic BPA for the synthesis of epoxy, bio-based thermosets are widely studied. During the last decades many research works have therefore been dedicated to the specific domain of bio-based epoxy thermosets pointing out the potential of rosin,<sup>4</sup> cardanol,<sup>5</sup> iso-eugenol,<sup>6,7</sup> eugenol,<sup>8-12</sup>, vegetable oil,<sup>13</sup> modified lignin,<sup>14-16</sup> vanillin,<sup>17-19</sup> terpene derivatives,<sup>20</sup> daidzein,<sup>21</sup> ... to replace DGEBA epoxy monomer. Among the most promising feedstocks to replace BPA, building blocks derived from lignin, have been particularly pointed out.<sup>22-28</sup> Indeed, with the objective of future industrial developments, accessibility and availability of the raw components are crucial parameters and have to be considered. Thus, lignin which is produced in large quantities (about 50 million tons of lignin per year are produced worldwide by the pulp and paper industry) constitutes the most abundant natural resource able to provide components containing rigid aromatic rings precursors of high performances thermosets<sup>29</sup>. Within this framework, Barta et al<sup>30</sup> recently reviewed new platform chemicals arising from lignin.

In parallel, different studies also underlined the importance of substituting conventional toxic polyamine hardeners by anhydrides curing agents, potentially bio-based and containing rigid ring leading to a reinforcement of the thermosets mechanical properties.<sup>6,19,20,31,32</sup>

However, epoxy polymers are known to be highly flammable and to release a large amount of toxic smoke during their combustion.<sup>33</sup> Thus, different approaches more mindful of health and of the environment were recently studied to improve the flame retardancy of epoxy thermosets in order to fulfill the flammability requirements for epoxy thermosets applications.

The conventional approach is to add flame retardants (FRs) to epoxy thermosets. Most of these FRs contain phosphorus rather than halogen due to their toxic effect. Phosphorus FR act as char promoter in condensed phase and/or as flame inhibitor in gas phase.<sup>34</sup> They can be phosphine oxide, phosphinate, phosphonate or phosphate and it is generally considered that their modes-

of-action strongly depend on their oxidation level.<sup>35, 36</sup> Especially phosphate and phosphonate are rather effective in condensed phase by promoting charring of polymers.

Among these additives, DOPO and its derivatives have been extensively studied.<sup>37,38</sup> Recently most of the studies were dedicated to the synthesis of new bio-based phosphorus-containing FRs. Howell and Sun<sup>39</sup> have reported the beneficial effect of different bio-based FRs (molecules and oligomers) synthesized from tartaric acid and used as additive in a DGEBA matrix. Recently Wang et al.<sup>5</sup> also used a cardanol-based phosphate as bio-based FR (Phosphaneanthrene groups-containing TrisCardanol Phosphate PTPC). The resulting thermosets (DGEBA-PTPC-aromatic diamine) showed that the FR addition accelerated the thermal degradation process and improved the char yield.

Nevertheless, the addition of non-reactive organophosphorus FRs of low molecular weight presents several drawbacks in connection with their potential volatility, causing their partial loss from the polymer by migration to the surface during the processing or ageing or in the early stages of combustion. Moreover, they can also act as plasticizers, leading to a decrease of the thermoset mechanical properties and/or also give rise to the formation of a segregated microphase.<sup>40</sup> Furthermore, in the case of DOPO addition, an unintended consumption of epoxy group was reported, leading to a lower network density and a subsequent decrease of the thermoset T<sub>g</sub>.<sup>41,42</sup>

Thus, the route which uses reactive organophosphorus FRs monomer is preferred in order to maximize the fire retardancy and to reduce the evolution of toxic species. Wang et al.<sup>43-45</sup> have prepared epoxy resins containing phosphorus groups and some other groups (piperidine, imidazole, benzimidazole) to combine char promotion, radical quenching and fuel dilution effects. Phosphorus ionic liquids were also incorporated into epoxy network to impart flame retardancy.<sup>46,47</sup> To increase the biobased content of materials, bioresources were also functionalized with phosphorus groups to be incorporated as reactive flame retardants. Various

phosphorus groups were grafted to biomolecules derived from carbohydrates<sup>48</sup>, lipids<sup>49</sup>, terpenes<sup>49</sup>, phenols<sup>50-53</sup> and so on. Biophenols are especially interesting due to the presence of aromatic rings. Ecochard et al.<sup>54</sup> have shown that phosphorylated eugenol was more suitable than cardanol to improve the flame retardancy of epoxy resins, due to the aliphatic chains of cardanol limiting the charring. Within this framework, Wang and co-authors<sup>13</sup> have synthesized two phosphorus-containing dicarboxylic acids FRs used as hardeners with epoxidized soybean oil with the aim of obtaining Pressure-Sensitive Adhesives having low T<sub>g</sub>. The curing reaction was carried out at 180°C and authors reported a lower reactivity of the phosphorus-containing hardeners. Ma and coauthors<sup>48</sup> have also reported the benefit in using a reactive phosphorus-containing diepoxy FR synthesized from itaconic acid (EADI). This diepoxy FR was then copolymerized with DGEBA using MHPA as hardener. Both these studies highlight the enhanced fire retardancy properties of the resulting thermosets induced by a charring effect, and this behavior depends on the environment of the phosphorus compound (aliphatic/aromatic/ester bonds).

It is noteworthy that the lower reactivity of both these phosphorus containing derivatives needs more severe curing conditions with higher curing temperature ( $T > 180^{\circ}\text{C}$ ) making these matrices no longer compatible with the use of vegetable fibers.

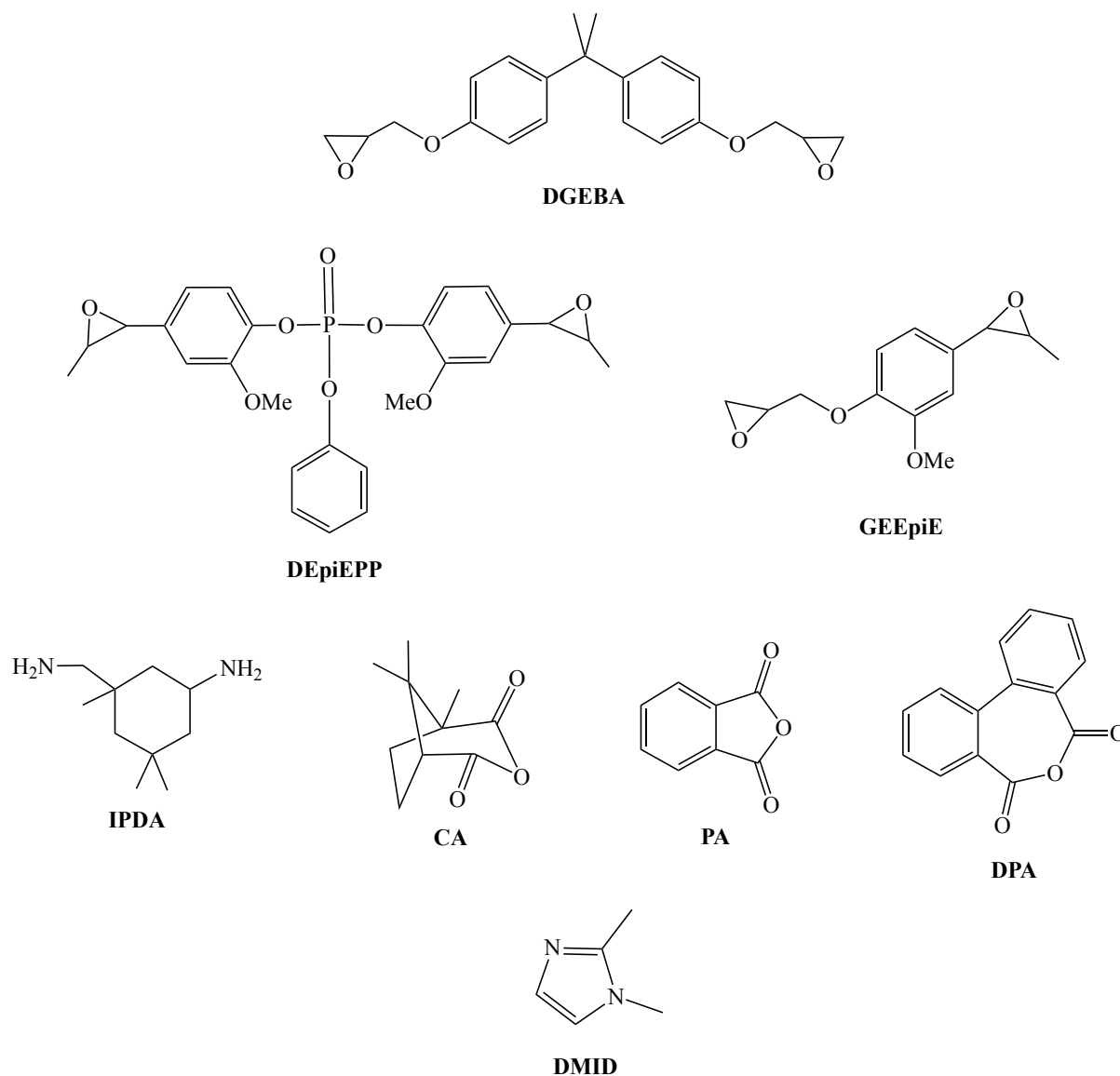
Furthermore, some of the solutions currently proposed consist in using intrinsic fire retardancy properties of partially bio-based epoxy thermosets. Wan and al.<sup>10,55</sup> demonstrated that high performance flame retardancy epoxy thermosets were obtained using an eugenol derivative epoxy resin with an aromatic polyamine. Li and co-authors<sup>56</sup> also showed enhanced flame retardancy when silicon bridged difunctional epoxy monomers derived of eugenol were also cured with an aromatic polyamine (diamino diphenyl sulfone DDS). The high curing temperatures are incompatible with the use of vegetable fibers (2h at 150°C and then 2h at 190°C).

Thus, with the aim of developing fully bio-based composites containing vegetable fibers, our research team recently described a bio-based epoxy thermoset made of glycidylether epoxy iso-eugenol (GEEpiE) resin cured with camphoric anhydride (CA) at 150°C. This totally bio-based material well fits with the mechanical properties required for structural composite applications using vegetable fibers as highlighted by their high instantaneous modulus of about 4.8 GPa and curing temperatures.<sup>6</sup> In our ongoing studies on the bio-based epoxy thermosets for high performance composite applications, we herein report the two steps synthesis of a reactive bio-based phosphate flame retardant derived from iso-eugenol. Phenyldichlorophosphate first reacts with bio-based iso-eugenol according to a Williamson reaction. The second step consists of an oxidation of the alcene function by Oxone® as previously reported<sup>6</sup> leading to the formation of the diepoxy-iso-eugenol phenylphosphate (DEpiEPP). The chemical structure of DEpiEPP was confirmed by <sup>1</sup>H, NMR, <sup>13</sup>C NMR, <sup>31</sup>P NMR, FTIR and mass spectrometry (MS). Then the copolymerization of DEpiEPP either with conventional diglycidylether of bisphenol A resin (DGEBA) or with the totally bio-based glycidylether epoxy iso-eugenol (GEEpiE) using the bio-based camphoric anhydride (CA) as hardener was monitored by differential scanning calorimetry DSC and confirmed by FTIR. Finally, thermal (DSC) flame retardant (TGA, PCFC, cone calorimeter) and mechanical (nanoindentation, 3-point bending test) properties of the resulting thermosets containing varying rates of flame retardant (1 to 4.3 w% phosphorus) were investigated.

