Valorization of Industrial Lignin as Biobased Carbon Source in Fire Retardant System for Polyamide 11 Blends

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Abstract: In this study, two different types of industrial lignin (i.e., lignosulphonate lignin (LL) and kraft lignin (DL)) were exploited as charring agents with phosphorus-based flame retardants for polyamide 11 (PA11). The effect of lignins on the thermal stability and fire behavior of PA11 combined with phosphinate additives (namely, aluminum phosphinate (AlP) and zinc phosphinate (ZnP)) has been studied by thermogravimetric analysis (TGA), UL 94 vertical flame spread, and cone calorimetry tests. Various blends of flame retarded PA11 were prepared by melt process using a twin-screw extruder. Thermogravimetric analyses showed that the LL containing ternary blends are able to provide higher thermal stability, as well as a developed char residue. The decomposition of the phosphinates led to the formation of phosphate compounds in the condensed phase, which promotes the formation of a stable char. Flammability tests showed that LL/ZnP ternary blends were able to achieve self-extinction and V-1 classification; the other formulations showed a strong melt dripping and higher burning. In addition to this, cone calorimetry results showed that the most enhanced behavior was found when 10 wt % of LL and AlP were combined, which strongly reduced PHRR (−74%) and THR (−22%), due to the interaction between LL and AlP, which not only promotes char formation but also confers the stability to char in the condensed phase.

Keywords: industrial lignin; Polyamide 11; phosphinate; thermal decomposition; fire retardancy

1. Introduction

Over the past two decades, the sustainable development of flame retardant material has become a major topic of concern in industry and academia, due to the toxicity associated with flame retardants and increasing environmental and health safety regulations. This issue has driven the development of flame retardant systems made from renewable resources as possible alternatives to the non-halogenated ones [1]. Nowadays, Polyamide 11 (PA11), a renewable natural high-performance polymer derived from castor seed oil, has gained much attention because of its high mechanical strength and chemical resistance properties [2]. PA11 has been exploited in automotive, aerospace, sports applications, and textile industries [3–5]. However, low flame-retardant properties and extended dripping of PA11 limits its potential applications in high-performance textiles. In order to confer flame retardancy to PA11, intumescent systems based on bioresources are possible substitutes to halogen-free flame retardants. It is believed that the use of materials based on renewable resources can reduce the
carbon footprint in our environment. Lignin consists of non-toxic polyaromatic polyols, one of the most abundant biopolymers. The majority of industrial lignin is produced annually by the pulp and papermaking industries and the lignocellulosic ethanol industries; approximately 70 million tonnes of lignins are produced per year worldwide [6,7]. However, around 2% is isolated and comprehensively utilized, and the rest is primarily incinerated for energy recovery. In recent years, the valorization of lignin compounds has attracted growing interest for its potential applications in polymers, due to its competitive price, abundant availability, reactive functional groups, high aromatic carbon content, and tailored capability for structural modification [7–10].

In particular, lignin has been employed to improve the thermal stability of thermoplastic polymers [11–13], exploiting the thermal decomposition of lignin, which takes place over a broad temperature range, since various aromatic functional groups have different thermal stability. More recently, lignin has shown great potential as a biobased carbon source in an intumescent system with traditional flame retardants [14,15] because of its high char yield (about 50–60 wt %) upon thermal decomposition in inert atmosphere [16]. The char-formation can reduce the heat release rate of the polymeric material during the combustion process. Furthermore, char forming ability of lignin can be exploited for the design of environmentally friendly intumescent systems with phosphorus-based non-halogenated flame retardants. In this context, among various phosphorus flame retardants, metal phosphinates are considered very efficient and have several advantages, being mostly non-hygrosopic, non-toxic, thermally stable (>350 °C), and resistant to hydrolysis. In particular, AlP and ZnP are the most widely used and are found to be very effective flame retardants, especially for polyesters and polyamides [17]. However, they show satisfactory flame-retardant properties only on rather a high loading of about 30 wt % [18]. Phosphinate loading was conceivably reduced in the presence of synergistic or catalytic compounds. Furthermore, it was found that some nitrogen-containing compounds such as melamine polyphosphate (MPP) and melamine cyanurate (MC) show synergism when combined with phosphinate [19,20]. The efficiency was further improved by the addition of zinc borate (ZnB) to AlP and MPP system [21]. The presence of ZnB led to the formation of stable residue and a dominant barrier effect. In another study, Didane et al. exploited the synergistic effects of Polyhedral Oligomeric Silsesquioxane (POSS) with ZnP in developing flame retardant PET fibres [22]. The addition of synergistic agents enhances the ZnP efficiency, by reducing its required loading in the polymer matrix.

In order to promote the utilization of biobased lignin as flame retardant additive, our previous study has elucidated the use of lignin (pure grade) as a carbon source with ZnP [23]. Lignin used in this work was low sulfonate containing (4% sulphur) alkali kraft lignin. Different formulations of pure lignin and ZnP were prepared and their fire retardancy effect and thermal decomposition behaviour were investigated. It was found that the blends containing 10 wt % of both additives showed most enhanced flame-retardant properties, due to the formation of a stable char layer with barrier features. Besides, in the present work, we have focused on the use of chemically different industrial lignin (i.e., kraft lignin and lignosulfonate lignin) combined with two different phosphinates, i.e., aluminum phosphinate (AlP) and zinc phosphinate (ZnP). Lignins exploited in this work were chemically different and less pure than the same used in the previous study. The purpose of this study is to compare the flame-retardant effectiveness of industrial lignin with phosphinate in PA11. To this aim, the different formulations based on phosphinates and lignins were prepared by melt extrusion and thoroughly investigated. The morphology of prepared blends examined by Scanning Electron Microscopy (SEM) analyses. Their effects on the thermal stability and char-forming ability of blends were thoroughly investigated by thermogravimetric (TG) experiments. Fire behaviour of prepared blends was assessed by the vertical flame spread and cone calorimetry tests. Furthermore, the morphology of char residues after cone calorimetry tests was also investigated by SEM.
2. Experimental

