Intumescent polypropylene in extreme fire conditions 1 Serge Bourbigot^{a*}, Johan Sarazin^a, Tsilla Bensabath^a 2 ^aUniv. Lille, ENSCL, UMR 8207 - UMET - Unité Matériaux et 3 Transformations, France, serge.bourbigot@ensc-lille.fr 4 5 *Corresponding author 6 7 **Highlights:** Intumescent polypropylene (PP) can resist to burnthrough test 8 • Long time of piercing using zinc borate as synergist 9 • 10 • 11 Abstract: 12 The paper deals with intumescent polypropylene (PP) undergoing extreme 13 fire (burn-through test with heat flux higher than 100 kW/m²). The purpose 14 of this unusual approach is to explore the possibility to design intumescent 15 plastic (here PP) resisting to burn-through test. A combination of 16 commercial intumescent flame retardants (ammonium polyphosphate-based 17 compounds containing a char former; AP766 (AP) and FlameOff (FO) of 18 the companies Clariant and FlameOff Inc) with zinc borate (supplied by US 19 Borax, ZB) or Kemgard (combination of ZB and molybdate supplied by 20 21

- Huber, KZ) was incorporated in PP. Use of ZB and KZ as synergists in FO formulations increases dramatically the time of piercing (formation of hole 22 through the plaque of polymer at 80 s without ZB or KZ vs. 280 s with KZ) 23 at the burn-through test (heat flux = 116 kW/m^2 , propane burner) while the 24 combination with AP does not show any benefit. Analyses of the residues 25 obtained at different times of combustion by solid state nuclear magnetic 26 resonance (NMR) of ³¹P, ¹¹B and ¹³C shows the formation of 27 borophosphates creating a glass reinforcing the intumescent char: it acts as 28 a 'glue' providing flexibility and cohesion to the char. 29
- 30

Keywords: intumescence, fire chemistry, burn-through, polypropylene 31

32

1. Introduction 33

- There is not a commonly accepted definition of the concept of 'extreme fire'. 34
- According to its common use the expression 'extreme fire' is a complex 35

Borophosphates forming a glass reinforcing the intumescent char

entity as it involves different realities and ways of looking. It contains at 36 least three concepts [1]: (i) an idea of extension in the sense that 'extreme 37 fire behavior' is very commonly associated to very large fires or fires that 38 extend in large areas during extended periods of time; (ii) an idea of intensity 39 in the sense that some properties of fire spread, namely its rate of spread or 40 its rate of energy release acquire very large values; (iii) a third idea that is 41 associated to 'extreme fire behavior' is related to rapid change in fire 42 behavior conditions that is also linked to some degree of uncertainty in its 43 prediction and danger. In this paper, we only kept the idea of high rate of 44 45 energy.

It is common practice by fire scientists to quantify the intensity of a fire by 46 the radiant heat flux rather than flame temperature [2]. There is an 47 approximate relationship between fire type and heat flux but we should 48 49 recognize it is a crude assumption because the incident heat fluxes depend 50 on the spatial arrangement between the flame and the receiving target. The examples give a direct measurement of the heat flux from the source and 51 they are: (i) small smoldering fire: 2-10 kW/m²; (ii) trash can fire: 10-50 52 kW/m²; (iii) room fire, open pool fire: 50-100 kW/m²; (iv) post-flashover 53 room fire, confined pool fire: $>100 \text{ kW/m^2}$, (v) jet fuel fire, open jetfire: 100-54 200 kW/m² and (vi) confined jetfire, BLEVE: >200 kW/m². In this research, 55 heat flux corresponding to open jetfire or jet fuel fire was considered (it is 56 high energy and hence it can be considered as extreme fire). Higher heat 57 fluxes were not selected because the purpose was to investigate the response 58 of an intumescent polypropylene (PP) in unconventional conditions for fire 59 testing. 60

