Alkyl sulfone bridged phosphorus flame-retardants for polypropylene

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Abstract. Three novel alkyl sulfone bridged phosp' orus (P) compounds namely 6,6'-(sulfonylbis(ethane-2,1-diyl))bis(dibenzo[c,e][1,2]oxaphc sphi ine 6-oxide) (SEDOPO), sulfonylbis(ethane-2,1-diyl))bis(diphenylphosphine oxide) (3EDPPO), and tetraphenyl (sulfonyl bis(ethane-2,1-diyl)) bis(phosphonate) (SEDPP) (i.e. phosphine oxide, phosphinate and phosphonate, respectively) were synthesized via a Micha 1 ac dition reaction with good yields (≥ 85%) at a 200gram scale. They exhibited thermal stability above 250 °C, which allowed them to be melt-processed with polypropylene (PP) and formed into thin films (~ 0.6 mm). Rheological measurements of the PP blends exhibited a typical shear thin null behavior and provided evidence for the synthesized compound's thermo-oxidative stabilizing effect. This was also confirmed by thermal analysis showing that the thermo-oxidative stability of PP-SEDOPO and PP-SEDPPO blends was higher (~25 °C) than the blank PP; how ever PP-SEDPP had a smaller impact. Small scale fire tests of the PP-FR blends confirmed the flame retardant efficacy of the new P-compounds. Cone calorimetry on PP-SEDOPO blends showed a reduction in the heat release rate (HRR) (~48%) compared to blank PP. Further thermal and evolved gas analysis of the PP blends confirmed that the new P-compounds are primarily active in the gas-phase.

Keywords: Phosphorus, flame retardant, bridged sulfone, Sulphur, alkyl sulfone, Polypropylene, thermal analysis, fire tests, LOI.

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1. Introduction

Polypropylene (PP) is a common thermoplastic polymer, which provides a good balance of physical, chemical, mechanical, electrical, and thermal properties with easy processability. This combination of attractive features drives its application in various industries such as electronics, automotive, textiles, construction, and packaging [1-3]. PP is highly combustible due to its aliphatic chemical composition [4]. Combustion of PP yields high amounts of toxic gases that threaten human safety and restrict its usage in fire-safe applications [5, 6]. A commonly used method to reduce thermoplastic flammability is introducing a flame retardant (FR) additive into the bulk of material via melt processing. Typically, flame retardancy is not the only requirement to be fulfilled by such an FR additive. The FR additive cost of production is a crucial factor for a successful commercial application. Hence, selecting suitable FR additives or the development of new FR systems is a challenging task and mostly depends on the final product specifications and potential fire scenario. Compared to bulk thermoplastic products, flame retardation of fibers and thin films face additional processing and durability challenges[7]. For thin film and fiber applications, the FR additive needs to be either very small in particle size (<1 µm) or in 1 liquid (meltable) form during its passage through small slits or spinneret holes. The latter ty, or additive is preferred for fiber applications as nonmelting solids tend to agglomerate and clog the spinneret.

The conventional flame-retardants for Pr include halogen-based FRs, inorganic FRs, organic halogen-free FRs, phosphorus-convaining intumescent FRs (IFR), and metal hydroxide FRs [8, 9]. Blends of halogenated compounds with antimony trioxide (Sb₂O₃) can be used as FRs for PP; however, the blend can either produce toxic gases in combustion [10] or are themselves toxic [11]. Inorganic flame retardants such as aluminum hydroxide Al(OH)₃ and magnesium hydroxide (Mg(OH)₂ are used as FP for PP, but their high loading requirements (~40 wt%) can result in materials with inferior rheological and mechanical properties [12-15] and thus are not suitable for film and fiber fabrication.

N-alkoxy hindered amines (NORs) are commonly used in fiber and thin-film applications [16-20]. They possess excellent flame retardancy due to the dissociation of NORs forming free radical scavengers that interrupt and suppress the ignition process of polyolefins [21, 22]. Moreover, NORs can interact with conventional FRs via radical reactions to improve their flame-retardants efficiency and reduce the significant loading of traditional FRs [21, 22]. Ciba[®] FLAMESTAB[®] NOR[®] 116 (now BASF) is disclosed in patents as a synergist with conventional brominated and/or phosphorus-based

FRs for polyolefins [23, 24]. The synergistic effect of FLAMESTAB® NOR 116 [25] in combination with ammonium polyphosphate (APP) and small amounts of nanoclay in PP is also reported [26].

The effectiveness of sulfur(S)-containing compounds as FRs has been investigated [27]. They have also been used as co-additives with other FRs, most notably phosphorus compounds, to boost the FR performance of materials [28-35]. Several S-containing compounds have been studied as FRs in polycarbonates (PC) [36], polyamides 6 [37, 38], cotton and wool [39], polyurethanes [40], and polystyrene or polymethyl methacrylate [41]. The mechanism for FR of S-containing compounds has been found to vary based on different polymer systems. The thermal decomposition of elemental sulfur, sulfide, and disulfide generates sulfur radicals, which may promote crosslinking at the polymer surface [31, 32]. Moreover, S-containing FRs during the nal decomposition generates sulfur dioxide (SO₂) in the gas phase [30], which acts not only as a fuel-diluent, but may also act as a radical-scavenger to suppress combustion efficiency [42, 43]. Generally, the flame retardancy of the materials improves as the level of oxygenation at substractions (sulfone > sulfoxide > sulfide) [44]. Thus, a strategy that combines a phosphorus compound with a sulfur moiety for the development of a flame retardant is worthy of investigation. Development of meltable compounds for use as a flame retardant in polyolefins is reictively unknown and will be useful for the advancement of fire safe fibers and films.

In this study, sulfur and phosphorus-based oridged compounds (i.e., phosphine oxide, phosphinate, and phosphonate, Fig. 1) were synthesized by adding P(O)-H containing phosphorus precursors to divinyl sulfone through a straightforthard approach. The chemical structure and purity of synthesized FRs were characterized via Note and elemental analysis. The alkyl sulfone bridged phosphorus derivatives SEDOPO, SEDED and SEDPP were processed with PP using extruders and internal mixers and transformed into thin plates $(160 \times 60 \times 0.6 \text{ mm})$ and films (0.6 mm) via compression molding and extrusion, respectively. The PP-FR blends were then analyzed for their thermal, rheological, and flammability properties. Based on the various thermal and evolved gas analysis, a tentative mode of action of the bridged sulfone compounds was proposed.



Fig. 1. Structure of alkyl sulfone bridged phosphorus compounds

2. Experimental Section

2.1. Materials and methods

9,10-dihydro-9-oxa-10-phosphaphenanthrene-10-oxide (DOPO) was purchased from Metadynea GmbH (Austria), diphenylphosphine oxide (DPPO) was purchased from Combi-Blocks (Germany) divinyl sulfone was purchased from Fluorochem. Diphenyl phosphonate, diazabicyclo(5.4.0)undec-7-ene (DBU) and dry toluene (99.98%) were purchased from Sigma Aldrich (Switzerland) and used without any further purification. The polypropylene (PP) HSP165G (Braskem, The Netherlands) was purchased from Resinex Switzerland AG. 6,6'-(sulfonylbis(ethane-2,1-diyl))bis(dibenzo[c,e][1,2]oxaphosphinine 6-oxide) (SEDOFO), sulfonylbis(ethane-2,1diyl))bis(diphenylphosphine oxide) (SEDPPO) [45] and tetraphen via (sulfonyl bis(ethane-2,1-diyl)) bis(phosphonate) (SEDPP) were synthesized via optimized procedures. The details of optimized procedures and synthesis of FRs are described in the Supporting Information (SI), Sec. SI-1 2.2. Synthesis of bridged sulfone compounds.

2.2. Synthesis of bridged sulfone compounds

6,6'-(sulfonylbis(ethane-2,1-diyl))bis(dibenz [r,e][1,2]oxaphosphinine 6-oxide) (SEDOPO): A dry three-neck, round-bottom flask connected to a mechanical stirrer, nitrogen (N₂) inlet, thermometer, and a condenser was charged with DOFC (187.38 g, 867.5 mmol), divinyl sulfone (50 g, 423.2 mmol), and toluene (600 mL) at room emperature. After the addition was completed, the reaction mixture was heated at 130 °C for 15 hours and cooled to room temperature. The reaction mixture was filtered and the solids were washed with acetone and dried at 80 °C overnight under vacuum to obtain a white powder with a 92% yiel I (Scheme 1).

sulfonylbis(ethane-2,1-diy,1)bis(diphenylphosphine oxide) (SEDPPO): A dry three-neck, round-bottom flask connected to a mechanical stirrer, N₂ inlet, thermometer, and a condenser was charged with diphenylphosphine oxide (DPPO) (82.89 g, 410 mmol), divinyl sulfone (23.63 g, 200 mmol) and toluene (500 mL) at room temperature. After the addition was completed, the reaction mixture was heated to 130 °C for 16 hours and then cooled to room temperature. The solid residue was filtered and washed with acetone and dried at 70 °C overnight under vacuum to obtain a white powder with (187g) 91% yield (Scheme 1).