2.1. Materials and Processing

A biobased Polyamide 11 (PA11), Rilsan® BMNO-TLD; $M_n = 17,000$ g/mol, melt flow index (MFI) = 14–20 g/10 min at 235 °C, supplied from Arkema (Colombes, France), was chosen as the polymer matrix. Flame retardants zinc phosphate (ZnP), Exolit 950 and aluminum phosphinate (AlP), Exolit 1230, were supplied by Clariant, Muttenz, Switzerland. Two different types of industrial lignins were used as a carbon source, varying in chemical nature: the first one is a lignosulphonate lignin (coded as LL), provided from Domsjö Fabriker AB (Örnsköldsvik, Sweden) and the second one is an alkali kraft lignin (coded as DL) obtained from UPM Biochemicals, Helsinki, Finland (European distributor of Domtar BioChoice® lignin). LL contains mainly Na-lignosulfonate (about 70%) and small amounts of Mg and Ca lignosulfonates, and some impurities such as ash and carbohydrates (about 20%). DL comprises mostly alkali kraft with 90% purity level. All the materials were dried at 80 °C for 12 h before use.

2.2. Blends Preparation

The different blend formulations based on PA11, lignins (LL or DL) and phosphinates (ZnP or AlP) were prepared by melt extrusion and labelled as PA$_X$-L$_Y$-P$_Z$, where $X$ represents the amount of PA11, L the type of lignin used, Y the wt % lignin loading, P the type of phosphinate, and Z the wt % phosphinate content. A total of 20 wt % loading was used in blend formulation; phosphinate, being the major component, was varied from 10 to 15 wt % and lignin from 5 to 10 wt % (Table 1). Blends were prepared using a co-rotating intermeshing twin-screw extruder (Thermo Haake, screw diameter = 16 mm, L/D = 25); the temperatures of the five heating zones ranged from 170 to 220 °C, and the rotation speed was set at 100 rpm. In all cases, the extruded rods were pelleted for thermal analysis. 3 mm thick plates were manufactured by compression molding using a Collins Teach Line 200 hydraulic press, operating at 60 bars and 220 °C for 3 min. Specimens in accordance with cone calorimeter ($100 \times 100$ mm$^2$) and UL-94 flammability ($125 \times 12.5$ mm$^2$) tests were prepared and used.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Polyamide 11 (wt %)</th>
<th>Lignin (L in wt %)</th>
<th>Phosphinate (P in wt %)</th>
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<tr>
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</tr>
<tr>
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<td>80</td>
<td>20</td>
<td>-</td>
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<td>PA$<em>{80}$-L$</em>{7}$-P$_{13}$</td>
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<td>80</td>
<td>10</td>
<td>10</td>
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</table>

Samples name are coded as PA$_X$-L$_Y$-P$_Z$, where L is the kind of lignin, i.e., LL (lignosulphonate lignin from Domsjö Fabriker AB) or DL (kraft lignin from UPM Biochemicals), P is the type of phosphinate, i.e., ZnP (zinc phosphinate, OP950) or AlP (aluminium phosphinate, OP1230)

2.3. Morphology

Morphology and distribution of the fillers in polymer matrix were assessed by SEM; analyses were carried out using a LEO-1450VP apparatus (Carl Zeiss, Oberkochen, Germany), equipped with a X-ray probe (INCA Energy Oxford Instruments, Cu Ka X-ray source, k = 1.540 562 Å, Abingdon-on-Thames, UK) under the voltage of 20 kV. Energy dispersive X-rays (EDX) elemental mapping was also used for assessing the dispersion of phosphinate and lignin in PA11 matrix. Pellets of different blends were fractured in liquid nitrogen and then coated with a thin conductive gold layer.

2.4. Thermal Decomposition

Thermogravimetric (TG) analyses were carried out with a TA instruments thermal analyzer Q500 (New Castle, DE, USA), either under air or nitrogen atmosphere at a purge rate of 60 mL/min,
with a heating rate of 10 °C/min from 50 to 800 °C, using alumina pans and sample weight of 10 ± 0.2 mg. TG curves were recorded from experiments and dTG (derivative) curves were obtained from TA universal data analysis software for all the samples. The decomposition parameters, such as the temperature at 5% weight loss (T5%), and residue at 700 °C, were obtained from TG curve. Furthermore, the maximum mass loss rate (MMLR) and the corresponding temperature (Tmax) were obtained from dTG curves. During TG analyses in air, the main decomposition step (Tmax1) and the second thermo-oxidative degradation step (Tmax2) were evaluated.

The weight difference curves were plotted in order to determine the potential increase or decrease in the thermal stability of the blends. The weight difference curves were computed for the loaded samples, and correspond to the weight difference between the experimental and theoretical TG curves:

\[
\Delta(M(T)) = M_{\text{exp}}(T) - M_{\text{theo}}(T)
\]

Where, \(\Delta(M(T))\) is a residual weight difference, \(M_{\text{exp}}(T)\) is the experimental residual weight of blends (variation with temperature \(T\)), \(M_{\text{theo}}(T)\) is the theoretical residual weight of the same composition computed by a linear combination between the experimental weights of PA11 and additives. The curves show the subsequent interaction between additives and polyamide matrix by the observation of an increase or decrease of the thermal stability.

2.5. Fire Behavior

The flammability was evaluated on sample sheets (125 × 12.5 × 3 mm³) by vertical flame spread tests according to IEC 60695-11-10 [24], also known as UL 94 burning flame test, and used for plastic materials. This test is aimed at assessing the material capability to extinguish a flame. Materials were classified on the basis of burning rate, time to flame extinction, and dripping during burning.