## EXPERIMENTAL SECTION

**Materials.** Triethylamine (TEA, ≥99% purity, Sigma-Aldrich) and toluene (analytical grade, Carlo Erba) were purified by distillation. Dichloromethane (DCM, analytical grade, Carlo Erba) and acetone (analytical grade, Carlo Erba) were purchased and used as received. Phenyldichlorophosphate (PPDC, ≥95% purity, Sigma-Aldrich), sodium hydrogen carbonate

( $\text{NaHCO}_3$ ,  $\geq 95\%$  purity, Sigma-Aldrich), tetrabutylammonium hydrogen sulfate ( $\text{Bu}_4\text{NHSO}_4$ ,  $\geq 97\%$  purity, Fluka) and Oxone® (Sigma-Aldrich) were purchased and used as supplied. Diepoxy synthon derived from *iso*-eugenol (GEEpiE) was prepared from biobased *iso*-eugenol according to a previously published procedure.<sup>7</sup> Diglycidyl ether of bisphenol A (DGEBA, Sigma-Aldrich), isophorone diamine (IPDA,  $\geq 99\%$ , Sigma-Aldrich), camphoric anhydride (CA, 98% purity, Alfa Aesar) and 1,2-Dimethylimidazole (DMID, 95% purity, Sigma-Aldrich) were purchased and used without any further purification. Phthalic anhydride (PA), diphenic anhydride (DPA) were prepared according to published procedures.<sup>8,57</sup> Molecular representations of reagents used in the resin manufacturing process are presented in scheme 1.





**Scheme 1.** Molecular representations of reagents used. Epoxy monomers: diglycidylether of bisphenol A (DGEBA), glycidylether of phenoxy iso-eugenol (GEEpiE). Flame retardants: diepoxyiso-eugenolphenylphosphate (DEpiEPP). Curing agents: isophorone diamine (IPDA), camphoric anhydride (CA), phthalic anhydride (PA) and diphenic anhydride (DPA). Catalyst: 1,2-dimethylimidazole (DMID).

### ***Synthesis of glycidylether epoxy of iso-eugenol (GEEpiE)***

GEEpiE was prepared according to published procedure<sup>7</sup> as follows (Scheme 2(a)).

### ***Synthesis of Di(iso-eugenol)phenylphosphate (DiEPP)***

Toluene (25 mL), triethylamine (3.24 mL, 23.26 mmol) and iso-eugenol (3.71 mL, 24.4 mmol) were introduced under argon atmosphere at  $-10\text{ }^{\circ}\text{C}$  into a three-necked flask (100 mL) equipped with a digital thermometer, stirrer, refrigerant and dropping funnel. PPDC (1.76 mL, 11.74 mmol) was added dropwise for 15 min. Then, the reaction mixture was slowly brought to room temperature and stirring continued overnight. The mixture was then solubilized in 10 mL of toluene and washed thrice with water and brine (35 mL). Organic phase was dried on anhydrous  $\text{MgSO}_4$ . The toluene was driven off under reduced pressure to get a yellow viscous liquid. DiEPP was obtained with 74% yield.

IR (ATR,  $\text{cm}^{-1}$ ): 3019 (aromatic C-H), 1655 (C=C), 1589 (aromatic C=C), 1299 (P=O), 1030 (P-O-C).

$^1\text{H-NMR}$  (500 MHz,  $\text{CDCl}_3$ ):  $\delta$  ppm 1.79 (m,  $\text{H}_1$ ), 3.66 (m,  $\text{H}_7$ ), 6.09 and 6.25 (m,  $\text{H}_2$  and  $\text{H}_3$ ), 6.77 and 7.12 (m,  $\text{H}_{4-6}$ ,  $\text{H}_{8-10}$ ).

$^{13}\text{C-NMR}$  (125 MHz,  $\text{CDCl}_3$ ):  $\delta$  ppm 18.5 (d,  $\text{C}_1$ ), 55.8 (d,  $\text{C}_{10}$ ), 109.9 (s,  $\text{C}_9$ ), 118.3 (d,  $\text{C}_6$ ); 120.4 (d,  $\text{C}_{12}$ ), 121.3 (d,  $\text{C}_2$ ), 125.2 (s,  $\text{C}_{14}$ ), 126.1 (s,  $\text{C}_5$ ), 129.5 (s,  $\text{C}_{13}$ ), 130.3 (s,  $\text{C}_3$ ), 138.7 (d,  $\text{C}_8$ ), 150.6 (d,  $\text{C}_{11}$ ), 150.9 (d,  $\text{C}_7$  and  $\text{C}_{11}$ ).

$^{31}\text{P}$ -NMR (202 MHz,  $\text{CDCl}_3$ ),  $\delta$  ppm -16.7 (s, P).

***Synthesis of Diepoxy iso-eugenol phenylphosphate (DEpiEPP)***

DiEPP (4.00 g, 8.58 mmol) solubilized in DCM (50 mL), water (50 mL), acetone (6.6 mL),  $\text{NaHCO}_3$  (3.71 g, 44.16 mmol) and  $\text{Bu}_4\text{NHSO}_4$  (0.15 g, 0.44 mmol) were introduced into a bi-necked flask (100 mL) equipped with a digital thermometer, stirrer, refrigerant and dropping funnel. A solution of Oxone® (8.11 g, 26.38 mmol) in water (50 mL) was added dropwise at  $0^\circ\text{C}$  for 30 min to the biphasic mixture vigorously stirred. Then, the reaction mixture was slowly brought to room temperature and stirring continued overnight.

The organic phase was washed thrice with saturated NaCl water (50 mL) and dried on anhydrous  $\text{MgSO}_4$ . DCM was driven off under reduced pressure to get a yellow-brown very viscous liquid. DEpiEPP was obtained with 52% yield.

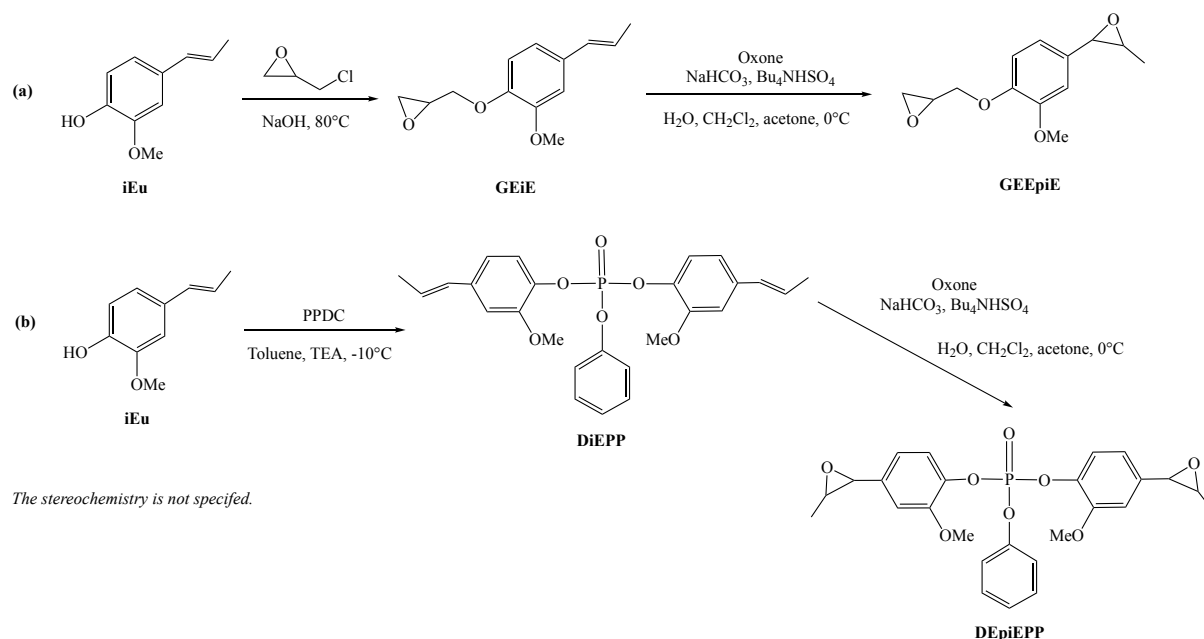
IR (ATR,  $\text{cm}^{-1}$ ): 3068 (aromatic C-H), 1599 (aromatic C=C), 1299 (P=O), 1025 (P-O-C), 904 (epoxy ring), disappearance of (C=C) characteristic band at 1655.

$^1\text{H}$ -NMR (500 MHz,  $\text{CDCl}_3$ ):  $\delta$  ppm 1.08 (d,  $\text{H}_1$ ), 1.45 (d,  $\text{H}_1$ ), 2.96 (m,  $\text{H}_2$ ), 3.31 (m,  $\text{H}_2$ ), 3.54 (d,  $\text{H}_3$ ), 3.74 (s,  $\text{H}_7$ ), 6.82 and 7.30 (m,  $\text{H}_{4-6}$ ,  $\text{H}_{8-10}$ ).

$^{13}\text{C}$ -NMR (125 MHz,  $\text{CDCl}_3$ ):  $\delta$  ppm 17.8 (s,  $\text{C}_1$ ), 55.9 (s,  $\text{C}_2$ ), 57.2 (s,  $\text{C}_3$ ), 59.1 (s,  $\text{C}_{10}$ ), 109.4 (d,  $\text{C}_9$ ), 118.0 (d,  $\text{C}_6$ ), 120.3 (d,  $\text{C}_{12}$ ), 121.3 (d,  $\text{C}_{14}$ ), 125.2 (s,  $\text{C}_5$ ), 129.5 (s,  $\text{C}_{13}$ ), 136.1 (s,  $\text{C}_4$ ), 139.5 (d,  $\text{C}_8$ ), 150.8 (m,  $\text{C}_7$  and  $\text{C}_{11}$ ).

$^{31}\text{P}$ -NMR (202 MHz,  $\text{CDCl}_3$ ):  $\delta$  ppm -16.8 (s, P).

MS (m/z, ES+,  $[\text{M}+\text{H}^+]$ ) calculated 498.468, found: 499.01.



**Scheme 2.** Description of the diepoxy monomers synthesis.

**NB:** It is noteworthy that epichlorohydrin, used for the GEEpiE synthesis and DCM, used as solvent are highly toxic chemicals. They must be used referring to hazard caution information and replaced if possible.

### ***Thermosets curing protocol***

Polyepoxy polymers were obtained *via* thermal curing according to the protocol previously described.<sup>6</sup> Diepoxy monomers containing different molar ratio of synthesized FR (DEpiEPP) were ground in a mortar at room temperature with anhydride hardeners (epoxide function/anhydride function molar ratio 1/0.6) until a homogeneous mixture was obtained. DMID catalyst was then added with a ratio of 0.025 eq. with respect to epoxide function. The mixture was cured in an oven at 150°C during 2 h. The curing protocol for diepoxy monomer (DGEBA) and diamine hardener (IPDA) was the same, with an epoxide/amine molar ratio of 1/0.5.

For cone calorimeter samples, the reactants were first melted at 120 °C until a homogeneous mixture was obtained and then DMID catalyst was added. Epoxy samples were obtained after curing in an oven at 120 °C during 45 min and post-curing at 150°C for 1h.

**Characterizations and instruments.** IR spectra were recorded using an FTIR Bruker Alpha spectrophotometer fitted with an ATR module ALPHA–P equipped with a mono-reflexion diamond crystal in the 4000 - 375 cm<sup>-1</sup> wavenumber range (32 scans at a spectral resolution of 4 cm<sup>-1</sup>).

<sup>1</sup>H, <sup>13</sup>C and <sup>31</sup>P Nuclear magnetic resonance (NMR) spectra were obtained using a Bruker Avance III 500 MHz spectrometer working at 500, 125 and 202 MHz, respectively, at room temperature and using CDCl<sub>3</sub> as solvent. External references were trimethylsilane (TMS) for <sup>1</sup>H and <sup>13</sup>C NMR and phosphoric acid (H<sub>3</sub>PO<sub>4</sub>) for <sup>31</sup>P NMR.