The intumescence process results from a combination of charring and 61 foaming at the surface of the substrate [3]. The result of this process is the 62 formation of a multicellular (alveolar) barrier, thick and non-flammable, 63 which protects the substrate or residual material from heat or flame action. 64 The charred layer acts as a physical barrier which slows down heat and mass 65 transfer between gas and condensed phase. The formation of an intumescent 66 char is a complicated process involving several critical aspects: rheology 67 (expansion phase, viscoelasticity of char), chemistry (charring) and 68 thermophysics (limitation of heat and mass transfer) [4]. This concept of 69 intumescence enables to make flame retarded (FR) polymeric materials 70 (including PP-based materials) exhibiting high performance in the case of 71 reaction to fire (contribution of the material to fire growth) [5]. FR PPs are 72 not used in the case of resistance of fire (ability of materials to resist the 73 passage of fire and/or gaseous products of combustion), i.e. fire scenarios 74 corresponding to burn-through, jetfire or structural response to fire, because 75 they are not designed for this: they soften upon heating and fire can spread 76

out. Nevertheless, intumescent coatings applied on steel or composite act as efficient fire barrier and we believe we should reach acceptable performance using intumescent thermoplastics in the case of resistance of fire. It is an unconventional testing for thermoplastics and no paper reports this type of approach. At this time, there is no specific application for this type of materials and the motivation of this paper is to explore the possibility to design intumescent plastic (here PP) resisting to burn-through test.

The paper is organized in three parts. The first part is devoted to the design of the intumescent formulation in PP considering potential synergists. Based on the results of the first part, the second part deals with the determination of mechanism of action using specific analyses by solid state nuclear magnetic resonance (NMR). Finally, the effect of the fillers content (loading) determined in the first part and incorporated in PP is examined in the third part.

91

92 2. Experimental

93 2.1. Materials

Commercial grade of PP was used in this work: PP (ISPLEN PP 089 Y1E) 94 was supplied by Repsol (Madrid, Spain). PP 089 Y1E has a melt flow rate 95 (MFR) for a load of 2.16 g at 230°C of 31 g/10 min. Modified ammonium 96 polyphosphate (AP) is the commercial grade of Clariant (Knapsak, 97 Germany) with the brand name Exolit AP766. It is an intrinsic intumescent 98 system containing 24 wt% phosphorus and 15 wt% nitrogen acting in 99 synergy. Another modified ammonium polyphosphate as a powder was used 100 from the company FlameOff (Raleigh, NC – USA) and hereafter called FO 101 102 (the composition is proprietary). Zinc borate (ZB) was added in combination with the intumescent additives as potential synergist. Its composition is 103 2ZnO3B₂O₃ and it was supplied by the company US Borax (Boron, CA -104 USA) under the brand name Firebrake. Kemgard 700Z is a complex mixture 105 of zinc molybdate/zinc borate (the composition is proprietary) from the 106 company Huber Engineered Materials (Atlanta, GA - USA): it was also used 107 in combination with the intumescent additives as potential synergist and it 108 109 is hereafter called KZ.

110

111 **2.2. Processing and formulations**

The strategy was to blend PP with FRs in a twin-screw extruder. The total loading of FRs in PP was between 30 and 50 wt% varying the ratio between

the intumescent additives (AP or FO) and the synergists (ZB or KZ). The

formulations prepared in this work are gathered in Table 1. Note the loading
of ZB and KZ is low compared to AP and FO to get the highest
reinforcement of the intumescent char [6, 7].

Table 1. Formulations of the intumescent PPs

Formulation	PP (wt%)	AP (wt%)	FO (wt%)	ZB (wt%)	KZ (wt%)
РР	100	0	0	0	0
PP(70)-AP	70	30	0	0	0
PP(70)-	70	28	0	2	0
AP/ZB(28:2)					
PP(70)-FO	70	0	30	0	0
PP(70)-	70	0	29	1	0
FO/ZB(29:1)					
PP(70)-	70	0	28	2	0
FO/ZB(28:2)					
PP(70)-	70	0	27	3	0
FO/ZB(27:3)					
PP(70)-	70	0	25	5	0
FO/ZB(25:5)					
PP(70)-	70	0	29	0	1
FO/KZ(29:1)	70	0	28	0	2
PP(70)-FO/KZ					
(20.2)					
$\Gamma\Gamma(70)-\GammaO/KZ$	70	0	27	0	3
(27.3) PP (70)- FO / K 7					
(25:5)	70	0	25	0	5
PP(50)-		_	10.0		
FO/ZB(29:1)	50	0	48.3	1.7	0
PP(50)-	50	0	46.7	3.3	0
FO/ZB(28:2)					
PP(50)-	50	0	45	5	0
FO/ZB(27:3)					
PP(50)-	50	0	41.7	8.3	0
FO/ZB(25:5)					
PP(50)-	50	0	48.3	0	1.7
FO/KZ(29:1)					
PP(50)-FO/KZ	50	0	46.7	0	3.3
(28:2)					
PP(50)-FO/KZ	50	0	45	0	5
(27:3)					
PP(50)-FO/KZ	50	0	41.7	0	8.3
(25:5)	-				