Cone calorimeter tests were carried out at a heat flux of 35 kW m⁻² to assess the forced combustion behavior of sheets (100 × 100 × 3 mm³) in accordance with ISO 5660 standard [25]. The distance between the sample and the heating cone was increased to 60 mm due to the material swelling. It was assumed that this swelling behavior was attributed to stress release under heat flux, which was formed during cone plate manufacturing. Before performing the tests, all specimens were conditioned at 23 °C and 50% RH for 72 h. Three tests were carried out on each formulation, and the results averaged. According to this analytical method, critical parameters were evaluated, namely: time to ignition (TTI), heat release rate (HRR), total heat release (THR); furthermore, parameters related to smoke release such as total smoke release (TSR), CO₂, CO, and CO₂/CO yield were also evaluated.

3. Results and Discussion

3.1. Morphology of Blends

The dispersion of the lignin and phosphinates in the PA11 matrix was assessed by SEM. The final properties of the blends, such as the flame-retardant performance, depend on the quality of dispersion of the fillers in the polymer matrix [26]. Figure 1a–i shows fractured cross-section images of PA11 and its binary and ternary blends with lignin and phosphinate. It was experienced that during thermo-mechanical compounding the additives size in the polymer matrix is reduced from their original size (Figure S1, Supporting information). From Figure 1b,c, the distribution of LL and DL can be observed for PA₈₀-LL₂₀ and PA₈₀-DL₂₀ binary blends, showing no aggregates, and particles are distributed in the entire surface. However, larger size (less than 10 µm) immiscible lignin particles can be observed due to a higher concentration of lignin in binary blends. In addition to this, PA₈₀-ZnP₂₀ binary blend exhibit distribution of ZnP particles in the polymer matrix. The larger size ZnP particles are observed to be due to broad size distribution in pristine ZnP. It is observed that blends containing ZnP show circular cavity, which can be ascribed to a fusible characteristic of ZnP [27]. However, a micrograph of the PA₈₀-AlP₂₀ binary blend (Figure 1d) shows homogenous distribution and good compatibility between AlP and polymer phase.
characteristic of ZnP [27]. However, a micrograph of the PA80-AlP20 binary blend (Figure 1(d)) shows homogenous distribution and good compatibility between AlP and polymer phase.

Figure 1. Scanning Electron Microscopy (SEM) micrograph of polyamide 11 (PA11) and its binary and ternary blends. (a) PA11; (b) PA80-LL20; (c) PA80-DL20; (d) PA80-ZnP20; (e) PA80-AlP20; (f) PA80-LL10-ZnP10; (g) PA80-DL10-ZnP10; (h) PA80-L10-LL10-ZnP10; (i) PA80-DL10-AlP10 at 5000× magnification.

In order to further assess the dispersion, EDX maps of Carbon and Phosphorus and EDX spectra of ternary blends containing 10 wt % loading of both the fillers are also presented in Figure 2a–e. In fact, it is commonly accepted that EDX mapping spectra can give the qualitative evidence of dispersion and distribution level of the elements [28]. As far as lignin distribution is concerned, it was observed during the blend preparation process that lignin particles are broken down and mixed well with PA11 matrix; furthermore, SEM micrographs show a homogeneous distribution of lignin particles in the polymer matrix, as no phase separation or agglomeration is observed; this indicates reasonable compatibility between lignin and PA11 [16]. In addition to this, Figure 2b,c shows the morphology and distribution of AlP in combination with LL and DL. AlP was found in the entire surface, though small agglomerations (<10 µm) are randomly observed. EDX mapping of phosphorus confirms the uniform distribution of the phosphinate in the blends. Carbon (C), Oxygen (O), Aluminium (Al) and Phosphorus (P) elements were mainly detected and confirmed the presence of AlP and lignin within the polymer matrix. In addition to this, the ternary blends of ZnP with DL/LL (Figure 2d,e) display a larger particle size of ZnP embedded in the matrix; this morphology was observed due to the broad range of size distribution in the pristine ZnP (Figure S1, Supporting information). EDX elemental mapping and corresponding spectra further confirm the identification of C, O, Zn, and P in the entire surface. However, apart from larger size, ZnP particles are uniformly dispersed in the entire surface, without agglomeration phenomena.
3.2. Decomposition Behavior