MS analysis was carried out in an HPLC Ultimate 3000 RSLC instrument from DIONEX coupled through an ElectroSpray Ionisation (ESI) interface to an ion-trap AmaZon SL mass spectrometer from Bruker for MS detection. Methanol was used as solvent.

Phosphorus elemental analyses were performed with inductively coupled plasma optical emission spectrometers ICP-OES (DUO) ICAP 7400. The cone calorimeter residue of the DEpiEu-CA series thermosets (0.0% - 2.0% - 3.0% P) containing DEpiEPP was mineralized by microwave using 4 mL of HNO<sub>3</sub> and water up to 10 mL. Thus, the sample solution was nebulized and introduced into the center of the plasma.

DSC analyses were performed on a TA Instruments Discovery DSC under nitrogen flow (50 mL min<sup>-1</sup>) with a sample masse of 10 ±3 mg. To study curing reaction, samples were heated from 0 °C to 230 °C at a heating rate of 5 °C min<sup>-1</sup>. The determination of glass transition temperatures (T<sub>g</sub>) was carried out by heating the cured resin from –20 °C to 240 °C at a heating rate of 20 °C min<sup>-1</sup> and using the TA Universal Analysis 2000 software.

Thermogravimetric analyses were carried out using a TA Instruments Discovery TGA under either N<sub>2</sub> atmosphere (75 mL min<sup>-1</sup>). Samples (8-15 mg) were heated from room temperature to 800 °C at a heating rate of 20 °C min<sup>-1</sup>. Weight loss percentages were determined using the TA Universal Analysis 2000 software. T<sub>5%</sub> and T<sub>max</sub> represent the temperature at which the material loses 5% of its initial mass and the temperature of the peak of mass loss rate respectively. Char content was measured at 600 °C.

Flammability was assessed using a Pyrolysis-Combustion Flow Calorimeter (PCFC) from Fire Testing Technology (FTT UK). About 3 mg was placed in the pyrolyzer, undergoing an increase of temperature from 20 °C to 750 °C at 1 °C s<sup>-1</sup> under a nitrogen flow. Decomposition gases were sent to a combustor heated at 900 °C under air flow (N<sub>2</sub>/O<sub>2</sub> = 80/20). At this temperature and with 20% of oxygen, combustion was considered to be complete.<sup>58</sup> Heat release rate was determined by oxygen depletion according to Huggett principle<sup>59</sup> (1 kg of consumed oxygen corresponds to 13.1 MJ of heat released). pHRR is the maximal value of the heat release rate. The total heat released (THR) was obtained by integration of PCFC curves. Heat of combustion ( $\Delta h$ ) is the ratio between THR and weight loss fraction.

Thermosets fire behavior was also studied using a cone calorimeter (Fire Testing Technology) which is a powerful tool to investigate the fire behavior of polymers. A horizontal cylindrical sample sheet of  $\phi$  77 × 3 mm was isolated by rock wool and placed at 6 cm below the conic heater because of the intumescent char formation observed. The samples were exposed to a 35 kW m<sup>-2</sup> heat flux in well-ventilated conditions (air rate 24 L s<sup>-1</sup>) in the presence of a spark igniter to force the ignition. HRR was determined according to oxygen depletion (Huggett's relation) as in PCFC. This test was performed according to the ISO 5660 standard.

Nanoindentation experiments were performed using an ultra nanoindenter from Anton Paar. A Berkovich indenter was used. The area function was calibrated by indenting in a fused silica sample with a known Young's modulus and Poisson's ratio. Before carrying out the tests, the

samples were polished with SiC paper, alumina particles until 0.03 mm and then colloidal silica. The instantaneous indentation modulus EI and hardness H at different indentation depths, the relaxed indentation modulus ER and finally the strain rate sensitivity m were determined from multicycle indentations, indentation at different constant strain rate  $\dot{h}/h$  (where h is the indentation depth) and trapezoidal holding force tests with linear loading and unloading segments. The indentation protocols were the same as used previously.<sup>7</sup>

3-point bending tests were also performed using a universal testing machine MTS criterion 45 equipped with a 1 kN full-range load sensor. Three samples of each batch were tested at a constant crosshead displacement rate of 1 mm/min. The total length of the specimen was approximately 50 mm, and their width and thickness 3.5 mm and 2 mm, respectively. The deflection was measured at mid-span, on the bottom face of the specimen, using a micrometer laser sensor. The support span was 40 mm. The ultimate stress ( $\sigma_{\max}$ ), the 3 points bending modulus (E) and the strain at failure ( $\epsilon$ ) were determined from the load-deflection curves.

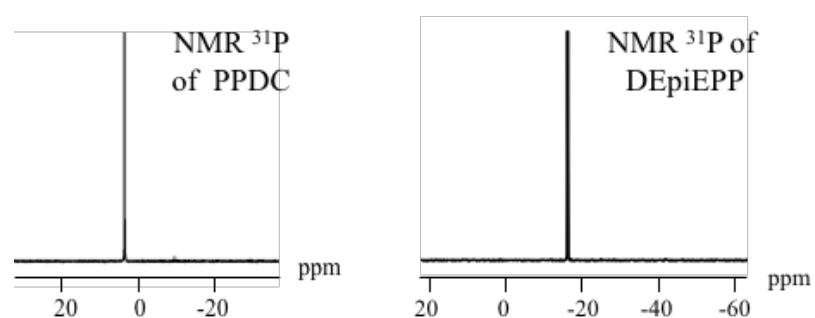
## RESULTS AND DISCUSSION

### *Synthesis of bio-based phosphorus-containing flame retardant*

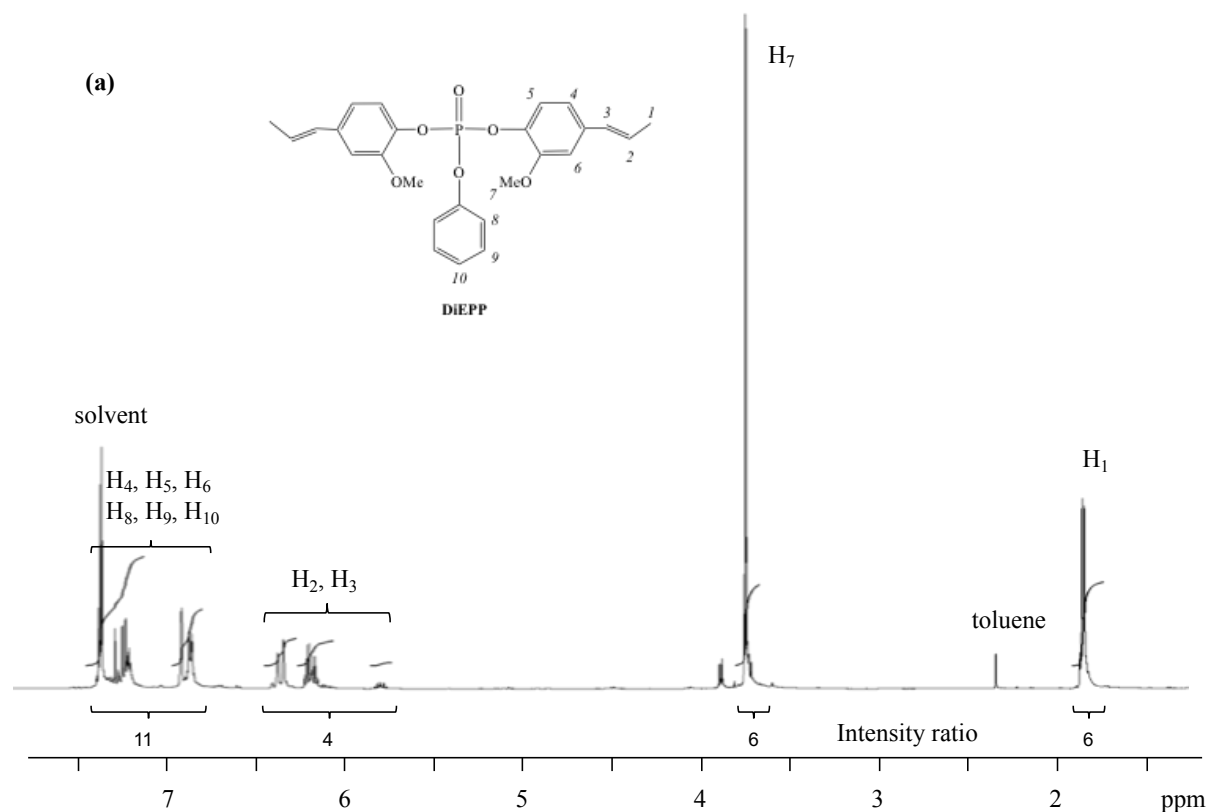
To the best of our knowledge, the synthesis of diepoxy-iso-eugenol phenylphosphate (DEpiEPP) is unprecedented. This monomer combines both the iso-eugenol (iEu) bio-based synthon, which can be catalytically extracted from the lignin,<sup>60</sup> and a phosphate group which can provide flame retardant properties.

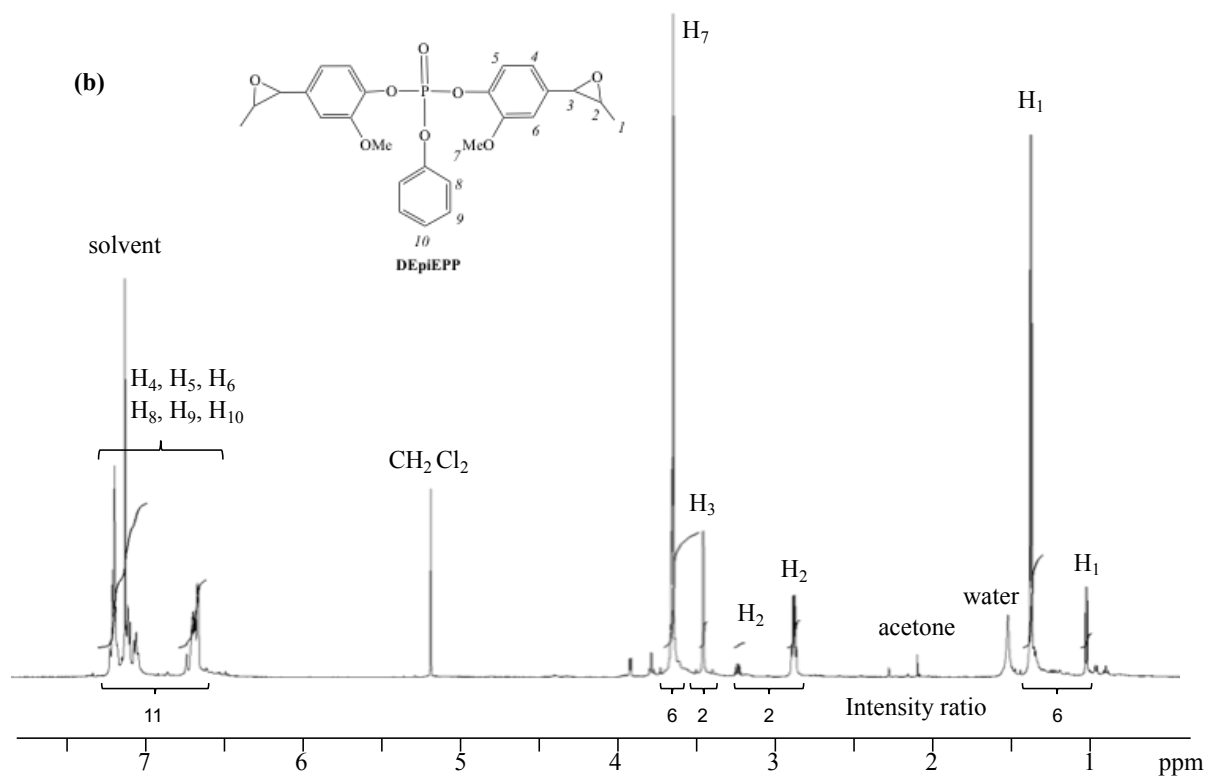
DEpiEPP is prepared in two reaction steps as shown in scheme 2b. First the condensation of iEu and PPDC in toluene using TEA as acid acceptor leads to DiEPP. Epoxidation<sup>61</sup> by Oxone® affords DEpiEPP in a 52 % yield. DiEPP and DEpiEPP structures are confirmed by FTIR-ATR, <sup>1</sup>H NMR, <sup>31</sup>P NMR, <sup>13</sup>C NMR (see Figures 1-3) and mass spectrometry.