Compounding was performed using HAAKE Rheomix OS PTW 16 twinscrew extruder. The extruder is a co-rotating intermeshing twin screw with a barrel length of 400 mm and screw diameter of 16 mm (L/D = 25) with 10 zones. PP and FRs were incorporated using two gravimetric side feeders into the extruder. Polymer flow rate is fixed to extrude about 500 g/h with a screw speed of 300 rpm. The temperature profile of the extruder from feeder

- to die was set at 200/ 200/ 200/ 200/ 200/ 170/ 185/ 180/ 200/ 200°C.
- 129

130 **2.3. Burn-through test**

In a previous work, a versatile fire test was developed with a complete set 131 of instrumentation to investigate the fire behavior of materials. The 132 description of this test is fully described in [8] and the reader could find all 133 information about the setup and the measurement. In this paper, only 134 required information to describe the operation of the test is given. This test 135 is a burn-through test which was designed to mimic the aeronautical fire test 136 defined in the standard ISO2685:1998. In this work, it was used as burn-137 through test and to create an extreme fire for fire-retarded thermoplastics. 138 The equipment is described as follows and is shown in Fig. 1: (i) Propane 139 burner from Bullfinch, (ii) High purity propane supplied by Air Liquide 140 (N35, purity 99.95%), (iii) Propane flowmeter from Bronkhorst High-tech, 141 (iv) Water-cooled heat flux gauge from Sequoia, (v) Cooling thermostat 142 from Lauda Brinkmann (Lauda Proline RP845), (vi) Infrared (IR) camera 143 from FLIR SystemsTM (ThermovisionTM A40M Researcher) calibrated 144 from 0°C to 1000°C and (vii) Fireproof panels composed of silicate of 145 calcium from Final Advanced Materials (Calsil) of 10mm thick. 146

The burner can deliver a propane-air flame characterized by a heat flux up 147 to 200 kW/m². The burner was placed at 75 mm from the material and the 148 heat flux was calibrated at 116 kW/m² using a heat flux gauge in the same 149 conditions as the tested sample in a separate box (error less than 5%). The 150 temperature of the flame was measured with 5 aligned thermocouples (along 151 the flame) at 1100°C. The sample size was 10x10 cm² and was put between 152 the two panels in Calsil (see above). Temperature was measured in the center 153 of the sample using a thermocouple embedded in the polymer at the surface 154 of its backside. Infrared camera was also used to estimate the surface 155 temperature assuming the emissivity of the surface constant equaling 0.92 156 (black paint of known emissivity on the backside of the polymer plaque). 157 Reasonable agreement was observed between the two measurements and 158 only temperature measured by the thermocouple was shown in the 159 following. All experiments were repeated at least twice and all 160

161 measurements were within 10% error (in [8], we showed the error on 162 temperature measurement was less than 10%).

163



164

Fig. 1. Picture (a, c, e) and scheme (b, d, f, g) of the experimental apparatus - (1) Test 165 166 bench frame, (2) Cooling system, (3) Copper coil cooler, (4) Propane thermocontroller, (5) Propane flowmeter, (6) Propane gas line, (7) Burner support, (8) Propane flame 167 burner, (9) Hood, (10) Ring sampler, (11) Sample holder, (12) Infrared (IR) camera, (13) 168 169 Precision scale, (14) Scale holder, (15) Data acquisition device, (16) Bolt, (17) Steel 170 support, (18) Fireproof boards, (19) Sample, (20) Fireproof boards, (21) Washer, (22) Screw, (23) Water-cooled calorimeter holder, (24) Water-cooled calorimeter, (25) 171 172 Fireproof panel, (26) Aligned thermocouples, (27) Thermocouple holder (adapted from 173 [8])

173 174

175 **2.4. NMR analyses**

¹¹B magic angle spinning – nuclear magnetic resonance (MAS-NMR) was performed at 256.6MHz on a 18.8 T Bruker Avance III spectrometer with a 3.2mm probehead operating at a spinning frequency (v_{rot}) of 20 kHz. The spectra were recorded with a 1 ms pulse length (corresponding to a $\pi/12$ flip angle determined on a liquid), a recycle delay (rd) of 10 s and 128 transients. ¹³C and ³¹P MAS-NMR experiments were performed on a 9.4 T Bruker