Tetraphenyl (sulfonyl bis(ethane-2,1-diyl)) bis(phosphonate) (SEDPP): A dry three-neck, round-bottom flask connected to the mechanical stirrer, N_2 inlet, thermometer, and a condenser was charged

with diphenyl phosphonate (203.16 g, 867.5 mmol), divinyl sulfone (50 g, 423.5 mmol) and toluene (1 L) at room temperature. The reaction mixture was cooled to 0 °C and DBU (12.88g, 84.6 mmol) was added dropwise over 1 hour. After the addition of DBU, the reaction mixture stirred for additional 2 hours at 0 °C. The solids were filtered and washed with methanol and dried at 60 °C under vacuum overnight to afford 221 g (89%) white powder (Scheme 1).

2.3. Polymer processing

Kneading trials of bridged sulfone compounds were performed by kneading in an internal laboratory mixer (HAAKE Rheomix 600, Thermo Fisher Scientific, Germany) equipped with Banbury type rotor design at a 50 g scale. The blends were prepared in the kneader at 30 rpm for 10 minutes with a theoretical mass ratio of 90:10 and 85:15 (wt/wt) % of PP: FR add fives. A processing temperature of 250 °C for SEDOPO, 220 °C for SEDPPO and SEDPP blends were used for their processing. These PP blends were used to perform thermogravimetric analysis (TCA), differential scanning calorimetry (DSC), and microscale combustion calorimetry (MCC) is oreover, PP plates (160 × 60 × 0.6 mm) were obtained using a compression molding machine Battenfeld Plus 350/75 (Battenfeld Technologies, Columbia, MO, USA), at 250 °C.

Extrusion trials: The main steps for polyme, film production included compounding and melt-extrusion of films. First, a 15% masterbatch of SEDOPO in a mixture of PP was prepared by melt extrusion in a co-rotating 36 L/D twin-straw extruder (Dr. Collin GmbH, Germany). To achieve a high SEDOPO loading in the matteratch, a side feeder (Dr. Collin GmbH, Germany) was used to add the FR powder to the polynor melt. The extruder was operated at 110 rpm with a constant throughput of 0.6-1 kg/h. The extrudate was quenched on a conveyor belt down to room temperature and subsequently pelletized. Second, film extrusion was carried out on a lab-scale extrusion line with a 25 L/D single screw extruder (Rheomex OS, Thermo Fisher Scientific, Germany). FR masterbatch (SEDOPO) pellets were premixed with the blank polymer and fed to the extruder. A static mixer was installed between the extruder and melt pump (Haake OS, Thermo Fisher Scientific) to promote additive dispersion. The extrudate film, pre-shaped by a custom-made 75 mm slit die, was quenched and molded by chill rolls and finally wound up. In both compounding and extrusion processes, the extruder's temperature profile ranged between 240-250°C.

To evaluate the dispersion of FRs in PP, energy dispersive X-ray (EDX) analysis was performed on the sample surface and in cross-sectional view, using a Hitachi S3700N scanning electron microscope (SEM) equipped with an EDAX Octane Pro SDD EDX detector. The SEM device was operated at 7

kV under low vacuum conditions (90 Pa) to avoid beam damage. As shown in Figure S4, phosphorus and sulfur are present everywhere in the 15% PP-SEDOPO (surface and bulk material showed in cross-sections) films. While the SEDOPO is relatively homogeneously distributed over the cross-section, areas with slightly increased P and S concentrations are found at the surface. Point analysis at different positions confirmed the FR distribution over the whole sample.

2.4. Characterization

NMR spectra were recorded on a Bruker AV-III 400 NMR spectrometer (Bruker Biospin AG, Fällanden, Switzerland) at 298 K. The spectral data and further details about NMR analysis are available in SI part sec. SI-2. NMR characterizations.

Phosphorus content of the PPs blends was measured with a 5110 inductively coupled plasma optical emission spectrometer (ICPOES, Agilent Switzerland AG, Parel, Switzerland). Sample preparation for ICP-OES consisted of mixing 200 mg of a sample with 2 mL HNO₃, followed by microwave digestion.

Differential scanning calorimetry (DSC): For DSC $^{\circ}$ and ysis, NETZSCH DSC Polyma 214 was used to analyze the additives melting point and gless t ansition temperature (T_g) of the PP matrix. For DSC experiments, ca. 5–8 mg of samples were used $^{\circ}$ n a low-pressure crucible and heated to 300 $^{\circ}$ C at a heating rate of 10 $^{\circ}$ C/min.

The thermogravimetric analysis (TCA) on the PP samples and synthesized FRs was carried out using a NETZSCH TG209 F1 Iris instrument. For TGA experiments, ca. 3-5mg of samples were heated from 25 °C to 800 °C at a heating rate of 10 °C/min. The measurements were performed under a nitrogen and air atmosphere with a total gas flow of 50 mL/min.

Limiting Oxygen Index (ICI) values of all plates were measured on an FTT oxygen index apparatus according to ASTM D2863-97. The dimension of specimens for the LOI test was $150 \times 50 \times 0.5$ mm and the samples (plates) were prepared via compression molding.

Vertical burning tests (small scale fire tests) were performed according to the Swiss Standard (BKZ-VB) as described in the literature [46]. PP plates in a vertical orientation were exposed to a flame (45°) for 15 s and the burn length and burn time were measured. Most thermoplastic materials drip when exposed to fire and adequate heat. Thus, in the vertical burning tests, the dripping behavior of the specimens was also observed. The dimension of the specimen for the Vertical burning tests was $160 \times 60 \times 0.6$ mm and the sample was prepared via compression molding.

Direct insertion probe mass spectrometry (DIP-MS) was performed in a Finnigan/Thermoquest GCQ ion trap mass spectrometer (Austin, TX, USA) equipped with a DIP module. ~1 mg of sample was placed in a quartz cup located at the tip of the probe and was inserted into the ionization chamber operating at an ionization voltage of 70 eV, a temperature of the ionic source of 200 °C and pressure $<10^{-6}$ mbar. The probe was heated from 30 to 450 °C at 50 °C/min.

TGA-FTIR was performed using around ca.10 mg of the samples and heated from 25 to 800 °C at a rate of 10 °C/min⁻¹ under a nitrogen purge at 300 ml min⁻¹. The FTIR (Fourier transform infrared spectroscopy) extracted spectra of the pyrolysis gases formed from the samples were measured by Bruker Tensor 27 Fourier transform spectrophotometer coupled with a TGA. The FTIR analysis was performed in the spectral range of 4000 to 550 cm⁻¹ at a resolution of 4 cm⁻¹.

Microscale Combustion Calorimeter (MCC) Heat release rate. (HRR) were determined using MCC (Fire Testing Technology Instrument, London, UK) following ASTM D7309. ~7 mg of sample was exposed to a heating rate of 1.0 °C/s from 150 to 750 °C in the pyrolysis zone.

Cone calorimetry (Fire Testing Technology, East Cirstead, London, UK) was performed with an irradiative heat flux of 35 kW/m² (ISO 5650 standard) on specimens (100×100×3 mm³) placed horizontally without any grids. Parameters such as heat release rate (HRR), peak of heat release rate (pHRR), total heat release (THR), total smoke release (TSR), CO production (COP), average specific extinction area (av-SEA), and CO2 cmput (CO2) were recorded for each sample. The cone calorimetry samples were prepared via compression molding.

Activation energy (Ea): Variation in Ea with the degree of conversion for blank PP and PP-15% SEDOPO, SEDPPO, and SEDPP blends were calculated using the Kissinger method using NETZSCH Kinetics Neo sor ware, version 2.4.6 [47]. For the kinetics experiment, 4 different heating rates (i.e., 2.5, 5, 7.5, and 10 °C /min) were used where samples were heated from 25 °C to 800 °C under air atmosphere with a total gas flow of 50 mL/min.

Rheological measurements were carried out on an Anton Paar Physica 301 MCR rotational rheometer (Austria). All experiments were performed at a constant temperature of 250 °C in the air using a parallel plate fixture (plate diameter of 25 mm and 1 mm gap). Samples were compression molded into plates using a hot press at 250 °C. Before starting a measurement, the samples were relaxed for 5 minutes in the rheometer to minimize the deformation history and get a homogeneous melt. Afterward, angular frequency sweep experiments were conducted at a constant strain of 1%.

