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Synthesis and characterization of a phosphorous/nitrogen based sol-gel coating as a novel

halogen- and formaldehyde- free flame retardant finishing for cotton fabric

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16	
17	Abstract
18	A novel formaldehyde- and halogen-free coating, containing phosphorus, nitrogen and silicon, was
19	synthesized with a promising approach to enhance flame retardancy of cotton fabric. To this aim, a
20	new sol-gel precursor, comprising in the same molecule P, N and Si, namely (3-
21	Glycidyloxypropyl)triethoxysilane modified N-(phosphonomethyl) iminodiacetic acid (PGPTES),
22	was co-hydrolysed and co-condensated with tetraethylorthosilicate (TEOS), as silane linker, and
23	used for producing a self-extinguishing cotton fabric coating. The structure of PGPTES was
24	characterized by ${}^{1}\text{H}/{}^{13}\text{C}/{}^{31}\text{P}$ nuclear magnetic resonance and the obtained coating was investigated
25	by FT-Infrared Spectroscopy and Scanning Electron Microscopy. The thermal properties of the
26	treated fabric were studied by Thermogravimetric Analyses and Cone Calorimetry Tests. The

obtained results show that the synthetized coating is able to catalyse the dehydration and char formation of cellulose based polymer at a lower temperature, thanks to the thermal decomposition of phosphate giving rise to acidic intermediates, able to further react with cellulose-based fabric, hence improving the flame retardant properties of the latter.

31

Keywords: Sol-gel; GPTES; N-(Phosphonomethyl)iminodiacetic acid; Textile finishing; flame
retardancy.

34

35 **1. Introduction**

36 Recently, nanotechnology has become a fast-growing area of research in the textile field because of 37 its many potential applications allowing the development and evolution of a new class of improved materials [1]. Advanced applications have been developing through textile or textile-based 38 39 materials, such as nanofibers, as well as nanocomposite fibres [2]. Meanwhile, nanoparticles are also successfully being used in conventional textiles to impart new functionalities and improved 40 41 performance [3, 4]. In fact, they have high surface energy and a large surface area-to-volume ratio, 42 which makes them easy to be linked to the treated substrate, increasing the durability of the 43 functions imparted to textile materials [5]. Currently, one of the biggest scientific and technological challenges is the design of new materials in order to develop innovative applications. In this 44 45 context, the role of preparative chemistry is to provide the compounds useful for obtaining 46 innovative materials: among them, hybrids exploit the peculiarities of both the organic and the 47 inorganic chemistries, hence giving rise to an almost unlimited number of applications. As an 48 alternative to chemistry employed for the surface modification, a range of solution techniques have 49 emerged, including co-precipitation, hydrothermal processing, solvothermal methods and sol-gel 50 chemistry [6]. Among these, the sol-gel approach shows some particular advantages, being centred 51 on the ability to produce a solid-state material from a chemically homogeneous precursor. The 52 "Design" of sol-gel materials - and their material properties - is to some extent possible by changing

53 the chemical composition and arrangement of the molecular building blocks and by deliberately tailoring their nano- and micro-structure. The sol-gel technique, consisting in hydrolysis and 54 55 condensation reactions, is based on hydrolysable precursors-building blocks - mostly metal or semimetal alkoxides (precursors); among them, the most widely studied are silicon alkoxides, 56 57 characterized by the strong covalent Si-O bonding and a hydrophobic behaviour that makes them immiscible with polar media. In textile applications, the most representative precursors are 58 59 organofunctional trialkoxysilanes (R'-Si(OR)₃), because of their unique structure bearing three 60 polimerizable groups, which enables the formation of a highly oriented polymer network structure 61 with an incorporated organic moiety.

The sol-gel process represents a simple method for the development of a coating with selected 62 63 protective properties, presenting many advantages, such as the possibility to achieve an environmentally-friendly surface functionalization of substrates and the easy adaptation of this 64 65 finishing process to the existing processing lines in industrial scale production [10–13]. Depending on the chemical structure of the network-modifying moiety, different functional properties can be 66 67 tailored on the material surface. Among these, it is worthy to mention antimicrobial [14], UV 68 radiation protection [15,16], biomolecule immobilization [17], dye fastness [16,18], anti-wrinkle 69 finishing [19] and super-hydrophobicity [20], as well as antistatic properties, odour control, stimuli-70 responsive performance [21] and strength enhancements [22]. In addition, since silica coatings exert 71 a thermal shielding effect on polymer surfaces improving the flame retardancy of the treated fabrics 72 [23], the use of sol-gel methods for conferring flame retardant properties to textile fabrics, in 73 particular cellulose-based fibres, has been documented by several research groups [24]. In fact, 74 cotton, thanks to its peculiarities such as strength, durability, flexibility and air permeability, as well 75 as good biocompatibility, low cost and good mechanical properties [25], is one of the most 76 important materials employed not only for producing apparel but also home furnishings and 77 industrial products, namely medical supplies, industrial thread and tarpaulins [26]. Unfortunately, 78 this cellulosic material has a low limiting oxygen index (LOI) and combustion temperature that