Avance spectrometer at 100.6 and 162MHz, respectively. The ${}^{13}C({}^{1}H)$ 182 cross-polarization (CP) NMR experiment was conducted with a 4mm 183 probehead at v_{rot} of 10 kHz with a 4 ms pulse length (corresponding to a $\pi/2$ 184 flip angle), a rd of 10 s, a contact time of 1 ms and 1024 transients. The ³¹P 185 NMR analysis was carried out with a 4 mm probehead under 1H decoupling 186 conditions. The spectrum was recorded with a v_{rot} of 12.5 kHz, a 2 µs pulse 187 length (corresponding to a $\pi/4$ flip angle) a rd of 120 s and 16 transients. ¹¹B 188 and ³¹P chemical shifts were referred to NaBH₄ and H₃PO₄ at - 42.06 ppm 189 and 0 ppm, respectively. 190

191

192 **3. Results and discussion**

193 **3.1. Design of the intumescent formulation**

Intumescent PPs are generally designed to exhibit high performance in terms 194 of reaction to fire but not in terms of resistance to fire. In this section, it is 195 our goal to examine the ability of intumescent PPs to resist to burn-through 196 test. This condition is unusual for commodity polymer such as PP but 197 intumescence is also used as protective coating for diverse substrates (e.g. 198 steel, wood or composites) to pass the burn-through test. Two types of 199 additives were incorporated in PP: (i) AP is a conventional intumescent 200 additive providing low flammability to PP for various fire scenarios such as 201 UL-94, glow wire or cone calorimeter and (ii) FO is a new intumescent 202 additive on the market and was made from intumescent ingredients of 203 204 intumescent paint. Based on previous works of this lab, zinc borate was selected as potential synergist because it is known to reinforce intumescent 205 char [6]. ZB is a pure zinc borate and was already used in previous 206 formulations [7] but KZ has never been evaluated in combination with 207 intumescent ingredients. This last product was selected because its main 208 composition is zinc borate and to take advantage of the presence of 209 molybdate (known as synergist with metal hydroxides). 210

Intumescent PPs filled at 30 wt% loading were first evaluated (Fig. 2). The 211 temperature rise in all materials is similar up to 40 s. Virgin PP softens when 212 it reaches 140°C on its backside (melting temperature of PP is in the range 213 214 150-170°C) and it pierces just after (sharp increase of temperature at 150°C). When AP and AP/ZB (ratio 28 to 2) are incorporated in PP, char formation 215 at the surface of the material can be observed but it cannot extend the time 216 to piercing. Visual observation suggests the expansion of the char is not fast 217 enough to provide a protection and it remains too soft to resist to the 218 219 impingement of the flame. In this case, there is no benefit to combine ZB with the intumescent additive. On the contrary, the incorporation of FO 220 slows the temperature rise after 40 s thanks to the formation of an 221

222 intumescent char. The char resists to the impingement of the flame up to 90 s and it pierces after because of the too high softening of the material. The 223 addition of ZB in the formulation dramatically improves the resistance to 224 piercing of the system. The time to piercing is reached at 225 s thanks to the 225 fast formation of an intumescent char which remains rigid (low softening) 226 for longer times. 227

228



229 230

232



Fig. 2. Temperature as a function of time on the backside of intumescent PPs containing AP and FO as main ingredient during a burn-through test at 116 kW/m².

From the results above, it is shown AP is not the right additive for PP to 233 perform the burn-through test. In the following, we only focus our work on 234 FO in combination with ZB and on determining the best ratio FO/ZB. We 235 kept the loading of ZB low compared to FO (Table 1) because our previous 236 work showed the ratio intumescent formulation over ZB should be high 237 enough (typically 28:2) to get the highest efficiency of ZB [7]. Fig. 3 –(a) 238 239 shows the temperature/time curves of the formulations varying the ratio FO/ZB compared to virgin PP and PP(70)-FO. The formulation having the 240 ratio 29:1 pierces at 125 s and so, it does not show any enhancement 241 compared to the ratio 28:2. Note the performance is highly sensitive to the 242 content of ZB: only 1wt% variation of ZB permits to gain 100s before 243 piercing. The two ratios 27:3 and 25:5 enhance the time to piercing at 250 s 244 compared to 225 s for the ratio 28:2. According to visual observation, the 245 addition of ZB permits keeping the rigidity of the intumescent char 246

undergoing the impingement of the flame and preventing too much 247 deformation at longer times (and hence the piercing of the sample). The 248 same experiment was done substituting ZB by KZ with the same ratio (Fig. 249 3 –(b)). The addition of KZ in the formulations PP(70)-FO/KZ permits for 250 all ratios to dramatically enhance the time to piercing compared to the use 251 of FO alone. The behavior looks similar and the longer time to piercing is 252 253 reached at about 300 s for PP(70)-FO/KZ(25:5). The same conclusion as above can be made namely the addition of KZ permits to reinforce the char 254 strength keeping its rigidity. 255





Fig. 3. Temperature as a function of time on the backside during a burn-through test at 116 kW/m² of (a) intumescent PP(70)-FO/ZB and (b) intumescent PP(70)-FO/KZ.