The thermal and thermo-oxidative stability of PA11 and blends was assessed by TG analyses in N₂ and air atmosphere, respectively. Thermogravimetric data are collected in Table 2. However, TGA curves of neat material are presented in Figure S2 (Supporting information). The thermal stability and degradation profile of the binary and the ternary blends are shown in Figures 3 and 4. In N₂ atmosphere, unfilled PA11 starts decomposing at 396 °C ($T_{5\%}$) and shows single decomposition step with a maximum mass loss at 430 °C ($T_{\text{max}}$) without leaving any residue at the end of the test: a similar behavior was reported in previous studies [16,23]. However, in air, PA11 decomposes in two steps; the first step is at around 454 °C ($T_{\text{max}1}$), giving rise to the formation of volatile products. The second step, at around 574 °C ($T_{\text{max}2}$), can be attributed to the further oxidation of hydrocarbon species with the formation of CO and CO₂, leading to a consistent weight loss until 800 °C.
The char residue from DL containing blends with respect to LL counterpart can be attributed to the loss and a lower char residue at the end of the experiment are observed. The lower production rise to a stable char [30]. Unlike PA–LL–ZnP blends, increasing DL content in its ternary blends of LL, the sulfonate compounds release SO$_4$ and ZnP combination used in our previous study [23]. It is assumed that, during the decomposition interaction within the additives. These blends generate slightly higher residue as compared to the lignin experimental char residue is higher with respect to the theoretical one, thus indicating the positive interaction within the additives. These blends generate slightly higher residue as compared to the lignin and ZnP combination used in our previous study [23]. It is noteworthy that, increasing the LL loading in PA-LL-ZnP blends strongly influences the T$_{5\%}$ and MMLR; as a consequence, the recorded experimental char residue is higher with respect to the theoretical one, thus indicating the positive interaction within the additives. These blends generate slightly higher residue as compared to the lignin and ZnP combination used in our previous study [23]. It is assumed that, during the decomposition of LL, the sulfonate compounds release SO$_2$ and transform it into thermally stable Na$_2$SO$_4$, giving rise to a stable char [30]. Unlike PA–LL–ZnP blends, increasing DL content in its ternary blends slightly influence the T$_{5\%}$ and the thermal stability of the ternary blends; in particular, a higher mass loss and a lower char residue at the end of the experiment are observed. The lower production of the char residue from DL containing blends with respect to LL counterpart can be attributed to.
As far as the thermo-oxidative stability is concerned, the ternary bends show a similar trend, with two decomposition steps: the first one is attributed to the main degradation occurring within 450 to 470 °C ($T_{\text{max1}}$), giving rise to the formation of volatile products such as phosphinate compounds and phosphinic acid; the second step, occurring at around 550 to 610 °C ($T_{\text{max2}}$) is attributed to the formation of thermally stable phosphate compounds in the condensed phase [32,33]. It was noticed that ternary blends containing AlP show a second decomposition step between 600 and 610 °C ($T_{\text{max2}}$) compared to ZnP containing ternary blends, which give $T_{\text{max2}}$ at lower temperatures (i.e., within 550 and 580 °C). This finding confirms that the phosphate compound formation from AlP is more favored than that from ZnP. Furthermore, the residue obtained in oxidative condition was lower compared to the theoretical one; this finding can be ascribed to the presence of an impurity in the lignin due to their industrial nature, which catalyzes a further oxidation with the formation of CO and CO$_2$.

The thermo-oxidative behavior of the prepared blends was assessed through weight loss difference curves $\Delta M$ (%). Those collected in Figure 5a,b, show the interactions among lignin, phosphinates, and PA11. Figure 5a shows stabilization and destabilization region in ternary blends of LL with AlP and ZnP. The blends show slightly destabilization ($\Delta M$ is below $-2\%$) region between 270 and 380 °C and then stabilization from 380 to 460 °C; afterwards, the destabilization region continues until 800 °C. In addition, an increase of $\Delta M$ and thermal stability was observed with increasing LL loading in blends, hence confirming the positive interaction of LL with fire retardant additives. For example, $\text{PA80-LL10-AlP10}$ blend shows stabilization from 415 to 482 °C, and then destabilization of transient char continues up to 800 °C. On the other hand, the presence of DL with AlP and ZnP decreases the thermal stability, as assessed by the destabilization at a lower temperature (within 270 and 400 °C, Figure 5b). Further, $\text{PA–DL–AlP}$ blends show small stabilization region ($\Delta M$ is below 5%) between 400 and 450 °C and large destabilization ($\Delta M$ is below $-16\%$) between 450 and 520 °C due to the evolution of more volatile compounds. Finally, stabilization of transient char continues until 800 °C due to the
formation of a stable phosphate layer in the solid phase. The replacement of AlP with ZnP widens
the stabilization region (ΔM is about 12%) between 400 and 500 °C; then, the destabilization (ΔM is
less than −5%) of transient char continues until 800 °C. Interestingly, increasing the DL content has no
positive influence on mass difference and stabilization region, thus indicating a poor interaction of DL
with fire retardant additive in PA11.

Figure 5. Curves of residual mass loss difference for polyamide 11 (PA11) ternary blends with
lignosulphate lignin (LL) (a) and with kraft lignin (DL) (b) in air.

Table 2. Thermogravimetric data for polyamide 11 (PA11) and its blends in N2 and air.

<table>
<thead>
<tr>
<th>Samples</th>
<th>T5% (°C)</th>
<th>Tmax (°C)</th>
<th>MMLR (%/min)</th>
<th>RExp 1 700 °C (%)</th>
<th>RCal 1 (%)</th>
<th>Tmax1 (°C)</th>
<th>Tmax2 (°C)</th>
<th>MMLR (%/min)</th>
<th>RExp 1 700 °C (%)</th>
<th>RCal 1 (%)</th>
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<td>454</td>
<td>574</td>
<td>1.3</td>
<td>2</td>
<td>-</td>
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1 RExp = experimental residue; RCal = calculated residue.

3.3. Flammability Behavior

The results of the UL94 tests for PA11 and the ternary blends are summarized in Table 3, and the
typical pictures of the specimens left after the tests are shown in Figure 6. However, the UL94 tests
data for the binary blends are reported in Table S2 (Supporting information). Dripping was observed
in all the blends, except PA80-AlP20, which improves the flame-retardant performance and achieves
self-extinction (V-0 rating); further, no melt dripping is observed. This finding can be ascribed to
the release of phosphinic acid and phosphinate compounds in the gas phase, which dilutes the fuel.
Conversely, 20 wt % ZnP loading in PA11 is not sufficient to improve flame retardancy and UL94
rating; in fact, PA80–ZnP20 specimens lead to high flammability and dripping (Table S2, Supporting
information).