DiEPP and DEpiEPP formation is evidenced by the shift of the  $^{31}\text{P}$  NMR signal from 4 ppm in PPDC to -17 ppm (Figure 1). Not surprisingly the  $^1\text{H}$  NMR spectrum of DiEPP (Figure 2a) combines the signals due to the iso-eugenol moiety and those of the phenoxy part. Assignments of the chemical shifts from  $^1\text{H}$  NMR analysis shows the disappearance of the relative protons to C=C linkage ( $\delta = 6.09\text{-}6.25$  ppm) and the appearance of the oxirane ring protons ( $\delta = 2.96\text{-}3.54$  ppm) in the DEpiEPP spectrum (Figure 2b).

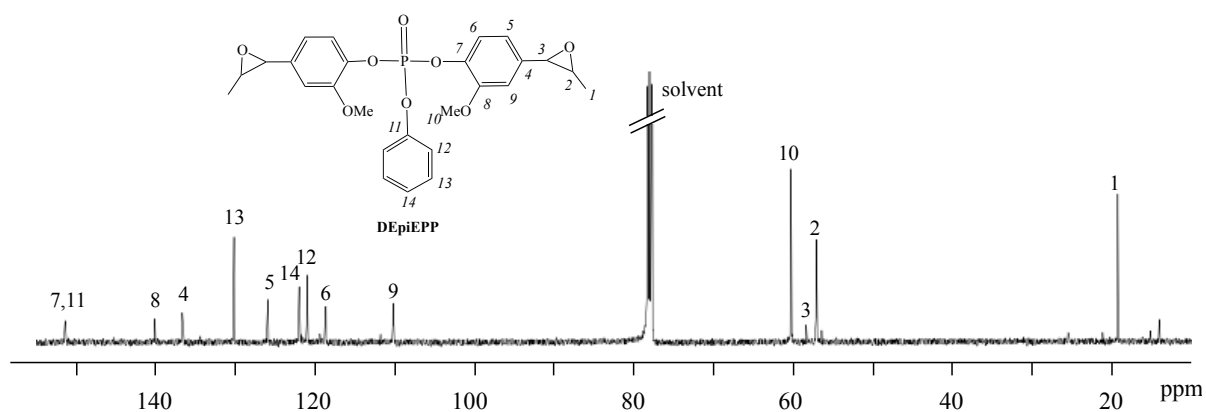


**Figure 1.**  $^{31}\text{P}$  NMR spectra of PPDC (left) and DEpiEPP (right)





**Figure 2.** <sup>1</sup>H NMR spectra of: (a) DiEPP and (b) DEpiEPP



**Figure 3.** <sup>13</sup>C NMR spectra of DEpiEPP.

The FTIR spectra of DiEPP and DEpiEPP are in accordance with the DiEPP and DEpiEPP proposed structures. The P-O-C linkage is evidenced by a peak at 1030 cm<sup>-1</sup> whereas the



disappearance of the C=C vinyl stretching band at 1655 cm<sup>-1</sup> and the appearance of the oxirane ring at around 904 cm<sup>-1</sup> confirm the DEpiEPP formation.

### ***Formulations studied***

With the aim of achieving 100% bio-based epoxy thermosets with flame retardancy properties, various formulations have been studied (Table 1).

Thermosets	P <sup>a</sup> (w%)	w% epoxy / P-FR / curing agent/DMID	Bio-based content (%)
DGEBA/ DEpiEPP / CA	0.0	60.4/0/38.8/0.9	36
	1.0	46.3/16.1/36.8/0.8	47
	2.0	32.3/32.1/34.8/0,8	57
	3.0	18.2/48.2/32.8/0.7	68
GEEpiE / DEpiEPP / CA	0.0	51.4/0/47.6/1	100
	1.0	39.4/16.1/43.5/1	96
	2.0	27.5/32.1/39.5/0.9	93
	3.0	15.5/48.2/35.5/0.8	89
DEpiEPP / CA	4.3	0/69.1/30.3/0.7	84
GEEpiE / DEpiEPP / PA	0.0	56.4/0/42.5/1.1	100
	1.0	44/16.1/38.9/1	96
GEEpiE / DEpiEPP / DPA	0.0	46.3/0/52.8/0.9	100
	1.0	34.8/16.1/48.3/0.9	97

<sup>a</sup>: Phosphorus content (w% P/thermoset)

**Table 1** Weight compositions data of the thermosets.

The reactive P-FR has been copolymerized with two types of epoxy resins, the petroleum-based DGEBA and the bio-based GEEpiE in the presence of two types of hardeners, the petroleum-

based IPDA and the bio-based anhydrides CA, PA and DPA. By this way epoxy materials ranging from petroleum-based 0% FR thermosets (DGEBA/IPDA) to totally bio-based FR thermosets (GEEpiE/DEpiEPP/CA) have been obtained.

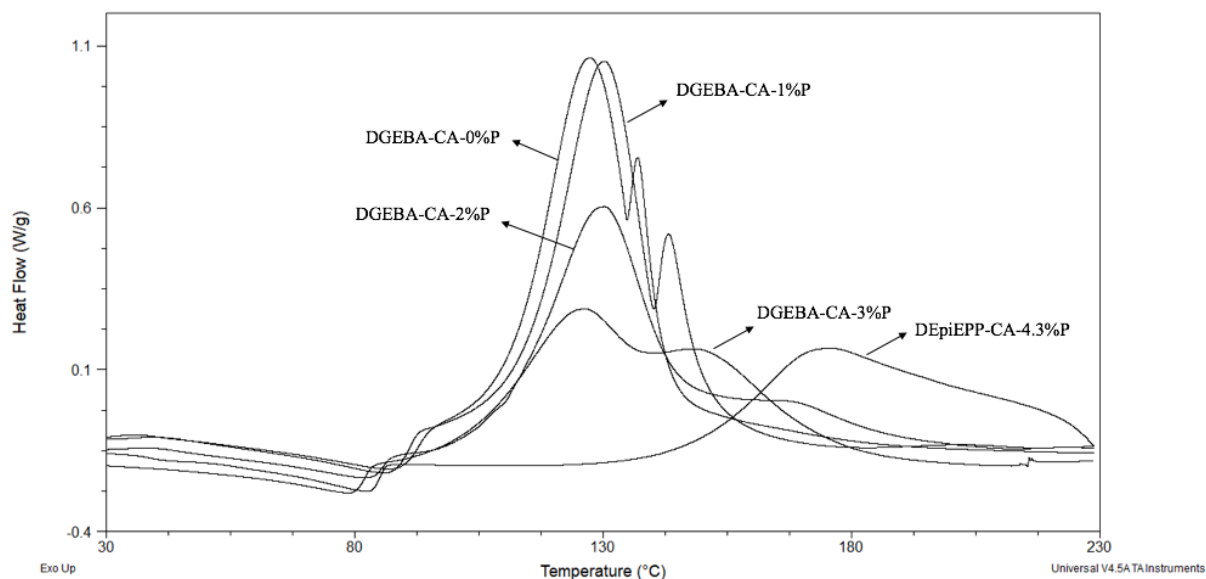
DGEBA was selected as it is the most used epoxy monomer in industry thanks to its aromatic structure that brings rigidity to the networks and therefore mechanical properties are excellent.<sup>62,63</sup> Anhydride curing agents (CA, PA, DPA) were chosen because they are bio-based and contain rigid ring leading to a reinforcement of the thermosets mechanical properties, whereas IPDA was selected as a model of polyamine hardener.<sup>13</sup>

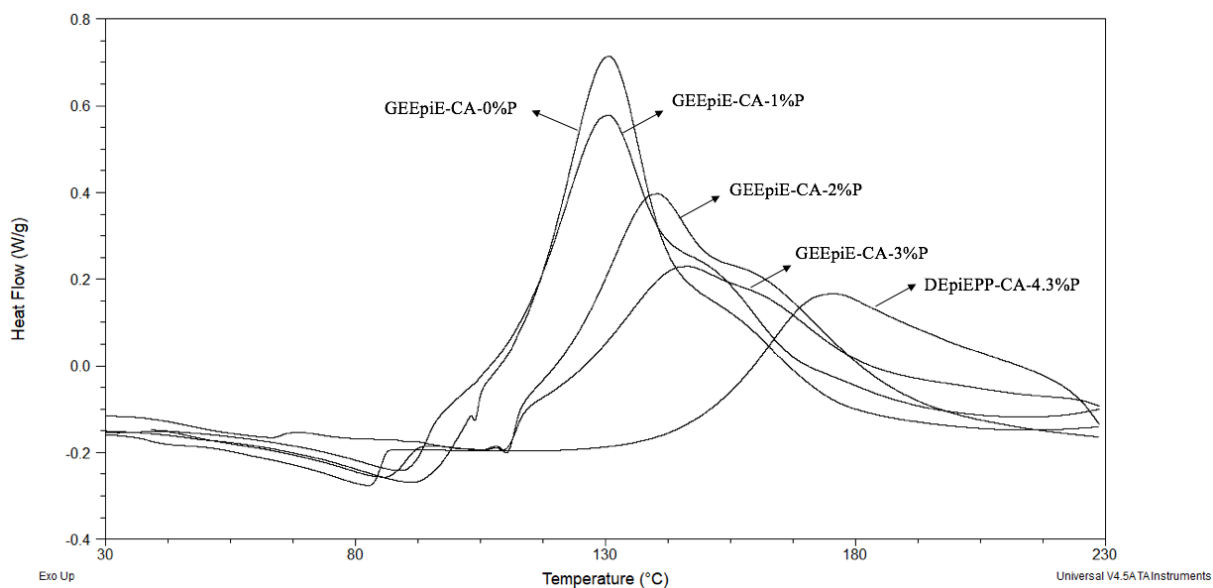
The diepoxy monomer/anhydride hardener and diepoxy monomer/polyamine molar ratio were equal to 1/1.2 and 2/1 respectively, as previously reported.<sup>6,64</sup>

Thermosets with varying rates of P-FR, from 0.0 to 4.3 w% phosphorus, have been produced (the 4.3 w% phosphorus sample corresponds to pure DEpiEPP).

### ***Curing step***

Curing of epoxy resins with CA hardener has been investigated by DSC. Samples were heated from 0 °C to 230 °C at a heating rate of 5 °C min<sup>-1</sup>. DSC thermograms are depicted in Figure 4.





**Figure 4.** DSC curves of dynamic curing reaction (above) DGEBA/CA (below) GEEpiE/CA series thermosets with varying DEpiEPP contents (0.0 to 4.3% P).