259

260 **3.2. Role of zinc borate**

The previous section showed the role of ZB reinforcing the intumescent 261 Specific chemical reactions should be responsible for char. this 262 reinforcement and were characterized by solid state NMR. This technique is 263 very useful because it permits the selection of the nucleus (here ³¹P, ¹¹B and 264 13 C) and to observe its surrounding. So, it gives the chemical species formed. 265 Four materials were selected based on the results above (best ratio of 266 FO/ZB=27/3): PP(70)-AP, PP(70)-AP/ZB(27:3), PP(70)-FO and PP(70)-267 FO/ZB(27:3). Note ZB was used instead of KZ because the main component 268 of KZ is ZB and we wanted to avoid other additional interactions. They were 269 270 submitted at the burn-through test and the combustion was stopped at characteristic times namely, 50 s (all samples), 100 s (samples containing 271 FO) and 215 s (only PP(70)-FO/ZB(27:3)). The samples before testing were 272 used as reference. 273

Ammonium polyphosphate (APP) is contained in both AP and FO and is 274 275 one of the main ingredients of the intumescent flame retardants (Fig. 4 at t=0 s). Upon heating, APP decomposes and yields acidic phosphates acting 276 as char promoter [3]. They play a significant role in the charring and in the 277 formation of an intumescent coating: the evolution of the phosphate species 278 should be revealed by ³¹P NMR. The four samples were then characterized 279 by DD-MAS ³¹P NMR as a function of burning time (Fig. 4). The number 280 of bridging oxygen atoms allows classifying the phosphate structure using 281 Qⁿ terminology where n represents the number of bridging oxygen atoms 282 283 per phosphorus tetrahedron. This terminology was used in the following [9]. The spectra of the neat materials (t = 0 s) exhibit a doublet located at -22 284 and -24 ppm assigned to Q^2 site. This doublet is characteristic of P in APP 285 as already reported in our previous work [10]. An additional band of low 286 intensity at 1 ppm can be distinguished on the two FO samples. It is assigned 287 to Q^0 site probably an orthophosphate linked to aliphatic species. 288

At t = 50 s, the two spectra of the samples without ZB exhibit three bands 289 but not located at the same chemical shift (Fig. 4 (a) and (b)). The bands at 290 0 and -12 ppm are common for the two materials and they are assigned to 291 Q^0 site (probably mainly phosphoric acid [11]) and to Q^1 site 292 (orthophosphates linked to aromatic species [12]) respectively. It is 293 noteworthy that the amount of species in Q^1 site is higher than that in Q^0 for 294 PP(70)-FO (ratio of the areas Q^{1}/Q^{0} is higher). It suggests FO promotes the 295 formation of phosphate linked to char. The band at -6 ppm (PP(70)-AP) is 296 attributed to Q^2 sites corresponding to pyrophosphates [12] and the broad 297 band centered at -27 ppm (PP(70)-FO) is attributed to the formation of 298 amorphous phosphate-type exhibiting Q^3 and Q^4 sites [13]. With the ZB in 299 the formulation, the two systems have similar spectra (Fig. 4 (c) and (d)). 300 The broad resonance (between 5 and -55 ppm) is the signature of a 301 distributed structure found in glasses or amorphous compounds while the 302 narrow resonances are characteristic of ordered phase. It can contain zinc 303 phosphate and borophosphate in addition to phosphate glass [9, 10]. The two 304 sharp bands at 0 and -6 ppm can be assigned as above. The band centered at 305 -30 ppm is assigned to borophosphate [14] and the broad band centered at -306 12 ppm might be assigned as above in a disordered structure and/or to 307 borophosphate glass [9]. Finally, at higher testing times for PP(70)-308 FO/ZB(27:3), the spectra are similar to those at 50s. Those results evidence 309 310 APP and its decomposition products react with zinc borate. It is not unusual in intumescent systems and it was already reported in previous work [9]. 311 The formation of borophosphate glass reinforces the char and acts as a 'glue' 312 providing flexibility and cohesion to the char. 313