In addition to this, the presence of LL in ternary blends decreases the total combustion time.
However, the best flame-retardant results were obtained with PA–LL–ZnP blends, specifically with
7 and 10 wt % loading of LL (i.e., for PA80–LL7–ZnP13, PA80–LL10–ZnP10 blends), which achieve
self-extinction and V-1 classification. Interestingly, PA80–LL10–ZnP10 shows the minimum ZnP loading. It is expected that the presence of sulfonate functionality in LL may lead to the formation of a thermally stable compound. In fact, increasing ZnP loadings deteriorate the performances, as ZnP promotes melt dripping phenomena [34]. As regards to the flame-retardant action, it was assumed that ZnP decomposition produces phosphinate compounds in the gas phase that can release P-O• radicals: this latter can act as radical scavengers and lead to flame inhibition through radical trapping, hence improving the flame retardancy of blends [35,36]. These results demonstrate that the proposed LL/ZnP combinations are potentially effective. Conversely, the presence of DL in ternary blends does not show significant improvement under UL94 tests, as all these ternary blends have a V-2 classification. This behavior was attributed to the rapid decomposition of DL in air; further, DL decomposed mainly (T\text{max1}) during the decomposition of PA11.

### Table 3. UL94 vertical flame spread test data for polyamide 11 (PA11) and its blends.

<table>
<thead>
<tr>
<th>Samples</th>
<th>1st Flame t1 (s)</th>
<th>2nd Flame t2 (s)</th>
<th>Combustion time (t1 + t2)</th>
<th>Cotton Ignition</th>
<th>Dripping</th>
<th>Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>PA11</td>
<td>11 ± 1</td>
<td>7 ± 1</td>
<td>18 ± 1</td>
<td>Yes</td>
<td>Yes</td>
<td>V2</td>
</tr>
<tr>
<td>PA80–LL5–ZnP15</td>
<td>11 ± 1</td>
<td>3 ± 1</td>
<td>14 ± 1</td>
<td>Yes</td>
<td>Yes</td>
<td>V2</td>
</tr>
<tr>
<td>PA80–LL7–ZnP13</td>
<td>4 ± 1</td>
<td>3 ± 1</td>
<td>7 ± 1</td>
<td>No</td>
<td>Yes</td>
<td>V1</td>
</tr>
<tr>
<td>PA80–LL10–ZnP10</td>
<td>6 ± 1</td>
<td>3 ± 1</td>
<td>9 ± 1</td>
<td>No</td>
<td>Yes</td>
<td>V1</td>
</tr>
<tr>
<td>PA80–LL5–AlP15</td>
<td>16 ± 1</td>
<td>6 ± 2</td>
<td>22 ± 1</td>
<td>Yes</td>
<td>Yes</td>
<td>V2</td>
</tr>
<tr>
<td>PA80–LL7–AlP13</td>
<td>24 ± 2</td>
<td>4 ± 1</td>
<td>28 ± 2</td>
<td>Yes</td>
<td>Yes</td>
<td>V2</td>
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<tr>
<td>PA80–LL10–AlP10</td>
<td>8 ± 1</td>
<td>4 ± 1</td>
<td>12 ± 2</td>
<td>Yes</td>
<td>Yes</td>
<td>V2</td>
</tr>
<tr>
<td>PA80–DL5–ZnP15</td>
<td>10 ± 1</td>
<td>4 ± 1</td>
<td>14 ± 2</td>
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<td>V2</td>
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<tr>
<td>PA80–DL7–ZnP13</td>
<td>11 ± 1</td>
<td>4 ± 1</td>
<td>15 ± 2</td>
<td>Yes</td>
<td>Yes</td>
<td>V2</td>
</tr>
<tr>
<td>PA80–DL10–ZnP10</td>
<td>7 ± 1</td>
<td>3 ± 1</td>
<td>10 ± 1</td>
<td>Yes</td>
<td>Yes</td>
<td>V2</td>
</tr>
<tr>
<td>PA80–DL5–AlP15</td>
<td>22 ± 2</td>
<td>5 ± 1</td>
<td>27 ± 2</td>
<td>Yes</td>
<td>Yes</td>
<td>V2</td>
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<tr>
<td>PA80–DL7–AlP13</td>
<td>19 ± 1</td>
<td>4 ± 1</td>
<td>23 ± 2</td>
<td>Yes</td>
<td>Yes</td>
<td>V2</td>
</tr>
<tr>
<td>PA80–DL10–AlP10</td>
<td>18 ± 1</td>
<td>3 ± 1</td>
<td>20 ± 1</td>
<td>Yes</td>
<td>Yes</td>
<td>V2</td>
</tr>
</tbody>
</table>

### Figure 6. Pictures of PA11 blends specimen left after UL 94 vertical flame test. (a) PA11; (b) PA80–ZnP20; (c) PA80–AlP20; (d) PA80–LL20; (e) PA80–DL20; (f) PA80–LL10–ZnP10; (g) PA80–DL10–ZnP10; (h) PA80–DL10–ZnP10; (i) PA80–DL10–AlP10.

### 3.4. Forced-Combustion Behavior

In order to simulate the fire hazards under a real fire scenario, cone calorimetry tests were performed using 35 kW/m² heat flux and 60 mm separation length. Since the distance from the sample surface to the spark igniter increases, the volatiles leaving from the heated sample and the oxygen from air have more time to mix before reaching the spark igniter. This is indicated by an increase in time to ignition. The heat release rate (HRR) and total heat release (THR) are potential fire parameters to evaluate the combustion behaviour of a material exposed to certain heat flux [37,38]. The HRR and THR curves are shown in Figures 7 and 8 and the cone calorimetry data for PA11 and the ternary
blends are collected in Table 4. However, heat release curves and the cone calorimetry data for the binary blends are presented in Figure S3 and Table S3 (Supporting information). Figure 7a–d shows the influence of LL with ZnP/AlP on HRR and THR. In particular, the ternary blends containing LL and ZnP showed lower TTI as compared to unfilled PA11, as anticipated by TG analyses in air: in fact, the adding of 5 to 10 wt % LL strongly reduces its $T_{5\%}$, promoting, at the same time, the formation of a thermally stable char residue. A similar trend was observed during combustion tests: TTI value decreases with increasing LL content from 5 to 10 wt % in blends; this finding is attributed to the rapid mass loss of LL taking place before the decomposition of PA11. Besides, PHRR and THR values substantially drop (by 64% and 22%, respectively) for PA80-LL10-ZnP10 blend, due to the formation of a protective char layer in the condensed phase that can effectively delay the heat release during combustion. LL contains sulfonate compounds, which are likely to decompose during combustion, releasing SO$_2$, thus limiting the heat release and originating thermally stable Na$_2$SO$_4$ in the condensed phase [33,39]. It is noteworthy that the LL and ZnP formulation used in this study showed the enhanced fire performance compared to the cone calorimeter results presented in our previous study with lignin and ZnP combination [23]. When ZnP is replaced with AlP, PHRR remarkably decreases up to 230 kW/m$^2$ for PA80-LL10-AlP10. In addition, HRR curve reveals a broad and single peak of HRR, hence indicating the formation of an effective protective char layer. This superior fire-retardant property in forced combustion tests is due to the presence of an efficient aluminum phosphate layer, able to confer stability to the char residue. The increased char residue at the end of the test (Table 4) further confirms the formation of the protective layer.