Tensile test: The load-strain behavior of dogbone film specimens (Fig. S5) cut in film production direction was evaluated using the Zwick Z100 (Zwick Roell GmbH, Germany) tensile testing machine, equipped with a 100 N load cell. The tensile tests were performed at 23°C and 50% relative humidity, with an initial gauge length of 50 mm and a constant elongation rate of 20 mm/min. The starting length of the neck was 25 mm, which was considered a reference length for assessing strain. Stress was calculated by dividing the measured load by the initial cross-section area, i.e., the film thickness multiplied with the neck's width (4 mm). The sample used for the tensile test was the film (0.6 mm) obtained via slit extrusion.

3. Results and Discussion

3.1. Synthesis of Flame Retardants and analysis

The addition of the P(O)-H bond to an alkene is a vital ransformation for the preparation of organophosphorus compounds with P-C bonds. This addition is often promoted by bases, radical initiators (benzoyl peroxide, AIBN), transition me'ai, and microwaves [45, 48-51]. FR additives consisting of two moieties based on sulfur dioxide and phosphorus were synthesized in this study via the addition of P(O)-H containing precurse so divinyl sulfone. The synthesis of SEDOPO and SEDPPO has been reported in the literature before, but the previous methodology was not feasible for industrial scale-up [45]. The synth sis of 1,2-bisphosphorylethane compounds using toxic trimethylphosphine as a catalyst is previously reported [48]. SEDPPO has been synthesized in solvent-free conditions in an alternative synthesis strategy by reacting one equivalent of divinyl sulfone and two equivalents of diphenylphosphine oxide using a microwave-assisted doublehydrophosphorylation of all ane, on a 50 mg scale [45]. The preparation of a phenylphosphine oxidebridged phosphorus compound from diethylphosphonic acid using a phosphine catalyzed synthesis is also reported in the literature [52]. With a view to industrial-scale production, the use of alkyl phosphine catalysts is highly undesirable considering their toxicity and potential fire hazard. On the other hand, the microwave heating of a solvent free reaction mixture is challenging, particularly when applied to a high-temperature melting reaction mixture. Accordingly, there is a need for an improved approach to preparing bridged phosphorus-based compounds as these compounds were never tested as flame retardants previously.

Scheme 1: Synthesis of alkyl sulfone bridged phosynous compounds SEDOPO, SEDPPO, and SEDPP.

The addition of DOPO (1) to divinyl sulfor, was chosen as a model reaction and detailed procedures of various reaction conditions are described in SI-1. By following the optimized reaction conditions (method 3, Table S1), SEDOPO and SEDPPO were further synthesized at ~200g scale with high yields (93% and 91%, respectively) (Scheme 1). Surprisingly, diphenyl phosphite reaction with divinyl sulfone and toluene as a solvent at reflux condition yielded no final product (SEDPP). However, when a catalytic amount of DBU (10 mol %) was added to the reaction mixture at 0 °C, the desired addition product (SEDPP) was obtained at 89% yield (Scheme 1).

3.2. Thermal processing and rheological behavior

A practical approach to preserving the desired mechanical and thermal properties of PP is to develop a blend with a minimum loading of the FR additive. Usually, a meltable additive can exhibit higher compatibility with PP as compared to non-meltable additives. To evaluate the effect of the bridged sulfone compound on the properties of PP, compounded blends were obtained in the kneader with a theoretical loading mass ratio of 90:10 and 85:15 (wt/wt) % of PP: additives. In general, the effectiveness of a phosphorus-based FR additive relies on phosphorus concentration in the blend. Therefore, it should be noted that the measured phosphorus content in the PP blends containing

SEDOPO and SEDPPO additives are very close to the theoretical predictions (Table 1). In contrast, the phosphorus content is almost 40% lower than the theoretical concentration in PP-SEDPP blends. This difference in the phosphorus content of PP-SEDPP blends could be due to polymer and additive's poor compatibility. Being polar (presence of P-O-C bond), SEDPP demixes and separates from PP during the kneading process (Table 1), which results in lower phosphorus contents.

Moreover, it is critical to understand the effects of FR additive on the rheological behavior of the polymer and on its melt stability. The complex viscosity of different PP blends with frequency sweep rheological experiments were characterized and are presented in Fig. 2a, b. Before kneading, the blank PP sample shows a typical Newtonian viscosity at low frequencies followed by a typical shear thinning behavior at higher frequencies. After PP extruded under the conditions explained before, its viscosity reduced, indicating a loss of molecular weight due to the rmomechanical degradation. The presence of SEDOPO in PP prevents its degradation; hence a higher viscosity for the blend is achieved, Fig. 2a. Similar behavior was also observed for PP blends containing other bridged sulfone compounds (SEDPPO and SEDPP) with some differences in effectiveness, Fig. 2b. It has been shown in earlier studies that this protection mechanism of phosphorous additives arises due to a combination of (i) molecular lubrication that reduces stees, on polymer chains and (ii) released phosphorous radicals that prevent oxidative attacks on polymer [46, 53].

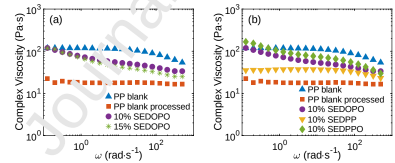


Fig. 2: Rheological behavior of melts at 250 °C in air.

3.3. Small scale fire tests

PP films and fibers need to be fire-resistant in specific applications, for instance, electronics, transportation, and textiles. PP-FR blends were fabricated into plates (0.6 mm thickness) resembling textile fabrics and evaluated for their flammability via a small-scale vertical fire test (BKZ-VB) and limiting oxygen index (LOI) (Table 1). [54, 55]. Videos of vertical burning tests and digital photos of residues after burning (Fig. S6) test for blank PP and PP-FR blends are available in

SI for blank PP, PP-10% SEDOPO, PP-15% SEDOPO, PP-10% SEDPPO, PP-10% SEDPPO, PP-10% SEDPP and PP-15% SEDPP. The blank PP burned readily without leaving any char in the vertical burning test, as shown in Fig. S6. However, on incorporating bridged sulfone derivatives in PP, a decline in the burning length of PP was observed compared to the blank PP (Fig. S6). No further ignition was observed for the plates containing 10 or 15% SEDOPO after removing the flame, which confirms their self-extinguishing behavior. An increased concentration of bridged sulfone compounds in PP reduced the burning length and duration of the burning. Not much char was formed for the samples in the vertical burning test; it is more likely that the bridged sulfone derivatives primarily work in the gas phase. Furthermore, the PP blends with10% or above loading exhibited no or few drips than under the plank PP. The melt dripping was suppressed completely for PP samples containing a minimum of 10% SEDOPO and 15% for other bridged sulfone compounds. PP blends containing 5% bridged sulfone compounds were also manufactured; however, they didn't pass the BK.7-VB test and thus were omitted from any further analytics.

The limiting oxygen index (LOI) value for the Clank PP plate is 19.4%[56]; however, the LOI values for various PP blends in this work were significantly higher than the blank PP. Increased LOI values were observed for increased concentration of bridged sulfone compound in PP and the highest LOI was observed for PP containing 15% SEDOPO. As discussed earlier, it is likely that all bridged sulfone compounds work primarily in the gas phase as mostly gas-phase active FRs have a superior influence on promoting the LOI values of polymeric materials.

Table 1: Summary of small-scale fire tests and the limiting oxygen index (LOI).

			1			
Samples	Predicted P. [wt%]	Measured P. [wt%]	After Flame(sec) ±1	Burning Length(cm) ±1	Melt Drips	LOI % ±0.2
PP-Blank			40	15	Many	19.4
PP-10% SEDOPO	1.12	1.10	0	2.0	No	24.3
PP-15% SEDOPO	1.69	1.54	0	1.4	No	25.7
PP-10% SEDPPO	1.18	1.09	0	4.6	Few	22.7
PP-15% SEDPPO	1.78	1.54	0	4.2	no	23.5
PP-10% SEDPP	1.05	0.63	3.5	7.0	Many	22.4
PP-15% SEDPP	1.59	0.90	3	6.2	Few	24.9

^a Swiss standard vertical fire test.