314





316 317

MAS ¹¹B NMR spectra exhibit two bands at t=0s (Fig. 5). The first band 318 lying from 10 to 20 ppm, shows the presence of trigonal (BO₃) borate units 319 while the second band (band centered around 1 ppm) is assigned to 320 tetragonal (BO₄) borate species [9]. Those two polyhedra can be 321 characterized according to their different chemical shifts but also through 322 323 their quadrupolar constant (C_0). C_0 of BO₃ is indeed much larger than that of BO₄ because of the higher asymmetry of the planar BO₃ species (2.4-3.0 324 MHz and <1 MHz for the BO₃ and BO₄ units, respectively) [15]. This 325 explains why BO₃ signals are broader than BO₄ resonances. After burning 326 (t > 0s), the broad band assigned to BO₃ units disappears and the band 327 assigned to BO₄ units is shifted to -4 ppm (Fig. 5). The spectra are similar 328 for the two formulations and whenever the duration of testing. A main sharp 329 band can be distinguished at -4 ppm and a shoulder at -1.5 ppm which are 330 assigned to two types of borophosphates [16]. It is consistent with the 331 assignments of the DD-MAS ³¹P NMR spectra. Borates react with 332

phosphates upon heating and they are all consumed by the reaction: they are

no longer 'free' borates in the intumescent coating.

335



Fig. 5. MAS ¹¹B NMR of intumescent PPs containing ZB as a function of burning time

The basic principle of intumescence is to make a protective char. Visual 337 observation of the behavior of the materials evidences the charring of the 4 338 systems and hence, ¹³C NMR should provide information on carbonaceous 339 species formed upon burning. All CP-DD-MAS ¹³C NMR spectra exhibit 340 three resonance bands located at 21, 26 and 43 ppm (Fig. 6). They can be 341 assigned to polymeric chains of PP where the bands at 21, 26 and 43 ppm 342 are assigned to CH₃, CH and CH₂ groups respectively [17]. For the sake of 343 brevity, the spectra at t = 0 s were not shown because they only exhibit the 344 three mentioned bands and no additional insight was provided. It is 345 noteworthy the bands of PP are detected whenever the duration of testing. It 346 makes sense because the test is stopped when piercing occurs due to the 347 softening of PP. PP is therefore not completely decomposed and so, it can 348 be detected by NMR. 349

Intumescent char is constituted by condensed polyaromatic species 350 containing mainly carbon and sometimes heteroatoms like nitrogen and 351 oxygen [18, 19]. They are then detected by solid state NMR of carbon by a 352 broad band centered around 130 ppm corresponding to sp² hybridized 353 aromatic carbon atoms. Except for the samples containing FO, this band 354 cannot be detected with our experimental conditions (even when zooming 355 in) (Fig. 6). Two phenomena could explain this: (i) CP was used for the 356 acquisition of the spectra and because of the low number of protons on 357 aromatic rings, the magnetization transfer is low and hence, the intensity of 358 the band is low or undetectable and (ii) the formation of carbon free radicals 359 on aromatic ring creates a strong anisotropy of magnetic susceptibility and 360 then the loss of NMR signal [12]. Charring occurs for each sample and when 361 detectable, a broad band centered at 125 ppm can be distinguished. On the 362

spectra recorded on FO containing samples at 50 s (Fig. 6-b) and at 100 s 363 (Fig. 6-d), a tail to the higher ppm is observed. It suggests the formation of 364 oxidized carbons and of aromatic carbons bound to phosphates [20]. 365 Additional bands can be distinguished on the spectrum of PP(70)-366 FO/ZB(27:3) at t = 215 s (Fig. 6-d) located at 111, 121, 125 and 143 ppm. 367 They are relatively sharp on the broad band corresponding to the aromatic 368 369 carbons. The band at 111 ppm is assigned to protonated aromatic carbon, those at 121 and 125 ppm are assigned to non-protonated aromatic carbons 370 and that at 143 ppm is assigned to aromatic carbons bound with phosphates 371 372 [20]. The shape of the 4 bands indicate carbon atoms are in an environment of higher symmetry suggesting the formation of some crystalline species. 373 Overall, it is shown phosphates can be bound to the charred structure. The 374 presence of phosphate prevents the oxidation of 'carbons' [21] and provides 375 additional mechanical properties to the char (higher char strength and 376 flexibility) [22]. 377