Furthermore, Figure 8 shows the influence of DL and ZnP/AlP on HRR and THR. It is worthy to note that, unlike LL lignin, the addition of DL with ZnP does not show a significant reduction in PHRR and THR. However, a certain reduction in PHRR ($-43\%$) is seen when DL content achieves 10 wt %. Besides, the PA–DL–AlP blends show a remarkable reduction of PHRR ($-64\%$) and THR ($-22\%$) for PA80-DL10-AlP10. This is attributed to the presence of AlP, which promotes the formation of a stable char layer acting as a protective barrier, limiting the heat and mass transfer from and to the underlying polymer.

![Figure 7](image_url)

**Figure 7.** Heat release rate (HRR) and total heat release (THR) curves of polyamide 11 (PA11) blends. (a) and (b) for PA–LL–ZnP blends; (c) and (d) for PA–LL–AlP blends.
Flame retardant action can also be confirmed by the effective heat combustion (EHC), which is expressed by the ratio of heat release to mass loss during combustion. EHC reflects the combustion efficiency of flammable volatiles. An obvious gas phase flame retardant action would lead to a noticeable reduction in EHC value as compared to the neat material [40]. Data collected in Table 4 show EHC for PA11 and the ternary blends, while the ternary blends containing 7 and 10 wt % of the lignins show lower EHC values; in this case, the flame inhibition occurs with an incomplete combustion, thus increasing the smoke and the CO release (Table 4) in the gas phase.

Table 4. Cone calorimetry data for polyamide 11 (PA11) and its blends.

<table>
<thead>
<tr>
<th>Samples</th>
<th>TTI (s)</th>
<th>PHRR (kW/m²)</th>
<th>Reduction (%)</th>
<th>THR (MJ/m²)</th>
<th>EHC (kJ/g)</th>
<th>TSR (m²/m²)</th>
<th>CO Yield (g/kg)</th>
<th>CO₂ Yield (kg/kg)</th>
<th>CO₂/CO Residue (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PA11</td>
<td>154 ± 3</td>
<td>884 ± 4</td>
<td>-</td>
<td>92 ± 4</td>
<td>33.8 ± 0.6</td>
<td>1033 ± 1</td>
<td>33 ± 1</td>
<td>2.6 ± 0.1</td>
<td>79 ± 0.6</td>
</tr>
<tr>
<td>PA80-DL10-ZnP10</td>
<td>112 ± 12</td>
<td>560 ± 40</td>
<td>37</td>
<td>79 ± 2</td>
<td>29.3 ± 0.6</td>
<td>1799 ± 48</td>
<td>88 ± 1</td>
<td>1.9 ± 0.1</td>
<td>22 ± 0.2</td>
</tr>
<tr>
<td>PA80-DL7-ZnP13</td>
<td>92 ± 9</td>
<td>443 ± 21</td>
<td>50</td>
<td>77 ± 4</td>
<td>28.8 ± 0.7</td>
<td>1652 ± 151</td>
<td>92 ± 1</td>
<td>2.1 ± 0.1</td>
<td>23 ± 0.2</td>
</tr>
<tr>
<td>PA80-DL7-ZnP15</td>
<td>86 ± 9</td>
<td>315 ± 11</td>
<td>64</td>
<td>73 ± 2</td>
<td>28.4 ± 0.4</td>
<td>1691 ± 26</td>
<td>72 ± 2</td>
<td>2.0 ± 0.1</td>
<td>28 ± 0.3</td>
</tr>
<tr>
<td>PA80-DL10-AlP10</td>
<td>142 ± 11</td>
<td>554 ± 33</td>
<td>37</td>
<td>78 ± 6</td>
<td>30.3 ± 1</td>
<td>1959 ± 28</td>
<td>98 ± 3</td>
<td>1.9 ± 0.1</td>
<td>19 ± 0.3</td>
</tr>
<tr>
<td>PA80-DL7-AlP13</td>
<td>124 ± 10</td>
<td>420 ± 27</td>
<td>52</td>
<td>77 ± 3</td>
<td>29.6 ± 0.3</td>
<td>2034 ± 37</td>
<td>85 ± 3</td>
<td>2.0 ± 0.1</td>
<td>24 ± 0.3</td>
</tr>
<tr>
<td>PA80-DL10-AlP15</td>
<td>108 ± 12</td>
<td>230 ± 14</td>
<td>74</td>
<td>72 ± 4</td>
<td>29.4 ± 0.4</td>
<td>1995 ± 18</td>
<td>71 ± 6</td>
<td>2.1 ± 4</td>
<td>30 ± 0.3</td>
</tr>
<tr>
<td>PA80-DL5-ZnP10</td>
<td>150 ± 18</td>
<td>740 ± 23</td>
<td>16</td>
<td>79 ± 2</td>
<td>29.6 ± 1.3</td>
<td>1720 ± 36</td>
<td>71 ± 3</td>
<td>2.2 ± 1</td>
<td>31 ± 0.8</td>
</tr>
<tr>
<td>PA80-DL7-ZnP13</td>
<td>128 ± 14</td>
<td>678 ± 36</td>
<td>23</td>
<td>77 ± 3</td>
<td>30.2 ± 0.4</td>
<td>1800 ± 88</td>
<td>73 ± 4</td>
<td>2.1 ± 0.1</td>
<td>29 ± 0.2</td>
</tr>
<tr>
<td>PA80-DL7-ZnP15</td>
<td>116 ± 13</td>
<td>500 ± 48</td>
<td>43</td>
<td>75 ± 7</td>
<td>30.3 ± 1.2</td>
<td>1745 ± 62</td>
<td>61 ± 6</td>
<td>2.1 ± 0.1</td>
<td>34 ± 0.2</td>
</tr>
<tr>
<td>PA80-DL7-AlP10</td>
<td>174 ± 14</td>
<td>424 ± 39</td>
<td>42</td>
<td>76 ± 4</td>
<td>28 ± 1</td>
<td>1819 ± 40</td>
<td>99 ± 2</td>
<td>2.0 ± 0.1</td>
<td>20 ± 0.3</td>
</tr>
<tr>
<td>PA80-DL7-AlP13</td>
<td>140 ± 12</td>
<td>406 ± 34</td>
<td>54</td>
<td>74 ± 5</td>
<td>31 ± 1</td>
<td>1929 ± 45</td>
<td>90 ± 3</td>
<td>2.0 ± 0.1</td>
<td>22 ± 0.3</td>
</tr>
<tr>
<td>PA80-DL5-AlP15</td>
<td>95 ± 7</td>
<td>320 ± 10</td>
<td>64</td>
<td>72 ± 1</td>
<td>29 ± 1.3</td>
<td>1967 ± 68</td>
<td>86 ± 5</td>
<td>2.1 ± 0.1</td>
<td>24 ± 0.4</td>
</tr>
</tbody>
</table>