79 makes it highly flammable [27]. In order to meet fire safety regulations and expand the use of 80 cotton in textile applications that require flame resistance, a significant number of flame retardant 81 treatments has been developed in the last century [28], among which formaldehyde-based and 82 halogenated compounds have been the most employed. Although they show high performance with 83 excellent washing fastness, in most of them, the presence of active hydroxymethyl units causes the 84 release of formaldehyde from the treated fabrics both during the fabric application and throughout 85 the lifetime of the garment, which is not environmentally compatible [29]. For what concerns 86 halogenated compounds, recently, research studies regarding their persistence, ability to bio-87 accumulate, and potential for toxicity have led to increasing restrictions and regulations on the 88 production and use of some of them. In particular, such compounds as polybrominated biphenyls, 89 penta and octobromodiphenilethers have been banned, as they could generate corrosive and toxic 90 combustion products (e.g. dioxins and furans). [30]. Given the negative impact of formaldehyde and 91 bromine-based finishes on human health, since they are carcinogenic and bio-accumulative, 92 respectively, it is a primary focus for public safety to develop equivalent compounds, without 93 formaldehyde and halogens. Replacing the above-mentioned flame retardants finishes with 94 environmentally-friendly compounds represents an ecological step forward, in agreement with IPPC 95 European Directive [31]. Phosphorus-based flame retardants seem to be a valid alternative for the 96 above-mentioned FRs: in fact, unlike the halogen-containing compounds, which generate toxic 97 gases, corrosive smoke, or harmful substances [30], they act in condensed phase, by converting into 98 phosphoric acid or metaphosphoric acid during combustion or thermal degradation. Thus, non-99 volatile polyphosphoric acids can react with the decomposing polymer by esterification and 100 dehydration to promote the formation of protective char [32]. The development of the latter provide 101 lower flammability to fabric by protecting the underlying polymer from attack by oxygen and 102 radiant heat. Besides, it was found that organophosphorus containing active nitrogen have 103 multifunctional advantages and show higher effectiveness if compared with pure phosphorus 104 counterparts: a) low toxicity during combustion, b) high efficiency measured by cone calorimeter,

105 and c) low smoke development in fire accidents [6]. Some nitrogen-containing compounds seem to 106 accelerate phosphorylation of cellulose through the formation of a phosphorus-nitrogen polymeric 107 species, and thus synergize the flame retardant action of phosphorus. The reason could be attributed 108 to the type of bond between elements: P-N bonds are more polar than the already present P-O 109 bonds, and the enhanced electrophilicity of the phosphorus atom increases its ability to 110 phosphorylate the C(6) primary hydroxyl group of cellulose. By this way, the intra-molecular C(6)-111 C(1) rearrangement reaction forming levoglucosan is blocked. Meanwhile, the auto-crosslinking of 112 cellulose promotes and consolidates the char formation derived by the action of the same flame 113 retardants [33,34]. In addition to the above-mentioned compounds, silicon is demonstrated to be 114 used as a flame-retardant element because it is able to produce a continuous layer of silica that 115 retards char oxidation.

In this paper, we take advantage of the synergistic effect between silica and phosphorous in conferring flame retardant properties: in fact, the concurrent presence of P and Si elements in the same precursor can be exploited for preparing a hybrid coating that behave, at the same time, as a char promoter (with the same above-mentioned mechanism) and thermal shield, due to the metaloxide ceramic network [35–40].