379

380

381 **3.3. Effect of loading**

The first section showed that ZB and KZ in combination with FO provide 382 superior performance at the burn-through test. The purpose is to increase the 383 384 total loading of fillers (at 50 wt%) in PP to investigate its effect in terms of time to piercing. The time/temperature curves of all formulations are shown 385 on Fig. 7. Surprisingly, the curves of the formulations containing ZB does 386 not show any improvement compared to those at 30 wt% loading (Fig. 3 vs. 387 Fig. 7). It is also true for the formulations containing KZ (times to piercing 388 are even shorter) except for the ratio FO/KZ at 25 to 5 (Fig. 7-b). In this last 389 case, the time to piercing reaches 330 s compared to the others exhibiting a 390 time to piercing at 175 s. At high loading, it is observed higher charring but 391 at the same time, the material looks softer. So, it implies that the 392 impingement of the flame creates the piercing at shorter times because of its 393 lower viscosity even if its efficiency as heat barrier might be higher. 394



Fig. 7. Temperature as a function time on the backside during a burn-through test at 116
 kW/m² of (a) intumescent PP(50)-FO/ZB and (b) intumescent PP(50)-FO/KZ.

397

398 **4. Conclusion**

This paper showed that intumescent PP could be designed to resist to 399 burnthrough test. The selection of the intumescent system (or in another 400 words the chemistry of the system) is essential to get long time to piercing 401 (e.g. 100 s vs. 40 s when using FO instead of AP) and a synergist can 402 dramatically extend the time to piercing (e.g. 215 s vs. 100 s when using 403 FO/ZB instead of FO alone). Analyzing the residues obtained at different 404 times of combustion by solid state NMR of ³¹P, ¹¹B and ¹³C, it is shown that 405 phosphates were bound to the charred structure (in the case of FO containing 406 systems) and that the formation of borophosphates created a glass 407 reinforcing the intumescent char. This combination provides flexibility and 408 cohesion to the char. The unusual fire resistance of FR plastics paves the 409 way to other applications where burnthrough scenario can be involved. Such 410

scenario might happen in plants if leak of flammable products occurs
creating a torch impinging surrounded plastics. The fire resistance of our
formulations might bring therefore additional fire safety.

414

415 **5. Acknowledgement**

- 416 This work has received funding from the European Research Council (ERC)
- under the European Union's H2020- the framework programme for
 Research and Innovation (2014-2020) ERC Grant Advances Agreement
- ⁴¹⁸ Research and Innovation (2014-2020) ERC Grant Advances Agreemen
 ⁴¹⁹ N°670747-ERC 2014 AdG/FireBar-Concept for FireBar Concept project.
- 420

421 6. References

- 422 [1] Viegas DX. Extreme fire behaviour. In. Extreme fire behaviour. Nova Science
- 423 Publishers, Inc., 2012, pp. 1-56.
- 424 [2] La Delfa G, Luinge JW, Gibson AG, Integrity of composite aircraft fuselage materials
- 425 under crash fire conditions, Plastics, Rubber and Composites, 2009;38: 111-17.
- 426 [3] Alongi J, Han Z, Bourbigot S, Intumescence: Tradition versus novelty. A
- 427 comprehensive review, Progress in Polymer Science, 2015;51: 28-73.
- 428 [4] Jimenez M, Duquesne S, Bourbigot S, Multiscale Experimental Approach for
- 429 Developing High-Performance Intumescent Coatings, Industrial & Engineering Chemistry
 430 Research, 2006;45: 4500-08.
- 431 [5] Bourbigot S, Sarazin J, Bensabath T, Samyn F, Jimenez M, Intumescent
- 432 polypropylene: Reaction to fire and mechanistic aspects, Fire Safety Journal, 2019;105:433 261-69.
- 434 [6] Casetta M, Delaval D, Traisnel M, Bourbigot S, Influence of the recycling process on
- the fire-retardant properties of PP/EPR blends, Macromolecular Materials and
- 436 Engineering, 2011;296: 494-505.
- 437 [7] Fontaine G, Bourbigot S, Duquesne S, Neutralized flame retardant phosphorus agent:
- Facile synthesis, reaction to fire in PP and synergy with zinc borate, Polymer Degradationand Stability, 2008;93: 68-76.
- 440 [8] Tranchard P, Samyn F, Duquesne S, Thomas M, Estèbe B, Montès JL, Bourbigot S,
- 441 Fire behaviour of carbon fibre epoxy composite for aircraft: Novel test bench and
- 442 experimental study, Journal of Fire Sciences, 2015;33: 247-66.
- 443 [9] Hansupo N, Tricot G, Bellayer S, Roussel P, Samyn F, Duquesne S, Jimenez M,
- Hollman M, Catala P, Bourbigot S, Getting a better insight into the chemistry of
- decomposition of complex flame retarded formulation: New insights using solid stateNMR, Polymer Degradation and Stability, 2018;153: 145-54.
- [10] Samyn F, Bourbigot S, Duquesne S, Delobel R, Effect of zinc borate on the thermal
- 447 [10] Saniyi F, Bourbigot S, Duquesne S, Derober K, Effect of Zinc borate on the therman 448 degradation of ammonium polyphosphate, Thermochimica Acta, 2007;456: 134-44.
- 449 [11] Sut A, Greiser S, Jäger C, Schartel B, Synergy in flame-retarded epoxy resin:
- 450 Identification of chemical interactions by solid-state NMR, Journal of Thermal Analysis
- 451 and Calorimetry, 2017;128: 141-53.
- 452 [12] Bourbigot S, Le Bras M, Delobel R, Decressain R, Amoureux JP, Synergistic effect
- 453 of zeolite in an intumescence process: Study of the carbonaceous structures using solid-
- 454 state NMR, Journal of the Chemical Society Faraday Transactions, 1996;92: 149-58.
- 455 [13] Mercier C, Montagne L, Sfihi H, Palavit G, Boivin JC, Legrand AP, Local structure
- 456 of zinc ultraphosphate glasses containing large amount of hydroxyl groups: 31P and 1H