3.4. Flammability: pyrolysis combustion flow calorimetry (MCC) measurements

MCC was employed to understand the combustion behavior of the PP blends (Table 2). Fig. S7 illustrates the heat release rate (HRR) profile versus temperature for the blank PP and PP blends. For the blank PP, a peak heat release rate (pHRR) of 1120 W·g⁻¹ at 484 °C and a THR of about 45.33 kJ·g⁻¹ was observed. The addition of SEDOPO to PP decreased the pHRR by 21% and 27% for 10 and 15% of SEDOPO, respectively. Similarly, a 21% and 25% decline in pHRR was observed for 10 and 15% loadings of SEDPPO in PP, respectively. Furthermore, for 10% loadings of SEDPP in the PP matrix, pHRR (1070 W g⁻¹) values decreased by ~4.5%; THR (46.3 kJ g⁻¹) values were higher than the corresponding blank PP data. For 15% loading of SEDPP in PP, a small decrease in the pHRR of 3.4% and THR around 2.9% was observed. Blank PP and the PP-SEDPP blends were characterized by a one-step thermal decomposition in MCC data. In contrast, a shoulder noticeable for the PP-SEDOPO and PP-SEDPPO blends at 3.50 °C to 370 °C can be ascribed to the bridged sulfone derivatives that are released due to the matrix decomposition and volatilization. A similar observation for the same blends was also matrix in the TGA analysis performed under an N₂ environment (Fig. S10).

Table 2: Results om MCC measurements.

Sample Name	$T_{M}[^{\circ}C]$	pHRR [y7.g ⁻¹]	THR [kJ.g ⁻¹]	Residue (wt%)
PP-Blank	484.0 ± 0.4	1120.3 ± 16.4	45.3 ± 0.2	0.42 ± 0.30
PP-10% SEDOPO	483.5 ± 0.6	890.7 ± 7.9	43.2 ± 0.4	0.87 ± 0.11
PP-15% SEDOPO	483.4 ± 1.2	809.4 ± 13.9	42.9 ± 0.4	0.26 ± 0.01
PP-10% SEDPPO	482.8 ± 1.3	884.7 ± 12.3	43.7 ± 0.3	0.15 ± 0.1
PP-15% SEDPPO	480.0 = 1.7	838.0 ± 22.8	43.0 ± 0.2	0.38 ± 0.3
PP-10% SEDPP	4,1.0:13	1070.3 ± 13.8	46.3 ± 2.5	0.32 ± 0.31
PP-15% SEDPP	489.0 : 1.5	1082.0 ± 39.6	44.0 ± 0.3	0.41 ± 0.3

In general, a decrease of the pHRR values corresponds to a drop in the heat produced during the combustion of the flame retarded polymeric material. A reduction in the pHRR and an increase in char formation is often associated with condensed phase activity of the FR additive. In such a case, the FR additive reacts with the polymer matrix and decreases flammable volatiles' formation. However, for PP-SEDOPO, PP-SEDPPO, and PP-SEDPP blends, a decrease in pHRR with no significant increase in the residue was observed. This corresponds well with the gas phase flame inhibition activity of the FR additive [30, 57].

The PP-SEDOPO and SEDPPO blends were characterized by a decrease in THR of approx. 2-3 $k \cdot Jg^{-1}$ relative to the blank PP (Table 2). However, a reduction in the THR is not significant for SEDPP, which might be due to the lower loading and FR efficacy of the SEDPP. A decrease in THR values for the PP blends than blank PP reduces its flame spread and fire load [30].

It is well known that phosphorus content is an important factor in determining the flame retardant efficiency of a system. As seen in Table 1, the P-contents of the 10, 15% SEDOPO (measured P-contents of 1.10, 1.54%), and for the 10, 15% SEDPPO (measured P-contents of 1.09, 1.54%) blends are nearly double compared to the 10, 15% SEDPP blends (measured P-contents of 0.63, 0.90%). The pHRR data of PP-10% SEDOPO and SEDPPO can be compared with the PP-15% SEDPP blends as they have similar P content. The SEDOPO and SEDPPO blends have higher flame retardant efficacy compared to SEDPP at similar P-contents.

3.5. Cone calorimeter test

The cone calorimeter was used to investigate alkyl bidged sulfone derivatives combustion behaviors in the PP matrix. Parameters such as pHRR, THY, 15R, COP, av-SEA, and CO₂ output (CO₂) are summarized in Table 3. Generally, these parameters are used to calculate the fire performance of PP blends. The HRR, particularly the pHRR, represents the point at which fire typically spreads further or ignites neighboring objects; hence, it's care is essential for fire safety. It can be seen from Table 3 that the bridged sulfone/PP blends resulted in a reduction of THR and pHRR; however, TSR and CO production increased, which is typeral of flame retardant polymers (Fig. 3). Thus, it is clear from the results that sulfone bridged phesological flame-retardants undermine the combustion behavior of PP. All PP blends have a higher CO/CO₂ ratio than the blank PP, indicating their possible gas phase flame inhibition activity. The PP-15% SEDOPO blend has the highest CO/CO₂ ratio and the lowest PHRR compared to PP-15% SEDOPO over SEDPP systems. This indicates a higher gas-phase flame inhibition efficacy of the SEDOPO over SEDPP.

Table 3: Results for blank PP and PP-FR (15%) blends obtained from cone calorimeter.

Sample	pHRR (kW/m ²)	THR (MJ/m²)	TSR (m ² /m ²)	SEA (m²/kg)	MARHE [kW/m²]	mean CoP	mean CO ₂ (kg/kg)	CO/CO ₂
PP-Blank	469.5 ± 13.8	62.6 ± 2.0	62.6 ± 20.7	172.86 ± 52	175.3 ± 1.8	0.04 ± 0.01	1.41 ± 0.20	0.03
SEDOPO	241.5 ±2 4.2	40.0 ± 12.8	941.2 ± 16.9	462.15 ± 15	132.9 ± 7.8	0.15 ± 0.02	0.98 ± 0.02	0.15
SEDPPO	336.2 ± 24.5	46.9 ± 3.6	843.3 ± 15.8	422.4 ± 31	131.6 ± 1.9	0.04 ± 0.02	0.68 ± 0.40	0.06

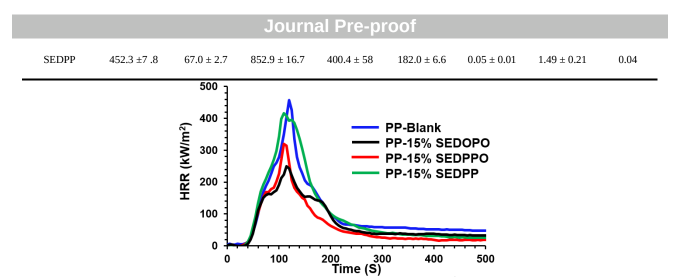


Fig. 3. Heat release rate (HRR) of the blank PP and PP/FRs measured by cone calorimeter.

To investigate the gas phase activity of alkyl bridged sulfone characteristic that combustion efficiency (X) of PP blends was calculated (Table 4). The combustion efficiency can be calculated from the ratio between effective heat of combustion (EHC) mer sund from cone calorimeter and heat of complete combustion (HCC) calculated from MCC (using TrIR and residue content) as described in the literature [58]. As shown in Table 4, the EHC of samples containing SEDOPO and SEDPPO decreased significantly compared to SEDPP and blank PP. The combustion efficiency value of all PP blends was lower than the blank PP, indicating a flame inhibition activity of the FRs [59].

Table 4: Determination of combustion efficiency Y) from cone calorimeter (CC) and MCC data.

Sample	EA'C _{CC} (Kg/g)	$HCC_{MCC}(Kg/g)$	X
PP-Blank	50.3	78.1	0.64
PP-15% SEDOPO	24.2	57.9	0.41
PP-15% 5.7D, PO	36.5	69.3	0.52
PP-15% SED 'P	46.6	74.5	0.62

 EHC_{CC} : effective heat of combustion, HCC_{MCC} : the heat of complete combustion, X: combustion efficiency (ratio between EHC and HCC).