3.5. Smoke and CO Release

The release of smoke and CO during combustion is not only a key fire hazard, but also an indication of the flame-retardant mechanism. For instance, a significant increase in the smoke release and CO amount is originated by incomplete oxidation of gaseous products, which indicates a flame inhibition action by radical trapping reactions taking place in the gas phase. Alternatively, a decrease in smoke release indicates a better-ventilated combustion process, in which the flame retardancy is dominated by fuel dilution and/or thermal barrier [35,41]. Therefore, total smoke release (TSR),
CO and CO$_2$ total yield, and CO$_2$/CO yield were assessed; the corresponding data are collected in Table 4. First of all, the CO$_2$ yield is practically similar for all the blends. As compared to PA11, binary blends with ZnP or AlP increase the smoke release and CO yield and lower the CO$_2$/CO yield, hence indicating the incomplete oxidation of gaseous products. Conversely, the incorporation of LL and DL lowers TSR and CO yield and increase CO$_2$/CO yield, also confirming the extended oxidation of the evolved gaseous products (Figure S4, Supporting information).

On the other hand, the combination of lignin and phosphinate worsens the smoke parameters, confirming the occurrence of incomplete combustion. However, increasing the lignin content reduces the CO release, which is attributed to the more complete oxidation of evolved volatile products. Figure 9 shows the influence of LL mixed with flame retardants on CO and CO$_2$ evolution. Interestingly, the blends containing 10 wt % of both fillers show the lowest CO release with respect to other ternary blends: more specifically, PA$_{80}$–LL$_{10}$–ZnP$_{10}$ blend significantly lowers the peak of CO release up to 300 ppm without compromising the other fire-retardant properties. This finding can be ascribed to the interactions taking place between LL and ZnP in the ternary blend, which leads to extended oxidation of the evolved gaseous products as well as to the formation of a protective char in the condensed phase. Conversely, the peak of CO evolution reported in our previous study with lignin and ZnP combination showed a slightly higher, up to 500 ppm, for the same blend (i.e., 10 wt % concentration of both additives) [23]. Based on results, it can be concluded that the combination of lignin and phosphinate promotes the smoke and CO release; however, high lignin loading remarkably reduces CO release, hence minimize the smoke toxicity.

![Figure 9](image.png)

**Figure 9.** CO and CO$_2$ evolution during combustion. (a) and (b) for PA–LL–ZnP blends; (c) and (d) for PA–LL–AlP blends.