- 457 solid state nuclear magnetic resonance investigation, Journal of Non-Crystalline Solids,
- 458 1998;224: 163-72.
- 459 [14] Jimenez M, Duquesne S, Bourbigot S, Intumescent fire protective coating: Toward a
- better understanding of their mechanism of action, Thermochimica Acta, 2006;449: 16-26.
- 462 [15] Hansen MR, Madsen GKH, Jakobsen HJ, Skibsted J, Refinement of borate structures
- 463 from11B MAS NMR spectroscopy and density functional theory calculations of11B
- 464 electric field gradients, Journal of Physical Chemistry A, 2005;109: 1989-97.
- 465 [16] Tricot G, Raguenet B, Silly G, Ribes M, Pradel A, Eckert H, P-O-B3 linkages in
- borophosphate glasses evidenced by high field 11B/31P correlation NMR, Chemical
 Communications, 2015;51: 9284-86.
- 468 [17] Laupretre F, Bebelman S, Daoust D, Devaux J, Legras R, Costa JL, NMR,
- 469 differential scanning calorimetry, and fourier transform infrared characterization of the
- 470 crystalline degree and crystallite dimensions of ethylene runs in isotactic
- 471 polypropylene/ethylene-propylene copolymer blends (iPP/EP), Journal of Applied
- 472 Polymer Science, 1999;74: 3165-72.
- 473 [18] Bourbigot S, Le Bras M, Delobel R, Breant P, Tremillon J-M, Carbonization
- 474 mechanisms resulting from intumescence part II. Association with an ethylene
- 475 terpolymer and the ammonium polyphosphate-pentaerythritol fire retardant system,
- 476 Carbon, 1995;33: 283-94.
- 477 [19] Bourbigot S, Le Bras M, Delobel R, Gengembre L, XPS study of an intumescent
- 478 coating II. Application to the ammonium polyphosphate/pentaerythritol/ethylenic
- 479 terpolymer fire retardant system with and without synergistic agent, Applied Surface480 Science, 1997;120: 15-29.
- 481 [20] Karrasch A, Wawrzyn E, Schartel B, Jäger C, Solid-state NMR on thermal and fire
- 482 residues of bisphenol A polycarbonate/silicone acrylate rubber/bisphenol A bis(diphenyl-
- 483 phosphate)/(PC/ SiR/BDP) and PC/SiR/BDP/zinc borate (PC/SiR/BDP/ZnB) Part I: PC
- charring and the impact of BDP and ZnB, Polymer Degradation and Stability, 2010;95:
 2525-33.
- 486 [21] McKee DW, Spiro CL, Lamby EJ, The inhibition of graphite oxidation by
- 487 phosphorus additives, Carbon, 1984;22: 285-90.
- 488 [22] Bourbigot S, Le Bras M, Delobel R, Trémillon JM, Synergistic effect of zeolite in an
- 489 intumescence process: Study of the interactions between the polymer and the additives,
- 490 Journal of the Chemical Society Faraday Transactions, 1996;92: 3435-44.
- 491