3.6. Thermal Stability

Understanding the thermal behavior of synthesized bridged sulfone compounds may help predict and understand their subsequent thermal processing, thermal, and fire behavior. The melting point (m.p.) of the bridged sulfone compounds were estimated by DSC (Fig. S8 in SI-5). All synthesized compounds have m.p. ranging from 148 to 248 °C and they are thermally stable up to 321 °C as seen in their TGA data (Table 5). Similarly, bridged DOPO compounds have been shown to exhibit m.p

greater than 250 °C [60, 61]. The low m.p. exhibited by compounds synthesized in this work allow for melt processing of their blends. Incorporating the bridged sulfone compounds into PP has negligible influence on melting point (Table 5) of PP. TGA evaluated the thermal decomposition behavior of the synthesized compounds and the PP blends. Their corresponding first order derivative (DTG) curves under both air and nitrogen environments are presented in Fig. S9b, d and f, respectively. The TGA data corresponding to the onset of weight loss (T_{donset}), the maximum decomposition rate (T_{dmax}), and the char yield at 800°C are summarized in Table 5. As shown in Fig. S9, bridged sulfone compounds exhibited high decomposition temperatures (T_{donset} >290°C). Among the synthesized bridged sulfone compounds, SEDOPO (T_{donset} ~32°C) showed the highest thermal stability, followed by SEDPP (T_{donset} ~317°C) and SEDPPO (T_{do set} ~2)8°C) under air atmosphere. The high decomposition temperatures of the compound indicate acreates suitability for the PP extrusion process, which is usually performed below 250 °C. SEDOP and SEDPPO decompose completely, leaving residue less than 2% at 800 °C; thus, likely, these compounds may primarily be active in the gas phase. In contrast, SEDPP yields a slightly higher residue of around 6% under the air atmosphere and 18.5% in the N₂ environment at 800 °C.

Table 5: Thermal properties of Nar', PP, FR additives, and the PP-FR blends.

Samples	T _{donset} [°C]		T _{dmax}	T _{dmax} [°C]		Residue at 800 °C [wt%]		Crystallization
Samples	N_2	Air] 12	Air	N_2	Air	Point [°C]	Point [°C]
PP-Blank	402	24	438	281	0.30	0.61	165	112
SEDOPO	322	^ 21	338	336	4.29	2.67	248	211
PP-10% SEDOPO	412	2.40	445	294	0.59	1.80	165	118
PP-15% SEDOPO	400	205	444	333	0.59	2.21	166	119
SEDPPO	2°+	∠98	309	314	2.20	1.25	202	131
PP-10% SEDPPO	407	250	436	309	1.55	0.30	167	118
PP-15% SEDPPO	576	265	437	314	0.30	1.5	167	115
SEDPP	32	317	341	338	18.5	5.96	148	80
PP-10% SEDPP	418	240	445	288	0.50	1.31	167	116
PP-15% SEDPP	427	246	450	294	0.18	1.23	166	118

 T_{donset} (°C): the temperature of weight loss begins and T_{dmax} (°C): the temperature of the maximum mass loss rate measured as the DTG peak maximum

In general, for all PP blends, higher thermo-oxidative stability was observed compared to blank PP. For PP-SEDOPO, SEDPPO blends, the T_{donset} and T_{dmax} shifted towards higher temperatures in an oxidative environment than blank PP (Table 5). A shoulder is visible in the DTG curve measured under the nitrogen environment for PP-SEDOPO and PP-SEDPPO blends (Fig. S9b, d), indicating the possible early decomposition of SEDOPO and SEDPPO to release volatiles and subsequent decay

of PP. Furthermore, a small shift in the thermal stability of PP in both inert and oxidative environments was noted with the incorporation of SEDPP into the PP matrix (Fig. S9e, f). To further understand the thermo-oxidative stability of the bridged sulfone/PP blends, thermal decomposition kinetics experiments were performed (Fig. 4) and activation energy of decomposition was evaluated. The variation in activation energy (E_a) with the degree of conversion for PP and PP-15% SEDOPO, SEDPPO, and SEDPP blends are presented in Fig. 4. The activation energy required to initiate the decomposition (α =0) of PP-SEDPP is 92 kJ.mol⁻¹and, relatively higher than the energy needed to initiate decomposition of PP-SEDOPO and PP-SEDPPO. At the initial stage of conversion $(\alpha=0.1)$, the activation energies for PP, PP-SEDOPO, PP-SEDPPC and PP-SEDPP are 78, 67, 62, and 62 kJ.mol⁻¹, respectively. For the PP-SEDOPO blend, the activation energy from 80 (α=0) to 109 kJ.mol $^{-1}$ (α =0.8) progresses through different conversion phases. There is a sharp rise in activation energy during the initial stages ($\alpha > 0.1$) of thermal decor, position, representing a decomposition mechanism where SEDOPO strongly interfered in the her nal decomposition process of PP and retard the release of volatiles. Furthermore, the increase in the activation energy of PP-SEDPP at the latter stages of conversion can be due to the limited care formation. This char acts as a thermal barrier and requires higher energy for decomposition. The activation energy for PP, PP-SEDOPO, PP-SEDPPO, and PP-SEDPP at the final stage of accomposition (i.e., α =0.8) are 69, 109, 91, and 112 kJ.mol⁻¹, respectively. Hence, the addca Fk3 in the PP matrix delayed the decomposition of PP. Studies on other FRs in PP have also acmonstrated an increase in activation energy of decomposition for PP [62].

The tertiary proton in PP is liable to the free radical degradation mechanism [63, 64]. As illustrated in Scheme S1, after removing the aertiary hydrogen, the carbon radicals are liable to react with oxygen (O₂), forming peroxide radicals. The subsequent decomposition of the peroxy radicals follows two possible mechanisms. The first possible degradation mechanism is a termination reaction, leading to a terminal aldehyde or unsaturated group in the case of primary and secondary peroxide radicals.

The second possibility is creating PP polymer chains with a reduced molecular weight by rearrangements at the tertiary carbon radical [65]. Thus, it is essential to have antioxidants or additives which can neutralize polymeric radicals and prevent oxidative degradation [66]. During melt processing at high temperatures, the alkyl sulfone bridged phosphorus compounds (SEDOPO, SEDPPO, and SEDPP) decompose to produce phosphorus and sulfone-based radicals. These radicals react with peroxide radicals, protect the PP polymer chain from fire, and delay the thermal-oxidative degradation (Scheme S1).

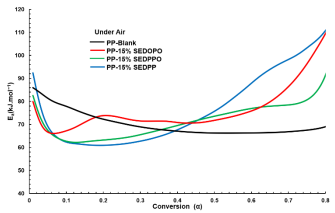


Fig. 4. Variation in activation energy with the degree of conversion for PP blank and PP-FR blends calculated using the Kissinger method.

3.7. TGA-FTIR analysis of evolved gases

To further understand the thermal decomposition behavior of the formulations and postulate a mode of action of the bridged sulfone compounds, evolved gases formed during pyrolysis experiments in TGA were analyzed with TGA-FTIR. The 3D plot of the TGA-FTIR spectra for the blank PP and PP-FR blends are shown in Fig. S11 and the except d FTIR curves of evolved volatiles under different temperatures are presented in Fig. S12. It can be seen in Fig.5 that the maximum absorbance intensities of all selected pyrolysis products of blank PP and FR-PP blends appear close to the decomposition temperature of the polymer matrix. The release of flammable volatiles like alkanes and alkenes is much lower for sulfone bridged/PP matrix, as shown in Fig. 5. Among PP-FR blends, PP-SEDOPO possesses the lowest the mable volatility, followed by the PP-SEDPPO blend. Thus, a reduced amount of flammable s_1 acres can lower the fuel response from underlying material and improve the fire resistance s_1 and s_2 acres can lower the fuel response from underlying material and improve the fire resistance s_2 and s_3 are solved and s_4 acres can lower the fuel response from underlying material and improve the fire resistance s_4 and s_4 are solved and s_4 and s_4 are solved and s_4 are solved and s_4 and s_4 are solved and s_4 and s_4 are solved and s_4 and s_4 are solved and s_4 and s_4 are solved and s_4 are solved and s_4 are solved and s_4 and s_4 are solved an

3.8. DIP-MS and mode of action

There is a limited char formation for the FR-PP blends under N₂ and oxidative conditions compared to the blank PP (Table 5). Bridged sulfone compounds may be active in the gas phase; hence, DIP-MS analysis was performed to evaluate the chemistry of volatiles produced during the thermal decomposition of the FR-PP blends. Fig. S13 shows the total ion chromatogram (TIC) for blank PP and FR-PP blends. SEDPP and SEDPPO blends show the presence of volatile components earlier compared to the blank PP and SEDOPO-blend. Extracted MS- TIC curves of the bridged sulfone-PP blends are presented in Fig. S14, S15, and S16 in the SI part.