121 In this study, (3-Glycidyloxypropyl)triethoxysilane (GPTES), one of the most used silica precursors 122 for hybrid silica-based textile finishing, was chosen since its ethoxysilyl groups and the reactive 123 epoxy group allow promoting the adhesion with the treated textile surface and the reaction with 124 other organic molecules, respectively. Furthermore, the epoxy ring can be simultaneously 125 crosslinked, via extended covalent bonds, to form poly- or oligo-(ethylene oxide) derivatives, thus 126 allowing the grow-up of a hybrid polyoxyethylene 3D-network. Carboxylic moieties of N-127 (phosphonomethyl) iminodiacetic acid (PMIDA), a nitrogen-containing carboxyphosphonate, react 128 with GPTES epoxy groups to form a β -hydroxy propyl ester, following the mechanism proposed in 129 Fig 1.



130

131

Fig. 1. Proposed mechanism of reaction between PMIDA and GPTES.

132

133 Combining the advantages of a facile synthesis with the use of a traditional application procedure, 134 the proposed phosphorus-functionalized sol-gel precursor (PGPTES) emerges as promising 135 candidate for next-generation of hybrid finishes based on the concurrent presence of Si, P and N, 136 with breakthrough performances. Considering that PGPTES possesses both an inorganic moiety and 137 a high phosphorus content, it is potential for serving as a facile, eco-friendly and efficient flame 138 retardant agent for cellulosic fibres. Furthermore, the mild reaction conditions of the process make 139 it very favourable for the deposition of a hybrid organic-inorganic coating during the finishing of 140 cotton fabrics. In this research, the structure and surface morphology of the untreated and treated 141 cotton fabric were investigated in detail by NMR and FT-IR spectroscopy and Scanning Electron 142 Microscopy (SEM), equipped with an energy dispersive X-ray spectroscopy (EDX). Then, the 143 surface chemical composition was investigated by X-ray Photoelectron Spectroscopy (XPS). 144 Thermogravimetric analysis (TGA), cone calorimetry (CC), horizontal and vertical flame spread 145 tests were exploited for evaluating the combustion behaviour, the thermal stability, as well as the 146 flammability of the treated cotton samples.

- 147
- 148 **2 Experimental part**
- 149
- 150 2.1. Materials
- 151

152 (3-Glycidyloxypropyl)triethoxysilane (GPTES, namely: $Si(OC_2H_5)_3C_3H_5O_2$), \geq 98%)) and

153 Tetraethoxysilane (TEOS, namely: Si(OC₂H₅)₄), \geq 98%) as sol-gel precursors, as well as N-154 (Phosphonomethyl)iminodiacetic acid hydrate (PMIDA, 97%, MW 227.11), monoethanolamine 155 (MEA), and HCl (all reagent grades) were purchased from Sigma Aldrich (Italy) and used without 156 any further purification. Scoured and bleached plain-weave cotton fabrics, with an areal density of 237 g/m², were supplied by Mascioni Spa, Varese, Italy. In order to remove impurities that would 157 158 scatter on the fabric surface randomly during manufacturing, before sol-gel treatments, all the 159 fabrics were carefully cleaned by washing in a 2% non-ionic detergent (Tergipal NRZ, linear 160 alcohol ethoxylate, kindly supplied by FTR SpA, Italy) at 40 °C for 20 min, then rinsed several 161 times with de-ionized water, and finally dried. Before all experiments, all the samples were placed 162 under standard laboratory conditions $65(\pm 4)$ % relative humidity and $20(\pm 2)$ °C temperature) for 24 163 h.

- 164
- 165 *2.2 Nanosol preparation and application process*
- 166

167 Pre-reacted precursor sol was initially prepared from a mixture of high purity GPTES and PMIDA. 168 To promote the reaction between the epoxy group and the hydroxyl groups of the phosphonate, the 169 reaction was carried out in the absence of water, with the goal of limiting the possibility that the 170 epoxy group can either undergo hydrolysis to form the corresponding diol or polyaddition reactions 171 and polyether linkages. Therefore, 10 g of PMIDA powder, finely grinded, were slowly added to 20 172 ml of a TEOS/GPTES mixture (precursors molar ratio 25:75) into a 100 ml flask. The mixture was 173 kept for 2 h at room temperature, under vigorous stirring, until the solution became clear. Finally, in 174 order to start stepwise hydrolysis and condensation of oligomeric intermediate, 18.5 ml of water 175 were added. Monoethanolamine (MEA) was used for adjusting the pH to 3 and to increase the 176 nitrogen content of the FR system. The so-obtained solution was stirred for 3 h to complete the 177 hydrolysis of both precursors. With the aim to produce the xerogel and investigate its chemical

structure, small amounts of the obtained sol were applied on glass slides, the solvent was removed
at 80°C for 1 h and the thin film was then cured at 170°C for 1 h.