Thermosets	DSC				
	$P^a$ (w%)	Polymerization Start-end T (°C)	Peak T (°C)	$\Delta H$ (J/g)	$T_g^b$ (°C)
DGEBA/DEpiEPP / IPDA	0.0	/	/	/	155
DGEBA/ DEpiEPP / CA	0.0	90-160	127 and 137	383	155
	1.0	90-170	130 and 143	371	137
	2.0	95-195	130	289	105
	3.0	95-185	126 and 150	261	89
GEEpiE / DEpiEPP / CA	0.0	95-180	131	346	162
	1.0	95-195	130	349	136
	2.0	110-220	140	312	129
	3.0	110-230	145	230	97
DEpiEPP / CA	4.3	150-230 <sup>c</sup>	175	194	71
GEEpiE / DEpiEPP / PA	0.0				115

	1.0	*	*	*	84
GEEpiE / DEpiEPP / DPA	0.0				123
	1.0	*	*	*	113

<sup>a</sup>: Phosphorus content (w% P/thermoset)

<sup>b</sup>: Glass temperature of thermosets cured in the oven

<sup>c</sup>: Beginning of the thermoset degradation à 230°C.

**Table 2:** DSC data of the thermosets.

Thermograms exhibit an exothermic peak related to the cross-linking reaction. As highlighted in table 2, when increasing the P-FR content, the exothermic peak is broadened and shifted to higher temperatures and the curing enthalpies concomitantly decrease, whatever the epoxy resin used. These phenomena reveal a lower reactivity of oxirane groups of the P-FR, in accordance with observations reported by Wan *et al.*<sup>55</sup> Indeed, the polarizability of the epoxy group progressively decreases in the order DGEBA, GEEpiE and DEpiEPP. In DGEBA and GEEpiE, the epoxy reactivity of the glycidyl ether groups is enhanced due to the presence of adjacent oxygen (induction effect). In addition, steric hindrance could also explain this lower oxirane reactivity. To confirm the lower reactivity of oxirane groups in the P-FR, complementary curing tests have been carried out at higher temperature (180°C) for DEpiEPP/CA (4.3% P). The resulting thermoset exhibits a higher T<sub>g</sub> of 84°C (instead of 71°C when curing temperature is 150°C). However, as the T<sub>g</sub> reflects both the crosslinking degree of the thermoset and its chemical structure, the P-FR chemical structure must also be considered.

With the aim of developing fully bio-based composites containing vegetable fibers, attention must be paid to the onset and exothermic peak temperatures in order to avoid the polymerization during the step of fiber impregnation and to prevent their damage.<sup>6</sup> It is noteworthy that all formulations well fit these requirements while a homogeneous paste is obtained at about 90°C

and the cross-linking reaction occurs below 160°C. Both thermoset series are thus suitable for manufacturing bio-composite reinforced by vegetable fibers.

### ***Glass transition temperature of the cured epoxy resins (DSC)***

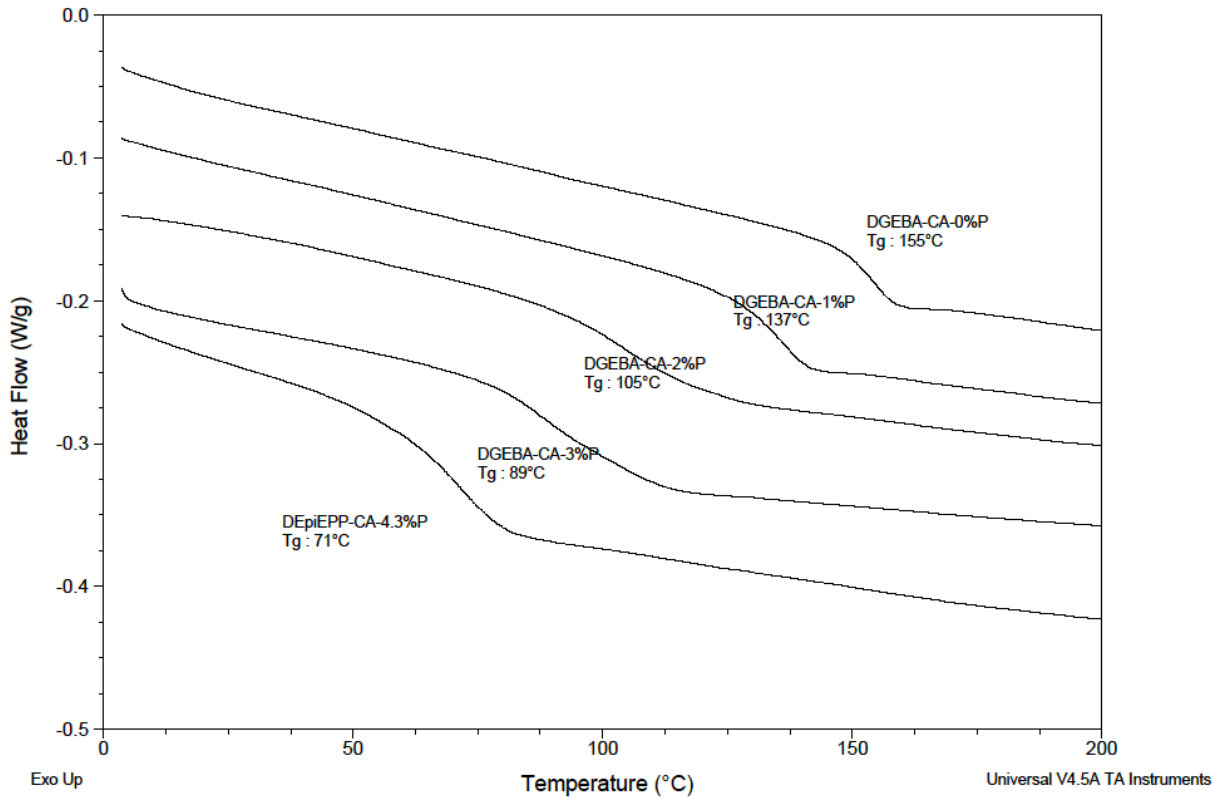
The DSC curves for the DGEBA/CA and GEEpiE/CA thermoset series are shown in Figure 5, and glass transition temperatures are reported in Table 2. It is noticeable that, for all the studied thermosets, no additional exothermal peak is observed in DSC after the curing schedule showing that the cross-linking reaction is ended. Moreover, as mentioned by Couture et al,<sup>20</sup> the initiator (DMID) could act as a thermoset plasticizer, decreasing thus the network Tg. Thus, higher Tg could be expected by precisely optimizing the required initiator quantity.

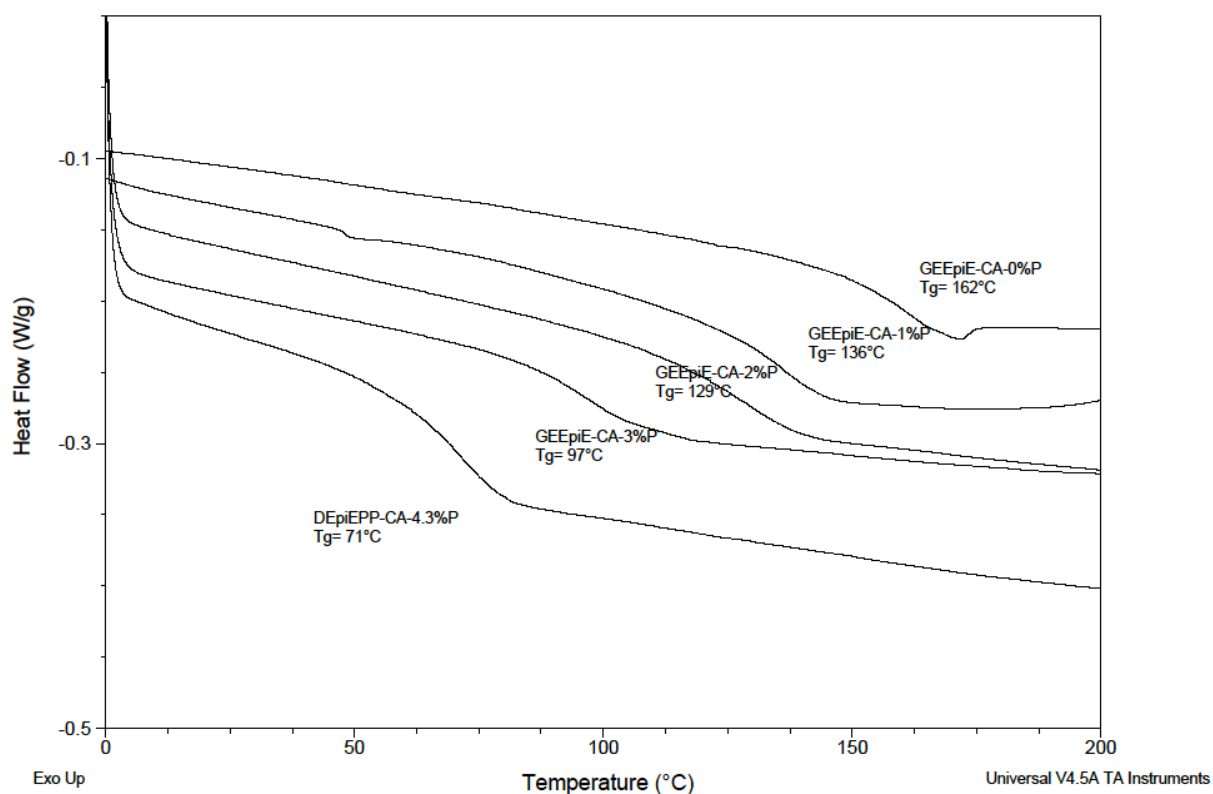
Not surprisingly, the Tg values progressively decrease as the P-FR content increases: the initial high Tg of about 160°C (free P-FR thermosets) drops to 71°C when the only P-FR (free DGEBA/GEEpiE) is cured. However, it is interesting to note that with a phosphorus weight content equal or lower than 2.0%, high Tg (> 100°C) are obtained with both GEEpiE-CA-DEpiEPP (95% bio-based content) and partially bio-based DGEBA-CA-DEpiEPP (57% bio-based content) epoxy thermosets. Interestingly the fully bio-based thermoset GEEpiE-CA-DEpiEPP, exhibits Tg of respectively 136°C and 129°C with the phosphorus weight contents of 1.0 or 2.0%.

As previously reported, phthalic and diphenic anhydrides were also found to be candidates as curing agents to obtain bio-based epoxy matrices for high performance composites. Thus, formulations of GEEpiE with both these anhydrides and with the green P-FR DEpiEPP were studied with a phosphorus weight content of 1.0% (see table 2). A Tg decrease of the same order (10 to 30°C) was again recorded when the epoxy resin was copolymerized with DEpiEPP, demonstrating that this effect remains the same whatever the hardener chemical structure. The Tg decrease has to be correlated with the lower reactivity highlighted by DSC recording during

the curing step, inducing then a lower cross-linking density. Similar effects were also reported and attributed to steric hindrance due to phosphate groups in the studies of Wang and co-authors<sup>13</sup> or Ma and coauthors.<sup>48</sup>

Thereby, both GEEpiE-CA-DEpiEPP (95% bio-based content) and partially bio-based DGEBA-CA-DEpiEPP (57% bio-based content) epoxy thermosets with the phosphorus weight contents of 2.0% seem to be good candidates for composite application in connection with their high Tg (respectively 129 and 105°C) compatible with further applications of the final material.





**Figure 5.** Determination of the glass transition by DSC (above) DGEBA/CA and (below) GEEpiE/CA thermoset series at different phosphorus content of DEpiEPP.

***Thermal degradation behaviour of the cured epoxy resins at microscale (TGA and PCFC)***

Anaerobic pyrolysis and complete combustion of flame retarded thermosets were characterized at microscale using TGA and PCFC (Table 3).