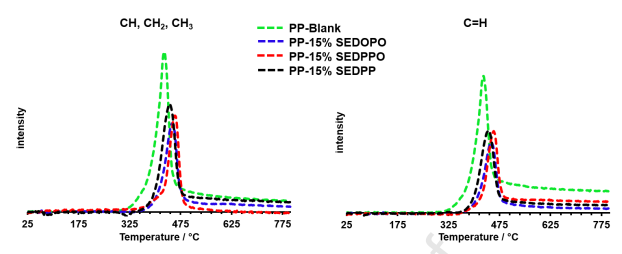
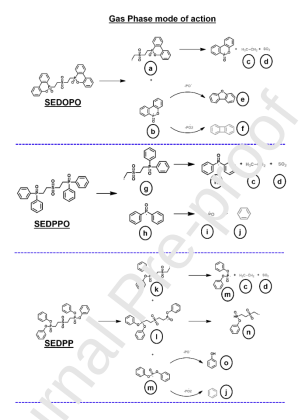


Fig. 5. TGA-FTIR spectra of the total pyrolysis products and intensity of characteristic peaks for selected pyrolysis products of PP blank, PP-15%SEDOPO, SEDPPO, SEDPP blends.

The decomposition mechanisms for SEDOPO, SEDPPC, and SEDPP generally involve elimination and hydrolysis reactions. From the evolved gas analysis by TGA-FTIR and DIP-MS measurements, a simplified decomposition pathway for the FR doctres is proposed in Scheme 2. For the PP-SEDOPO blend, species corresponding to m/z 168, 216, 243, and 335 were observed, representing dibenzofuran, DOPO, ethal DOFO and methylsulfonyl-ethyl DOPO, respectively (Fig. S14). Initially, the decomposition of SEDOPO leads to the formation of fragments a and b (Scheme 2) and subsequently, the accomposition of DOPO further leads to the formation of stable intermediates such as dibunzoiuran (e) and biphenyl (f) along with the production of PO and PO'₂ active species. Further necomposition of the ethylsulfonyl-ethyl DOPO (a) releases the sulfone containing alkare, vhic I further undergoes C-S bond cleavage and subsequent elimination of alkyl radicals (c) and SO₂ (c). Similarly, for SEDPPO blends, the identified fragments in DIP-MS are summarized in Fig. S15. The PP-SEDPPO thermal decompositions initially provide species such as (2-(ethylsulfonyl)ethyl)diphenylphosphine oxide (g) and diphenylphosphine oxide (h) and compound h undergoes subsequent cleavage to produce active PO' and benzene. The (2-(ethylsulfonyl)ethyl)diphenylphosphine oxide (g) further leads to the elimination of alkyl radicals (c) and sulfone (d), scheme 2. In case of SEDPP blends, cis-elimination leads to the production of diphenyl (2-(ethylsulfonyl)ethyl)phosphonate (k), (2-((2-(oxo(phenoxy)-14phosphaneyl)ethyl)sulfonyl)ethyl)phosphonate (l) along with diphenyl phosphonate (m) products. The compound I further decomposes to species n and diphenyl phosphonate decomposes to form phenol (o) and benzene (j), as shown in scheme 2. The decomposition of SO_2 containing compounds

is already known in the literature and the proposed scheme is in good agreement with the published data[57]. Based on these observations, a simplistic decomposition pathway for the bridged sulfone compounds and their possible mode of action is proposed in Scheme 2. As highlighted, the production of PO and SO₂ species for the alkyl sulfone bridged phosphorus FRs boosts their flame inhibition action.



Scheme 2: Proposed thermal decorposition pathways for FRs additives based on DIP-MS analysis. Note: Structures depicted in grey are unidentified species in DIP-MS.

Combining accumulated data acquired from TGA, MCC, and vertical fire test (BKZ-VB), a predominant gas-phase activity for all bridged compounds is thus predicted. Generally, an FR additive should decompose and evolve active species in the gas phase at a temperature close to the polymer matrix decomposition temperature.

3.9. Mechanical behavior for PP films of optimized blends

Melt extrusion of thin films requires high stability of the additives in a molten state and provides samples with partially oriented chains for mechanical tensile testing. As proof of concept, we extruded films from PP-SEDOPO blends, which showed to be the most promising compounds. Fig. 6

shows the exemplary load-strain curves of the films. The curve of the blank PP depicts a shape that is typical for a partially oriented PP film. A yield point indicates the expected transition from elastic to plastic deformation. Since deformation is irreversible after passing the yield point, the latter represents maximum elastic deformation (yield strain) and respective (yield) stress, described in Table 6.

Table 6: Tensile tests result of PP blank and PP-10, 15% SEDOPO films

Sample Type	Sample name	Nominal melt-pump through-put [cm³/min]	Yie, ¹ stress	Yield strain [%]
	PP-Blank	33.6	1.9±2.0	18.4±3.4
Films	PP-10% SEDOPO	33.6	28.2±0.9	11.5±1.1
	PP-15% SEDOPO	33.6	36.6±1.9	9.8 ± 1.0

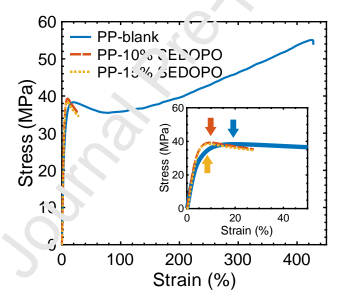


Fig. 6. Stress-strain curves of the PP blank and PP-SEDOPO films of optimized blends. The arrows in the inset plot indicate the yield points.

To assess the additive influence on the elastic deformation of PP film, yield stress and yield strain are compared (Table 6). Field stress slightly increased when FR was added, while yield strain was considerably reduced, indicating increased stiffness. On addition of SEDOPO to PP (10 and 15%), the plastic deformation of PP film is hindered as fillers can constrain polymer chain mobility [67]. As a result, the film breaks after having passed the yield point, which means that the extruded film

ductility is strongly afflicted by FR addition. Similar effects are also reported for other DOPO derivatives in mechanical flexibility tests[68].

4. Conclusions

In this study, a simplified synthesis procedure for three new alkyl sulfone bridged phosphorus compounds was developed, which has the potential to be upscaled in industry. The bridged sulfone compounds were incorporated in the PP matrix via the melt extrusion process and formed into plates and films. The rheological measurements of the FR/PP blends show that a higher viscosity was achieved for the blends; hence, the degradation mechanism during processing becomes less severe with the bridged sulfone compounds. Fire test results of the PP blends at 10% (around 1% phosphorus content) FR indicated that SEDOPO and SEDPPO have superior flame retardant (FR) performance compared to SEDPP. MCC measurements and cone calorimetry measurements demonstrated the flame retardant effectiveness of PP-SEDOPO and PP-SEDOPO blends as lower heat release rates were observed. Among all PP blends the lowest peak heat release of 27% and 49% (reduction compared to blank PP) in MCC and come calorimetry experiments were noted for the 15% PP-SEDOPO blend. The thermal Lecomposition kinetics experiments respectively were performed and the bridged sulfone derivatives how higher activation energy for the decomposition compared to the blank PP at higher conversion, thus further confirming the stabilizing effect of the bridged sulfone compounds under oxicative conditions. The TGA-FTIR and DIP-MS analysis confirmed that the alkyl sulfone briaged phosphorus derivatives are primarily active in the gas phase via SO₂ and PO species formation. To expand the scope of application of such bridged sulfone compounds, fiber spinning and and detailed mechanical analysis are planned in future work. In particular, the reduction of the content with the addition of a suitable synergist will be explored to reduce the adverse effect or high filler loadings on mechanical performance. Future experiments will also address the improvement of dispersion of bridged sulfone derivatives in PP matrix by use of suitable compatibilizers.

Supporting Information. Synthesis procedure for the optimization of the SEDOPO and NMR characterization of bridged sulfone compounds. The TGA and DTG curves in a nitrogen atmosphere, TGA-FTIR 3D spectra for SEDOPO, DIP-MS data- total ion chromatogram, and DIP-MS data for bridged sulfone compounds.

Notes: The authors declare no conflicting interests.