### 3.6. Morphology of Char Residue

Figure 10 shows the pictures of residues obtained after forced combustion tests. It is clearly observed (Figure 10a) that the residue from PA11 is very thin and practically negligible. Similarly, PA11 with ZnP (Figure 10b) shows a mechanically thin and non-charring characteristic. However, PA$_{80}$–AlP$_{20}$ slightly increases the residue (Figure 10c), although the presence of cracks and insufficient residue leads to the formation of a weak protective layer. Besides, the binary blends of lignin (LL
and DL) display a higher char residue formation: in particular, PA$_{80}$-LL$_{20}$ (Figure 10d) gives rise to a coherent and compact char layer formation, which protect the material against heat flux and reducing the release of flammable and non-flammable gases throughout the polymer. However, the presence of cracks deteriorates the performance. Similarly, the char residue of PA$_{80}$-DL$_{20}$ shows a loose and porous surface with many cracks (Figure 10e). Interestingly, the combination of lignin and phosphinate not only increases the char residue but also leads to the formation of a stable char layer with barrier features against heat flux and release of combustible gases. In particular, the addition of LL with ZnP/AlP (Figure 10f,g) shows a compact and protective layer due to sufficient char formation. Conversely, ternary blends of DL show a thin char layer and relatively loose structure with many cracks; more specifically, PA$_{80}$-DL$_{10}$-ZnP$_{10}$ (Figure 10i) gives a fragile and mechanically weak char layer. In order to further investigate the microscopic morphology and structure of the charred layer, the top surface of char residues was observed by SEM (Figure 11a–d). It can be observed that the surface morphology of residue from PA$_{80}$-LL$_{10}$-AlP$_{10}$ (Figure 11a) appears denser and more compact due to the increased char formation and interaction between LL and AlP leading to a stable protective char layer, which contributes to improved flame retardancy. Furthermore, EDX analysis reveals the presence of C, O, Na, Al, and P, which results in a more compact char residue. Similar char residue morphology with AlP is reported [42,43]. As shown in Figure 11b, PA$_{80}$-LL$_{10}$-ZnP$_{10}$ presents a relatively less dense and compact structure in comparison with PA$_{80}$-LL$_{10}$-AlP$_{10}$, although compact char layer is present with some holes on the surface. In contrast, the surface morphology of residue from DL containing blends (Figure 11c,d) does not form an intact char layer due to insufficient char formation during combustion; as a consequence, a loose and porous structure is formed.

**Figure 10.** Pictures of char residues collected at the end of cone calorimetry test: (a) unfilled polyamide 11 (PA11); (b) PA$_{80}$-ZnP$_{20}$; (c) PA$_{80}$-AlP$_{20}$; (d) PA$_{80}$-LL$_{20}$; (e) PA$_{80}$-DL$_{20}$; (f) PA$_{80}$-LL$_{10}$-AlP$_{10}$; (g) PA$_{80}$-LL$_{10}$-ZnP$_{10}$; (h) PA$_{80}$-DL$_{10}$-AlP$_{10}$ and (i) PA$_{80}$-DL$_{10}$-ZnP$_{10}$ blends.
were combined with phosphinate (ZnP and AlP) flame retardants in PA11. Through UL 94 vertical flame spread tests, it was possible to demonstrate that the combination of lignins and phosphinates up to 20 wt % minimizes the particle size due to the thermal mixing; in addition, EDX elemental mapping revealed a suitable dispersion of lignin and phosphinate within the polymer matrix. However, some small (<10 µm) agglomerates were observed in AlP containing blends. TG analyses showed that, regardless of its type, the incorporation of lignin in the ternary blends increased the thermal stability of PA11, promoting the obtainment of a stable char residue at the end of the tests. Through UL 94 vertical flame spread tests, it was possible to demonstrate that the combination of LL with ZnP effectively improves the flame-retardant properties by reducing total combustion time. More specifically, PA80–LL10–ZnP10 blends at 1000× and 5000× magnification.

**4. Conclusions**

So as to advance the exploitation of industrial lignin, in this work, different lignins, i.e., LL and DL, were combined with phosphinate (ZnP and AlP) flame retardants in PA11. Different flame retarded blends were prepared using melt extrusion. The influence of the presence of lignin with phosphinate on the thermal and fire behavior was thoroughly assessed by TG analyses, UL 94 vertical flame spread, and cone calorimetry tests. SEM analyses of the blends showed that the addition of lignin and phosphinates up to 20 wt % minimizes the particle size due to the thermal mixing; in addition, EDX elemental mapping revealed a suitable dispersion of lignin and phosphinate within the polymer matrix. However, some small (<10 µm) agglomerates were observed in AlP containing blends. TG analyses showed that, regardless of its type, the incorporation of lignin in the ternary blends increased the thermal stability of PA11, promoting the obtainment of a stable char residue at the end of the tests. Through UL 94 vertical flame spread tests, it was possible to demonstrate that the combination of LL with ZnP effectively improves the flame-retardant properties by reducing total combustion time. More specifically, PA80−LL10−ZnP10 and PA80−LL7−ZnP13 achieved self-extinction and V-1 rating. Furthermore, in forced combustion test the interactions between lignin and phosphinate promoted a remarkable reduction of PHRR and THR. In particular, the best fire-retardant performance was found by combining LL with AlP (i.e., PA80−LL10−AlP10), resulting in a strong reduction of PHRR (~74%) and THR (~22%) values. On the other hand, smoke parameters, namely TSR and CO yield, increased in ternary blends; however, increasing the lignin loading effectively reduced the CO and CO\textsubscript{2}/CO yield compared to PA11. Particularly, PA80−LL10−ZnP10 blend shows minimum CO evolution without affecting flame retardancy. The morphology of char residue showed that the formation of compact char layer is primarily responsible for the improved flame-retardant properties. In conclusion, from an overall point of view, the direct use of industrial lignin with phosphinate in PA11 seems to be quite promising as far as fire retardancy is concerned, also considering the “biobased” character of both the polymer matrix and the lignins.
Supplementary Materials: The following are available online at http://www.mdpi.com/2073-4360/11/1/180/s1, Figure S1: SEM images showing particle size distribution of pristine material, (a) ZnP, (b) AlP, (c) DL, (d) LL. Figure S2. TG curves of unfilled PA11 and neat materials in N2 (a) and air (b), Figure S3. HRR and THR curves of PA11 and the binary blends, (a) and (b) PA80-Ll20, (c) and (d) PA80-Dl20, (e) and (f) PA80-ZnP20, (g) and (h) PA80-AlP20 blends. Figure S4. CO and CO2 evolution during combustion for PA11 and the binary blends, (a) and (b) PA80-Ll20, (c) and (d) PA80-ZnP20, (e) and (f) PA80-AlP20 blends. Table S1: Thermogravimetric data for PA11 and its binary blends, Table S3. Cone calorimetry data for PA11 and its binary blends.


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Conflicts of Interest: The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision to publish the results.

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