Formulation	%P	TGA			PCFC			$\Delta h$ (kJ/g)
		T <sub>5%</sub> (°C)	T <sub>max</sub> (°C)	Residue (w%)	pHRR (W/g)	T <sub>pHR</sub> R (°C)	THR (kJ/g)	
DGEBA / IPDA	0.0	355	382	9.0	648	370	28.3	31.1
DGEBA / CA	0.0	386	435	7.4	477	426	25.1	26.4
DGEBA / DEpiEPP / CA	1.0	322	381	15.5	631	388	22.4	25.7
DGEBA / DEpiEPP / CA	2.0	308	382	16.8	687	372	21.3	24.8

DGEBA / DEpiEPP / CA	3.0	296	370	17.9	501	361	20.8	25.1
GEEpiE / CA	0.0	364	430	8.7	336	409	22.9	24.6
GEEpiE / DEpiEPP / CA	1.0	309	380	14.7	367	375	21.1	23.4
GEEpiE / DEpiEPP / CA	2.0	301	366	15.0	402	368	20.7	23.8
GEEpiE / DEpiEPP / CA	3.0	293	357	18.3	363	361	20.2	24.6
DEpiEPP / CA	4.3	274	312	19.6	181	367	17.6	22.9

**Table 3** Data of the TGA and PCFC analyses of all the prepared thermosets.

Figure 6 and Figure 7 show the thermograms and HRR curves performed for DGEBA/CA and GEEpiE/CA thermoset series at different phosphorus contents. Both P-free thermosets exhibit close thermal stability and char ability ( $T_{5\%}$  is slightly lower for the latter). Note that their temperature at maximum mass loss rate is significantly higher than for DGEBA/IPDA thermoset.

For both thermoset series, it is noted that the thermal stability is lower than that of corresponding P-FR-free matrices and it depends clearly on the phosphorus content (Figure 8). Higher is the phosphorus content, lower are the onset temperature  $T_{5\%}$  and temperature at peak of mass loss rate  $T_{\max}$  in TGA.  $T_{pHRR}$  in PCFC follows the same tendency and is generally close to  $T_{\max}$ . As an example,  $T_{5\%}$  is reduced from 386 °C for the 0 w% P (DGEBA/CA) thermoset to 322 °C for the 1.0 w% P (DGEBA / DEpiEPP / CA) and to 296 °C for the 3 w% P (DGEBA / DEpiEPP / CA). Concerning the GEEpiE-based thermosets,  $T_{5\%}$  is reduced from 364 °C for the 0.0 w% P (GEEpiE/CA) thermoset to 309 °C for the 1.0%P(GEEpiE / DEpiEPP / CA) and to 293 °C for the 3.0 w% P (GEEpiE / DEpiEPP / CA) (and to 274 °C for 4.3 w% P). These lower degradation temperatures are partly due to the reduced crosslink density and mostly to the early stage of decomposition for phosphorus-containing groups.<sup>5,65</sup> Indeed, phosphorus flame retardants decompose into phosphoric acids and dehydrate oxygen-rich

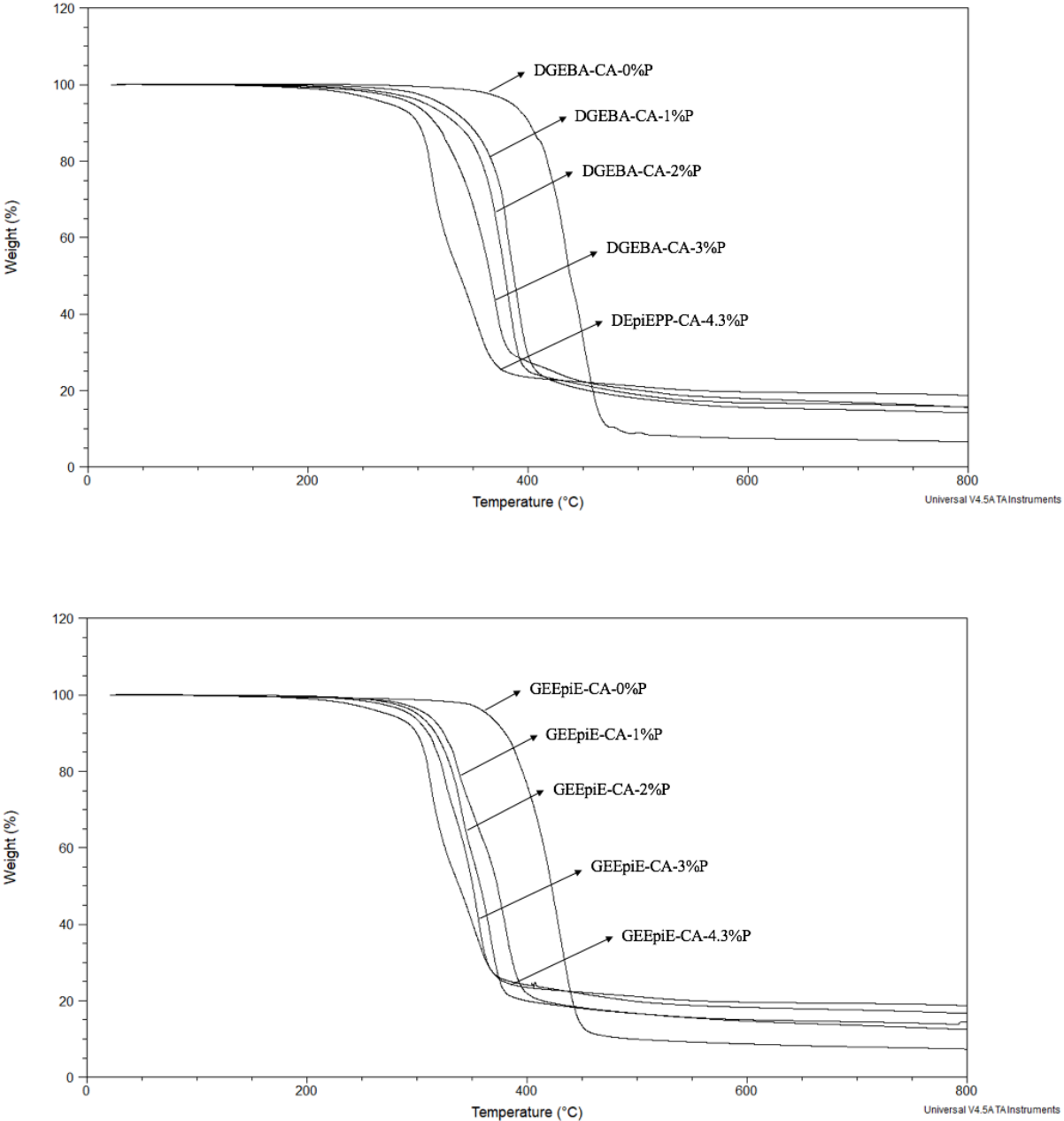


polymers promoting their charring. Note that the decrease is similar for both series (DGEBA and GEEpiE series).

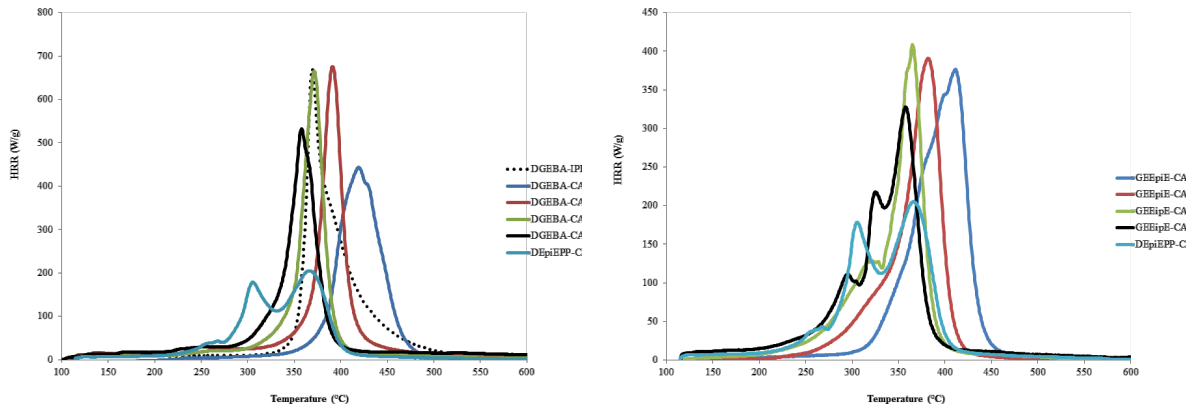
Peak of heat release rate is highly dependent on the network structure, i.e. on the chemical groups constituting the network. DGEBA/IPDA exhibits a higher pHRR ( $648 \text{ W g}^{-1}$ ) than DGEBA/CA ( $477 \text{ W g}^{-1}$ ) and GEEpiE/CA ( $336 \text{ W/g}$ ). Interestingly the incorporation of phosphorus does not lead to a reduction of pHRR. On the contrary, for low phosphorus content, the pHRR not only is reached at lower temperature but it also increases significantly and becomes narrower. This evidences that the phosphorus monomer (DEpiEPP) induces a thermal destabilization of the network. At higher content, the decomposition occurs in several well-identified peaks especially for GEEpiE/CA 3.0% P and DEpiEPP/CA 4.3% P. Therefore, the main pHRR decreases. There is no monotonous decrease in pHRR, as observed in other studies. THR of P-free thermosets is in the range  $22.9\text{-}28.3 \text{ kJ g}^{-1}$  depending on the network structure. Once again, GEEpiE/CA exhibits the lowest THR ( $22.9 \text{ kJ g}^{-1}$ ) while DGEBA/CA shows intermediate value ( $25.1 \text{ kJ g}^{-1}$ ). The increase of phosphorus content leads to a slight and continuous reduction in THR to  $20.8 \text{ kJ g}^{-1}$  and  $20.2 \text{ kJ g}^{-1}$  for 3.0 w% P (respectively DGEBA/CA and GEEpiE/CA). As explained below, this reduction is mainly due to an enhancement of char content rather than to a decrease in heat of combustion.

According to the char content, the DEpiEPP improves the charring of the matrix during thermal decomposition as expected. For DGEBA/CA and GEEpiE/CA thermoset series, the char content increases with the phosphorus content. It raises from 7.4 w% for the P-FR-free matrix to 15.5 w% with 1.0 w% P and to 17.9 w% with 3.0 w% P for DGEBA-based thermosets. Concerning the GEEpiE-based thermosets, the char content increases from 8.7 w% for the P-FR-free matrix to 14.7 w% with 1.0 w% P and to 18.3 w% with 3.0 w% P (and up to 19.6 w% with 4.3 w% P). These results reveal that DEpiEPP seems to be a good flame retardant since it promotes a significant charring of both epoxy thermosets.