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References

- [1] S.A. Arvidson, K.C. Wong, R.E. Gorga, S.A. Khan, Structure, molecular orientation, and resultant mechanical properties in core/ sheath poly(lactic acid)/polypropylene composites, Polymer 53(3) (2012) 791-800.
- [2] H. Palza, R. Vergara, P. Zapata, Composites of polypropylene melt blended with synthesized silica nanoparticles, Composites Science and Technology 71(4) (2011) 535-540.
- [3] R. Nazir, S. Gaan, Recent developments in P(O/S)–N cortaining flame retardants, Journal of Applied Polymer Science 137(1) (2020) 47910.
- [4] B. Li, M. Xu, Effect of a novel charring-foaming agent on flame retardancy and thermal degradation of intumescent flame retardant polypropyler; Polymer Degradation and Stability 91(6) (2006) 1380-1386.
- [5] A.F. Grand, Effect of Experimental Conditions On the Evolution of Combustion Products Using a Modified University of Pittsburgh Toxicity Test Approarus, Journal of Fire Sciences 3(4) (1985) 280-304.
- [6] M.M. Hirschler, Fire Hazard and Toxic I stericy of the Smoke from Burning Materials, Journal of Fire Sciences 5(5) (1987) 289-307.
- [7] R. Nazir, D. Parida, J. Borgstädt, S. Lehner, M. Jovic, D. Rentsch, E. Bülbül, A. Huch, S. Altenried, Q. Ren, P. Rupper, S. Annaler. S. Gaan, In-situ phosphine oxide physical networks: A facile strategy to achieve durable flame restraint and antimicrobial treatments of cellulose, Chemical Engineering Journal (2020) 128028.
- [8] S. Zhang, A.R. Horrocks, A review of flame retardant polypropylene fibres, Progress in Polymer Science 28(11) (2003) 1517-1538.
- [9] J. Wang, L. Wang, A. Yiao, Lecent Research Progress on the Flame-Retardant Mechanism of Halogen-Free Flame Retardant Folypropylene, Polym.-Plast. Technol. Eng. 48(3) (2009) 297-302.
- [10] P.J. Gale, Mass spectrometric studies of the reactivity of antimony trioxide with halogenated flame retardants in the processes of plastics, International Journal of Mass Spectrometry and Ion Processes 100 (1990) 313-322.
- [11] M. Alaee, P. Arias, A. Sjödin, Å. Bergman, An overview of commercially used brominated flame retardants, their applications, their use patterns in different countries/regions and possible modes of release, Environment International 29(6) (2003) 683-689.
- [12] Z.-Y. Wang, Y. Liu, Q. Wang, Flame retardant polyoxymethylene with aluminium hydroxide/melamine/novolac resin synergistic system, Polymer Degradation and Stability 95(6) (2010) 945-954.
- [13] A.A. Sener, E. Demirhan, The investigation of using magnesium hydroxide as a flame retardant in the cable insulation material by cross-linked polyethylene, Materials & Design 29(7) (2008) 1376-1379.
- [14] T. Xu, X. Huang, Y. Zhao, Investigation into the properties of asphalt mixtures containing magnesium hydroxide flame retardant, Fire Safety Journal 46(6) (2011) 330-334.

- [15] A.u.R. Shah, D.-w. Lee, Y.-q. Wang, A. Wasy, K.C. Ham, K. Jayaraman, B.-S. Kim, J.-I. Song, Effect of concentration of ATH on mechanical properties of polypropylene/aluminium trihydrate (PP/ATH) composite, Transactions of Nonferrous Metals Society of China 24 (2014) s81-s89.
- [16] R.C. Nicolas, C.-E. Wilén, M. Roth, R. Pfaendner, R.E. King III, Azoalkanes: A Novel Class of Flame Retardants, Macromolecular Rapid Communications 27(12) (2006) 976-981.
- [17] M. Aubert, M. Roth, R. Pfaendner, C.-E. Wilén, Azoalkanes: A Novel Class of Additives for Cross-Linking and Controlled Degradation of Polyolefins, Macromolecular Materials and Engineering 292(6) (2007) 707-714.
- [18] M. Aubert, C.-E. Wilén, R. Pfaendner, S. Kniesel, H. Hoppe, M. Roth, Bis(1-propyloxy-2,2,6,6-tetramethylpiperidin-4-yl)-diazene An innovative multifunctional radical generator providing flame retardancy to polypropylene even after extended artificial weathering, Polymer Degradation and Stability 96(3) (2011) 328-333.
- [19] D.C.O. Marney, L.J. Russell, T.M. Stark, The influence of an N-alkoxy HALS on the decomposition of a brominated fire retardant in polypropylene, Polymer Degradation and Stability 93(3) (2008) 714-722.
- [20] K. Cao, S.-l. Wu, S.-l. Qiu, Y. Li, Z. Yao, Synthesis of N-A'koxy Hindered Amine Containing Silane as a Multifunctional Flame Retardant Synergist and As Application in Intumescent Flame Retardant Polypropylene, Industrial & Engineering Chemistry Pesearch 52(1) (2013) 309-317.
- [21] R. Pfaendner, How will additives shape the future of plastics?, Polymer Degradation and Stability 91(9) (2006) 2249-2256.
- [22] R. Pfaendner, Nitroxyl radicals and nitroxyl a ers beyond stabilization: radical generators for efficient polymer modification, Comptes Rend is Chinie 9(11) (2006) 1338-1344.
- [23] S.M.A. Douglas Wayne Horsey, Leong a Harris Davis, Darrell David Dyas, Jr., Robert Leo Gray, Anunay Gupta, Bruce Vincent Hein, Joseph Stephen Puglisi, Ramanathan Ravichandran, Paul Shields, Rangarajan Srinivasan, Hindered amine Same retardant compositions., 1999.
- [24] M. Roth, Flame retardant compositions comprising sterically hindered amines., 2009.
- [25] D.A. Abdel-Ilah Basbas, Robert Con'ava, Michael Peter Difazio, Walter Fischer, Joseph A. Kotrola, Tiziano Nocentini, James Robeins, Kai-Uwe Schöning, Process for the preparation of sterically hindered nitroxyl ethers, 2018.
- [26] S. Zhang, A.R. Horrocks, K. Hull, B.K. Kandola, Flammability, degradation and structural characterization of fibre-forming polypropylene containing nanoclay–flame retardant combinations, Polymer Degradation and Statiluy 91(4) (2006) 719-725.
- [27] S.V. Levchik, E.D. Weil, Overview of recent developments in the flame retardancy of polycarbonates, Polymer International 54(7) (2005) 981-998.
- [28] W. Zhao, B. Li, M. Xu, L. Zhang, F. Liu, L. Guan, Synthesis of a novel flame retardant containing phosphorus and sulfur and its application in polycarbonate, Polymer Engineering & Science 52(11) (2012) 2327-2335.
- [29] C. Luo, J. Zuo, F. Wang, Y. Yuan, F. Lin, H. Huang, J. Zhao, High refractive index and flame retardancy of epoxy thermoset cured by tris (2- mercaptoethyl) phosphate, Polymer Degradation and Stability 129 (2016) 7-11.
- [30] U. Braun, U. Knoll, B. Schartel, T. Hoffmann, D. Pospiech, J. Artner, M. Ciesielski, M. Döring, R. Perez-Graterol, J.K.W. Sandler, V. Altstädt, Novel Phosphorus-Containing Poly(ether sulfone)s and Their Blends with an Epoxy Resin: Thermal Decomposition and Fire Retardancy, Macromolecular Chemistry and Physics 207(16) (2006) 1501-1514.
- [31] J. Wagner, P. Deglmann, S. Fuchs, M. Ciesielski, C.A. Fleckenstein, M. Döring, A flame retardant synergism of organic disulfides and phosphorous compounds, Polymer Degradation and Stability 129 (2016) 63-76.