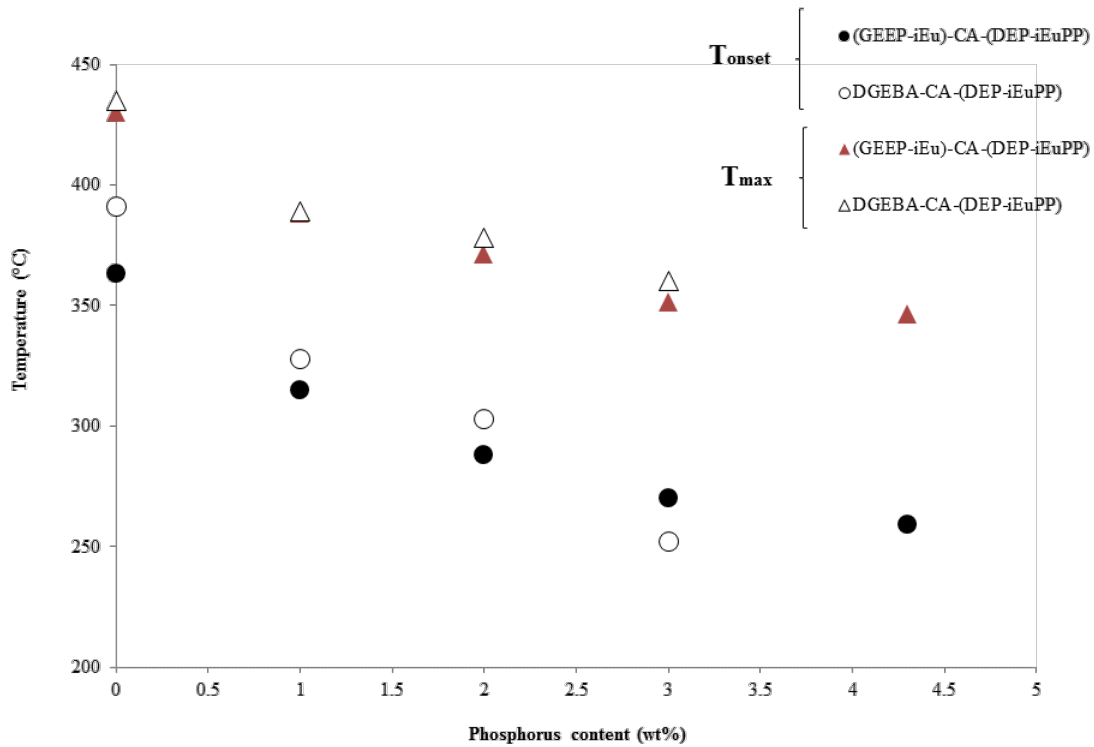
Heat of complete combustion was calculated from THR and residue content data. For both series, heat of complete combustion seems to decrease marginally. Indeed, char traps a large amount of carbon and reduces the energy released by pyrolysis gases. It has already been shown that the atomic composition of char is close to  $C_5H_2$  for a couple of polymers<sup>66</sup> corresponding to an energy released by char oxidation higher than  $30 \text{ kJ g}^{-1}$ . Nevertheless the heat stored in char containing phosphorus is generally lower.<sup>67</sup>



**Figure 6.** Thermograms of (above) DGEBA/CA and (below) GEEpiE/CA thermoset series at different phosphorus contents obtained by TGA at  $20 \text{ °C min}^{-1}$  under nitrogen atmosphere.



**Figure 7.** HRR curves of (left) DGEBA/IPDA and DGEBA/CA and (right) GEEpiE/CA thermoset series at different phosphorus contents obtained by PCFC in standard conditions.



**Figure 8.** Onset temperatures ( $T_{5\%}$ ) and temperatures at peak of mass loss rate  $T_{max}$  versus phosphorus content for DGEBA and GEEpiE series.

### Flame retardancy at bench-scale

Some samples were characterized using cone calorimeter tests at an external heat flux of 35 kWm<sup>-2</sup> (Figure 9 and table 4), which represent more realistic burning conditions than PCFC. All phosphorus-free resins exhibit a continuous increase of HRR after ignition up to a high peak of heat release rate (1800-2100 kW m<sup>-2</sup>) followed by a fast decrease due to fuel depletion. Residue content is limited and close to the values measured in TGA. The only difference concerns the time-to-ignition. DGEBA-CA appears to be more difficult to ignite. This thermoset was also the most thermally stable according to TGA.

The incorporation of phosphorus leads to a significant decrease of time-to-ignition. Such decrease is usual and can be assigned to the reduced stability of the resins as observed in TGA and PCFC. The ignition is followed by an increase of HRR up to a plateau during several dozens of seconds. Then HRR increases up to a pHRR which is twice as weak as the pHRR of the phosphorus-free counterpart. Char is significantly improved and in the same range as measured in TGA.

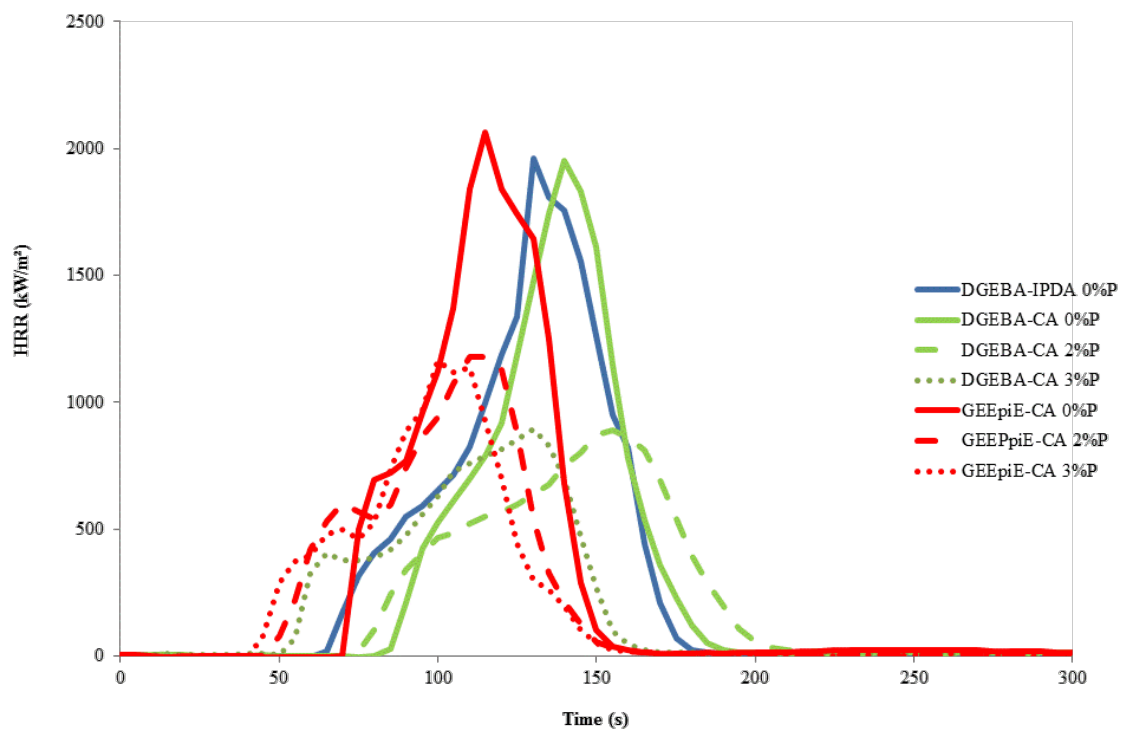
While char contents are similar in cone calorimeter and in TGA, it is possible to compare the effective heat of combustion (EHC) and the heat of complete combustion ( $\Delta h$ ). The ratio between both values corresponds to the combustion efficiency  $X$ . Phosphorus-free thermosets exhibit quite high combustion efficiency (0.79-0.95 depending on the thermoset). The incorporation of phosphorus leads to a significant decrease in combustion efficiency, which may be ascribed to a flame inhibition effect of phosphorus gases released in the flame.

Finally, total heat release is also significantly lower because a part of fuels is trapped into char and does not participate to combustion (charring promotion) but also because the combustion efficiency decreases.

Note that GEEpiE/CA with 2.0 and 3.0 w% P exhibit very close flammability behavior. No further improvement is obtained when phosphorus content increases from 2.0 to 3.0 w%. For

DGEBA-based series, there is no more improvement of pHRR when phosphorus content increases from 2.0 to 3.0 w%. But TTI decreases and char content increases significantly.

A proper comparison of flame retardancy with results from literature is quite difficult because of the major influence of the curing agent. For example, in a recent paper, several epoxy monomers based on eugenol and containing phosphate or phosphonate groups were cured with meta-xylylenediamine<sup>54</sup>. Performances at microscale were outstanding with very high char content and low pHRR and THR. Phosphorus contents were relatively higher (up to 5.8 wt%) than in the present study but in one case it reaches 4.5 wt%, which is similar to the value for DEpiEPP/CA. Nevertheless, the decrease in pHRR in cone calorimeter was similar: -55 to -60% versus -60% for DGEBA/DEpiEPP /CA with 2 wt% of phosphorus. Menard et al. also found a decrease of 55% in pHRR in cone calorimeter for a network based on DGEBA, IPDA and an epoxy monomer containing phosphorus and sulfur called P2EP1SP (Diglycicyl thiophosphonate phloroglucinol - phosphorus content in epoxy resin was 3 wt%)<sup>64</sup>.

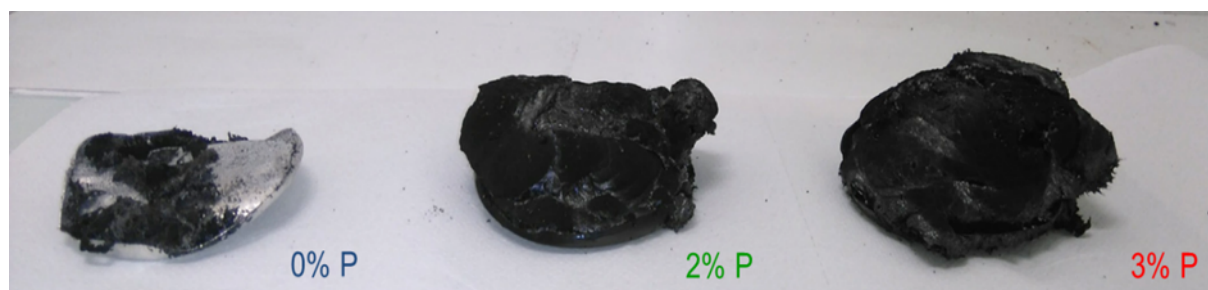


**Figure 9.** HRR curves obtained in cone calorimeter tests for several thermosets containing different phosphorus contents.

Formulation	% P	TTI s	pHRR kW.m <sup>-2</sup>	THR MJ.m <sup>-2</sup>	THR kJ.g <sup>-1</sup>	EHC kJ.g <sup>-1</sup>	X	Residue w%
DGEBA/IPDA	0.0	69	2145	99	22.7	24.5	0.79	6
DGEBA/CA	0.0	87	1848	90	20.4	22.3	0.84	9
DGEBA/DEpiEPP /CA	2.0	78	849	66	15.9	18	0.73	11.5
DGEBA/DEpiEPP /CA	3.0	53	890	58	13.8	17	0.68	19
GEEpiE/CA	0.0	71	2065	92	21.6	23.3	0.95	7.1
GEEpiE/DEpiEPP /CA	2.0	48	1302	67	15.7	19.3	0.81	18.3
GEEpiE/DEpiEPP /CA	3.0	43	1159	61	14.4	18.1	0.74	20.7

**Table 4** Data of the cone calorimeter analyses of all the prepared thermosets.

Pictures of some residues after cone calorimeter tests are shown in Figure 10. It is obvious than the incorporation of phosphorus groups leads to the formation of an expanded char. Such char may be able to prevent the heat and gas transfer between the flame and underlying material. Nevertheless, the shape of the HRR curves does not correspond to a thick charring behaviour and cannot prove the efficiency of the char as insulating layer.<sup>68</sup>



**Figure 10.** Pictures of the cone residues: GEEpiE/CA (0.0% P-2.0% P and 3.0% P) thermosets obtained by cone calorimeter.

The residual phosphorus content for the residues of the GEEpiE/CA series thermosets containing DEpiEPP was determined by ICP. Results are summarized in table 5. The phosphorus content is much higher in the char than in the initial corresponding resin, evidencing that a significant part of the phosphorus remains in the char and contributes to the formation of a thermally stable carbonaceous layer. For 3.0 w% of phosphorus initially in the resin, the phosphorus content in the char reaches 8.5 w%. This value is close to that found by Menard et al. in the char obtained from DGEBA-IPDA-P2EP1SP after thermogravimetric analysis under nitrogen flow.<sup>60</sup> Nevertheless, in the present work, only around 60 % of phosphorus remains in the char versus 82 % in the Menard's work. Note that the release of a significant amount of phosphorus in gas phase (i.e. around 40%) is compatible with the flame inhibition effect suggested from combustion efficiency data.

Notation	Initial phosphorus content in thermoset (w%)	Average phosphorus content in char (w%)	Average residual phosphorus content (%)
GEEpiE/CA	0.0	0	0
GEEpiE/DEpiEPP /CA	2.0	6.78	62
GEEpiE/DEpiEPP /CA	3.0	8.50	59

**Table 5** Determination of the residual phosphorus content in the cone calorimeter residue of the GEEpiE/CA series thermosets containing DEpiEPP.

### ***Mechanical properties of the cured epoxy resins.***

The GEEpiE-CA-DEpiEPP and DGEBA-CA-DEpiEPP epoxy thermosets with a phosphorus weight content equal to 2% were also mechanically characterized using nanoindentation and 3-point bending test. The nanoindentation properties obtained for these fully and partially bio-based epoxy thermosets (Table 6) are in the high range of the target bracket and are fully

suitable for structural applications. Noteworthy, they are in the same range as those previously reported for eugenol-based epoxy thermosets<sup>7 55</sup>. Interestingly, it can be observed that the highest properties are obtained for the fully bio-based epoxy with a relaxed modulus of 4.54 GPa, an instantaneous modulus of 5.1 GPa and a hardness of 278 MPa. The time-dependency is also slightly higher with a  $m$  value of approximately 0.032.

Samples (2%P)	$E_R$ (GPa)	$E_I$ (GPa)	H (MPa)	$m$ ( $\emptyset$ )
DGEBA/DEpiEPP/CA	4.45±0.03	4.8±0.03	264±14	0.020
GEEpiE / DEpiEPP / CA	4.54±0.13	5.1±0.05	278±3	0.032

**Table 6.** Properties measured with nanoindentation technique: average and standard deviation values of relaxed modulus ( $E_R$ ), instantaneous modulus ( $E_I$ ), hardness (H) and strain rate sensitivity ( $m$ ).

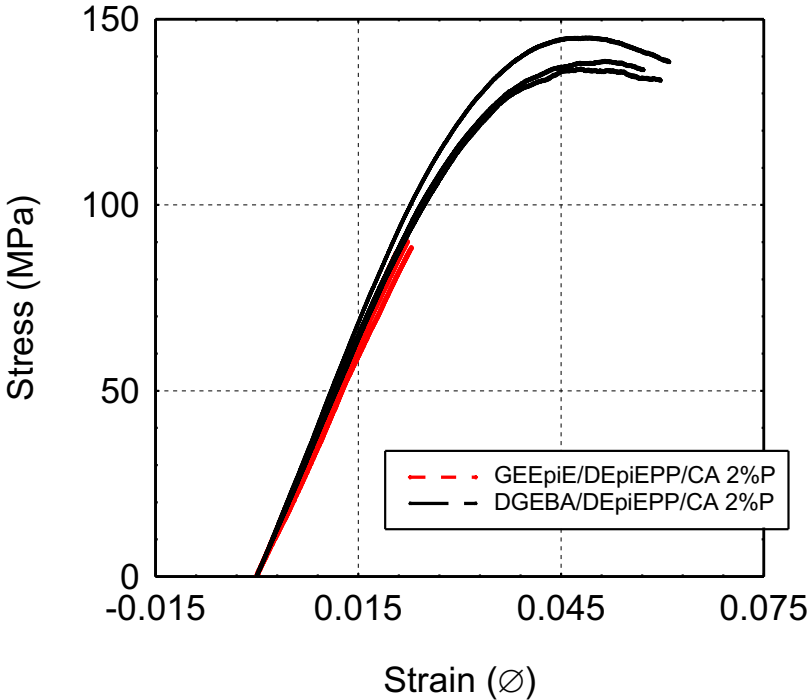
The 3-point bending test results are presented in table 7 and figure 11. For purposes of comparison, the mechanical characteristics of the commercial epoxy thermoset SR GreenPoxy 56®, according to Sicomin technical data sheet, are also given in table 7. The values obtained for the DGEBA-CA-DEpiEPP epoxy are in line with the values generally reported in literature<sup>12</sup> and technical datasheet for the DGEBA-based systems<sup>69</sup> and slightly higher than those of SR GreenPoxy 56®. The elastic modulus, maximum stress and strain at failure are equal to 4.3 GPa, 140 MPa and 5%, respectively. Concerning the rigidity, it can be observed that the same order of magnitude as the one obtained with nanoindentation is measured. The mean value for the GEEpiE-CA-DEpiEPP epoxy specimens is slightly lower (6%) than for the DGEBA-CA-DEpiEPP epoxy ones. In any case, such an elastic modulus of approximately 4 GPa is fully suitable for structural applications. A significant difference can be observed between the two epoxy systems for the maximum stress and strain at failure. The values measured for the GEEpiE-CA-DEpiEPP epoxy are 35% and 45% lower than for the DGEBA-CA-DEpiEPP epoxy polymer. Indeed, the stress-strain curves (Fig. 11) show two different failure behavior.



The DGEBA-CA-DEpiEPP epoxy polymer presents a ductile behavior with a yield point at approximately 85 MPa and 2% of strain and then a strongly non-linear behavior attributed to plasticity. On the contrary, the GEEpiE-CA-DEpiEPP epoxy polymer exhibits an elasto-brittle behavior with a sudden failure after exceeding the yield point.

	Modulus of elasticity (GPa)	Stress at failure (MPa)	Strain at failure (%)
DGEBA/DEpiEPP/CA	$4.3 \pm 0.2$	$140 \pm 4$	$5.0 \pm 0.1$
GEEpiE/DEpiEPP/CA	$4.1 \pm 0.2$	$90 \pm 1$	$2.2 \pm 0.1$
SR GreenPoxy 56®	$3.0 \pm 0.2$	$96 \pm 5$	$8 \pm 2$

**Table 7.** 3-point bending properties: average and standard deviation values of the elastic modulus, maximum stress and strain at failure.



**Figure 11.** Bending stress/strain curves obtained for the GEEpiE-CA-DEpiEPP and DGEBA-CA-DEpiEPP epoxy thermosets with a phosphorus weight content equal to 2%.

## CONCLUSION

A new bio-based epoxy phosphate FR derived from iso-eugenol was successfully synthesized and well characterized ( $^1\text{H}$ ,  $^{13}\text{C}$ ,  $^{31}\text{P}$  NMR, FTIR, MS). This diepoxy-iso-eugenol phenylphosphate (DEpiEPP) was then copolymerized either with conventional diglycidylether of bisphenol A resin (DGEBA) or with a totally bio-based glycidylether epoxy iso-eugenol (GEEpiE) resin in the presence of the bio-based camphoric anhydride (CA) used as hardener. The cured resins (2h at  $150^\circ\text{C}$ ) containing various contents of DEpiEPP were then mechanically and thermally characterized.

As the level of the DEpiEPP-FR monomer increases (0.0 to 4.3% wt P), the Tg of the cured resin decreases from  $155\text{-}162^\circ\text{C}$  to  $71^\circ\text{C}$  when the final material only contains the DEpiEPP-FR (4.3% wt P). TGA, PCFC and cone calorimeter results reveal that DEpiEPP is a good flame retardant since it promotes a significant charring of both epoxy thermosets (DGEBA-DEpiEPP-FR and GEEpiE-DEpiEPP-FR). Both formulations obtained with a ratio of 2%wt of P provide very promising green epoxy thermosets for developing high performance green composites (respectively 57% and 95% bio-based content). Indeed, the curing temperature of these epoxy thermosets remains compatible with the use of vegetable fibres and these epoxy matrices possess high Tg (respectively  $129$  and  $105^\circ\text{C}$ ), instantaneous elastic modulus of respectively 5.1 and 4.8 GPa, as well as high bending properties and demonstrate good flame-retardant properties due to char promotion.

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## For Table of Contents Use Only

**Scheme 1.** Molecular representations of reagents used. Epoxy monomers: diglycidylether of bisphenol A (DGEBA), glycidylether of phenoxy iso-eugenol (GEEpiE). Flame retardants: diepoxyiso-eugenolphenylphosphate (DEpiEPP). Curing agents: isophorone diamine (IPDA), camphoric anhydride (CA), phthalic anhydride (PA) and diphenic anhydride (DPA). Catalyst: 1,2-dimethylimidazole (DMID).

**Scheme 2.** Description of the diepoxy monomers synthesis.

**Figure 1.**  $^{31}\text{P}$  NMR spectra of PPDC (left) and DEpiEPP (right)

**Figure 2.**  $^1\text{H}$  NMR spectra of: (a) DiEPP and (b) DEpiEPP

**Figure 3.**  $^{13}\text{C}$  NMR spectra of DEpiEPP.

**Figure 4.** DSC curves of dynamic curing reaction (above) DGEBA/CA (below) GEEpiE/CA series thermosets with varying DEpiEPP contents (0.0 to 4.3% P).

**Figure 5.** Determination of the glass transition by DSC (above) DGEBA/CA and (below) GEEpiE/CA thermoset series at different phosphorus content of DEpiEPP.

**Figure 6.** Thermograms of (above) DGEBA/CA and (below) GEEpiE/CA thermoset series at different phosphorus contents obtained by TGA at  $20\text{ }^\circ\text{C min}^{-1}$  under nitrogen atmosphere.

**Figure 7.** HRR curves of (left) DGEBA/IPDA and DGEBA/CA and (right) GEEpiE/CA thermoset series at different phosphorus contents obtained by PCFC in standard conditions.

**Figure 8.** Onset temperatures ( $T_{5\%}$ ) and temperatures at peak of mass loss rate  $T_{\text{max}}$  versus phosphorus content for DGEBA and GEEpiE series.

**Figure 9.** HRR curves obtained in cone calorimeter tests for several thermosets containing different phosphorus contents.

**Figure 10.** Pictures of the cone residues: GEEpiE/CA (0.0% P-2.0% P and 3.0% P) thermosets obtained by cone calorimeter.

**Figure 11.** Bending stress/strain curves obtained for the GEEpiE-CA-DEpiEPP and DGEBA-CA-DEpiEPP epoxy thermosets with a phosphorus weight content equal to 2%.

**Table 1** Weight compositions data of the thermosets.

**Table 2** DSC data of the thermosets.

**Table 3** Data of the TGA and PCFC analyses of all the prepared thermosets.

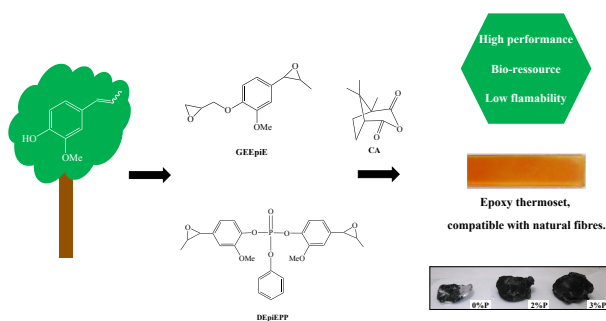
**Table 4** Data of the cone calorimeter analyses of all the prepared thermosets.

**Table 5** Determination of the residual phosphorus content in the cone calorimeter residue of the GEEpiE/CA series thermosets containing DEpiEPP.

**Table 6** Properties measured with nanoindentation technique: average and standard deviation values of relaxed modulus ( $E_R$ ), instantaneous modulus ( $E_I$ ), hardness (H) and strain rate sensitivity (m).

**Table 7** 3-point bending properties: average and standard deviation values of the elastic modulus, maximum stress and strain at failure.

### Graphical abstract:



**Synopsis (~20 words)**

High performance and fire-resistant green epoxy thermosets are prepared using iso-eugenol, a bio-ressource easily available from lignin.