- [32] W. Pawelec, A. Holappa, T. Tirri, M. Aubert, H. Hoppe, R. Pfaendner, C.-E. Wilén, Disulfides Effective radical generators for flame retardancy of polypropylene, Polymer Degradation and Stability 110 (2014) 447-456.
- [33] A. Ballistreri, G. Montaudo, E. Scamporrino, C. Puglisi, D. Vitalini, S. Cucinella, Intumescent flame retardants for polymers. IV. The polycarbonate—aromatic sulfonates system, Journal of Polymer Science Part A: Polymer Chemistry 26(8) (1988) 2113-2127.
- [34] S. Hou, Y.J. Zhang, P. Jiang, Phosphonium sulfonates as flame retardants for polycarbonate, Polymer Degradation and Stability 130 (2016) 165-172.
- [35] A. Nodera, T. Kanai, Thermal decomposition behavior and flame retardancy of polycarbonate containing organic metal salts: Effect of salt composition, Journal of Applied Polymer Science 94(5) (2004) 2131-2139.
- [36] J. Green, Mechanisms for Flame Retardancy and Smoke suppression -A Review, Journal of Fire Sciences 14(6) (1996) 426-442.
- [37] M. Lewin, J. Zhang, E. Pearce, J. Gilman, Flammability of polyamide 6 using the sulfamate system and organo-layered silicate, Polymers for Advanced Technolog es 18(9) (2007) 737-745.
- [38] M. Lewin, J. Brozek, M.M. Martens, The system polyamide surfamate/dipentaerythritol: flame retardancy and chemical reactions, Polymers for Advanced Technologies 13(10-12) (2002) 1091-1102.
- [39] M. Lewin, Flame Retarding of Polymers with Su¹ am tes. I. Sulfation of Cotton and Wool, Journal of Fire Sciences 15(4) (1997) 263-276.
- [40] Y. Chen, H. Peng, J. Li, Z. Xia, H. Tan, A rovel flame retardant containing phosphorus, nitrogen, and sulfur, Journal of Thermal Analysis and Coorimetry 115(2) (2014) 1639-1649.
- [41] J. He, G. Cai, C.A. Wilkie, The effects of everal sulfonates on thermal and fire retardant properties of poly(methyl methacrylate) and polystyrene, Polymers for Advanced Technologies 25(2) (2014) 160-167.
- [42] C.L. Rasmussen, P. Glarborg, P. Marsi all, Mechanisms of radical removal by SO2, Proceedings of the Combustion Institute 31(1) (2007) 33-347.
- [43] M.R. Zachariah, O.I. Smith, Experimental and numerical studies of sulfur chemistry in H2/O2/SO2 flames, Combustion and Flame 69(2) (1987) 125-139.
- [44] B.A. Howell, Y.G. Daniel, The impact of sulfur oxidation level on flame retardancy, Journal of Fire Sciences 36(6) (2018) 518-53-
- [45] R.A. Stockland, R.I. Ta, To, L.E. Thompson, P.B. Patel, Microwave-Assisted Regioselective Addition of P(O)—H Band, to Alkenes without Added Solvent or Catalyst, Organic Letters 7(5) (2005) 851-853.
- [46] K.A. Salmeia, A. Gomeie, P. Simonetti, R. Nazir, J.-P. Kaiser, A. Rippl, C. Hirsch, S. Lehner, P. Rupper, R. Hufenus, S. Gaan, Comprehensive study on flame retardant polyesters from phosphorus additives, Polymer Degradation and Stability 155 (2018) 22-34.
- [47] S. Vyazovkin, K. Chrissafis, M.L. Di Lorenzo, N. Koga, M. Pijolat, B. Roduit, N. Sbirrazzuoli, J.J. Suñol, ICTAC Kinetics Committee recommendations for collecting experimental thermal analysis data for kinetic computations, Thermochimica Acta 590 (2014) 1-23.
- [48] Y. Saga, D. Han, S.-i. Kawaguchi, A. Ogawa, L.-B. Han, A salt-free synthesis of 1,2-bisphosphorylethanes via an efficient PMe3-catalyzed addition of >P(O)H to vinylphosphoryl compounds, Tetrahedron Letters 56(38) (2015) 5303-5305.
- [49] T. Hirai, L.-B. Han, Air-Induced anti-Markovnikov Addition of Secondary Phosphine Oxides and H-Phosphinates to Alkenes, Organic Letters 9(1) (2007) 53-55.
- [50] L.L. Khemchyan, J.V. Ivanova, S.S. Zalesskiy, V.P. Ananikov, I.P. Beletskaya, Z.A. Starikova, Unprecedented Control of Selectivity in Nickel-Catalyzed Hydrophosphorylation of Alkynes:

- Efficient Route to Mono- and Bisphosphonates, Advanced Synthesis & Catalysis 356(4) (2014) 771-780.
- [51] L.-B. Han, C.-Q. Zhao, Stereospecific Addition of H–P Bond to Alkenes: A Simple Method for the Preparation of (RP)-Phenylphosphinates, The Journal of Organic Chemistry 70(24) (2005) 10121-10123.
- [52] Y. Lin, D. Bernardi, E. Doris, F. Taran, Phosphine-Catalyzed Synthesis of Unsymmetrical 1,3-Bis- and Trisphosphorus Ligands, Synlett 2009(09) (2009) 1466-1470.
- [53] A. Gooneie, P. Simonetti, K.A. Salmeia, S. Gaan, R. Hufenus, M.P. Heuberger, Enhanced PET processing with organophosphorus additive: Flame retardant products with added-value for recycling, Polymer Degradation and Stability 160 (2019) 218-228.
- [54] P. Simonetti, R. Nazir, A. Gooneie, S. Lehner, M. Jovic, K.A. Salmeia, R. Hufenus, A. Rippl, J.-P. Kaiser, C. Hirsch, B. Rubi, S. Gaan, Michael addition in reactive extrusion: A facile sustainable route to developing phosphorus based flame retardant materials, Composites Part B: Engineering 178 (2019) 107470.
- [55] S. Liang, M. Neisius, H. Mispreuve, R. Naescher, S. Gan, Tame retardancy and thermal decomposition of flexible polyurethane foams: Structural influence of organophosphorus compounds, Polymer Degradation and Stability 97(11) (2012) 2428-2440.
- [56] M.E. Üreyen, E. Kaynak, G. Yüksel, Flame-retardant et exists of cyclic phosphonate with HALS and fumed silica in polypropylene, Journal of Applied Pc¹vme: Science 137(4) (2020) 48308.
- [57] A. Battig, J.C. Markwart, F.R. Wurm, B. Schartel, Sun r's role in the flame retardancy of thioether-linked hyperbranched polyphosphoesters in er oxy resins, European Polymer Journal 122 (2020) 109390.
- [58] J.A.P. Solorzano, K.A.M. Moinuddin. S. Tatsiakova-McNally, P. Joseph, A Study of the Thermal Degradation and Combustion Characteristics of Some Materials Commonly Used in the Construction Sector, Polymers (Basel) 11(11) (2019).
- [59] A. Przystas, M. Jovic, K.A. Salmeia, D. Rentsch, L. Ferry, H. Mispreuve, H. Perler, S. Gaan, Some Key Factors Influencing the Figure Retardancy of EDA-DOPO Containing Flexible Polyurethane Foams, Polymers 10(10) (2018).
- [60] A. Buczko, T. Stelzig, L. Bonner, D. Rentsch, M. Heneczkowski, S. Gaan, Bridged DOPO derivatives as flame retardants for NA6, Polymer Degradation and Stability 107 (2014) 158-165.
- [61] I. Butnaru, P.M. Fernández-Ponco, J. Czech-Polak, M. Heneczkowski, M. Bruma, S. Gaan, Effect of Meltable Triazing DopO Additive on Rheological, Mechanical, and Flammability Properties of PA6, Polymer. 7(5) (2015).
- [62] Q. Dong, Y. Ding, B. Ven, F. Wang, H. Dong, S. Zhang, T. Wang, M. Yang, Improvement of thermal stability of polypropylene using DOPO-immobilized silica nanoparticles, Colloid and Polymer Science 290(14) (2012) 1371-1380.
- [63] A.C. Kolbert, J.G. Didier, L. Xu, Mechanochemical Degradation of Ethylene–Propylene Copolymers: Characterization of Olefin Chain Ends, Macromolecules 29(27) (1996) 8591-8598.
- [64] D.M. Mowery, R.L. Clough, R.A. Assink, Identification of Oxidation Products in Selectively Labeled Polypropylene with Solid-State 13C NMR Techniques, Macromolecules 40(10) (2007) 3615-3623.
- [65] H. Hinsken, S. Moss, J.-R. Pauquet, H. Zweifel, Degradation of polyolefins during melt processing, Polymer Degradation and Stability 34(1) (1991) 279-293.
- [66] A.V. Tobolsky, P.M. Norling, N.H. Frick, H. Yu, On the Mechanism of Autoxidation of Three Vinyl Polymers: Polypropylene, Ethylene-Propylene Rubber, and Poly(ethyl acrylate), Journal of the American Chemical Society 86(19) (1964) 3925-3930.

- [67] A. Gooneie, R. Hufenus, Hybrid Carbon Nanoparticles in Polymer Matrix for Efficient Connected Networks: Self-Assembly and Continuous Pathways, Macromolecules 51(10) (2018) 3547-3562.
- [68] A. Gooneie, P. Simonetti, P. Rupper, R. Nazir, M. Jovic, S. Gaan, M.P. Heuberger, R. Hufenus, Stabilizing effects of novel phosphorus flame retardant on PET for high-temperature applications, Materials Letters 276 (2020) 128225.

Rashid Nazir: conceived, planned and carried out the experiments and took the lead in writing the manuscript.

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Conflicts of Interest

The authors declare no conflict of interest.

Highlights

- Three meltable alkyl sulfone bridged phosphorus flame retardants were synthesized in a straightforward approach.
- They have thermal stability > 250 °C and melt processable with polypropylene.
- The incorporation of FRs in PP achieves a significant reduction in heat release rate (~48%).
- Alkyl sulfone bridged phosphorus compounds delay the thermal-oxidative degradation and prevent chain crosslinking of polypropylene.
- At low phosphorus loadings (~1% P content), polyptopylone successfully passed the vertical fire test with increased LOI values (~31%).