The cotton fabrics (20 cm x 30 cm) were dipped in the hybrid sol and then passed through a tworoll laboratory padding machine at nip pressure of 3 bar with about 75% of wet pick-up. After drying at 90°C for 5 min, the fabric sample was cured at 170°C in a laboratory oven for 5 min. The treated cotton sample was coded as CO_T. The amount of coating deposited, calculated as add-on on the untreated sample (A, wt% owf), was calculated weighing the sample before (W₀) and after the padding-curing treatment (W₁), using a Mettler balance (10⁻⁴ g):

186 $A = \frac{W_1 - W_0}{W_0} \times 100$ Eq. 1

187 The value obtained from Eq. 1 represents the average of five independent replicates, with the 188 standard deviation always lower than $\pm 2\%$.

189

190 2.3 Characterization

191

192 FTIR spectra of treated and untreated cotton samples were recorded using a Thermo Avatar 370 193 spectrophotometer equipped with an attenuated total reflectance (ATR) device for solids analysis. 194 Spectra were analysed using Omnic 7.3 software. Fabrics were stored at room temperature for 48 h 195 in a stabilized atmosphere at 20°C and 60% RH. The analysis was performed with the samples placed onto a diamond crystal, within 4000 and 650 cm⁻¹, with 64 scans and a resolution of 4 cm⁻¹. 196 The collected spectra were normalized to the 1314 cm⁻¹ band, associated with the C-H bending 197 198 mode of cellulose. Since infrared absorption bands of the silica-based coating applied onto the 199 fabric surface are covered by the strong vibrational peaks of cellulose, FTIR analysis was carried 200 out also on pure xerogel in order to characterize it, thereby avoiding other influences. In addition, 201 based on the intensity and shift of vibrational bands of FTIR spectra, the treated sample was 202 compared with pristine cotton in order to assess the presence of the coating. The morphologies of

203 treated and untreated cotton fabrics, including the char residues after horizontal flame spread tests, 204 were observed using scanning electron microscopy (LEO-1450VP, with beam voltage fixed at 5 kV), equipped with an X-ray probe (INCA Energy Oxford, Cu-Ka X- ray source, k_{α} = 1.540562 Å), 205 206 which was utilized for performing elemental analysis. Thermogravimetric analyses (TGA) were 207 carried out on a TAQ500 apparatus, using a heating rate of 20°C/min in nitrogen and air 208 atmosphere (gas flow: 60 mL/min for both the atmospheres). The experimental error was ±0.5% on 209 the weight and $\pm 1^{\circ}$ C on the temperature. Combustion tests of square fabric samples (50 mm x 50 210 mm x 0.5 mm) were carried out on a Fire Testing Technology Ltd Cone Calorimeter, under ventilated conditions, using a 35 kW/m² irradiative heat flow in horizontal configuration. The 211 212 experiments were repeated four times for each material investigated to ensure reproducible and 213 significant data; the experimental error was within 3%. The following parameters were registered: time to ignition (TTI, s), peak of heat release rate (pkHRR, kW/m²), total heat release (THR, 214 assessed at the end of the test, MJ/m^2), ratio of carbon dioxide and carbon monoxide yields, and 215 216 final residue (%). The accuracy was up to 3 and 10% for CO and CO₂ yields, respectively. A Fire 217 Performance Index (FPI, %/s) was also calculated as final residue to TTI ratio and employed as an 218 evaluating parameter: the higher the FPI value, the better the flammability performance.

The flammability of the cotton samples in the presence of a flame spread was measured both inhorizontal and vertical configurations.

221 In the first case, the flame was applied on the short side of the specimen (50 mm) for 10 s and then 222 removed rapidly. Two horizontal marks were drawn on the specimens (at 25 and 75 mm from the 223 side, on which the flame was applied) and the time $(t_1 \text{ and } t_2)$ required to the flame to reach them 224 was measured. Besides, other relevant parameters, such as total burning time and final residue, were 225 evaluated. Alternatively, when the test was performed in vertical configuration, a methane flame 226 was applied for 5 s at the bottom of a fabric specimen (50 mm x 100 mm), repeating the test 3 times 227 for each formulation in order to get reproducible data. A Flammability Performance Index (FPI, 228 %/s) was also calculated as the ratio of final residue to the total burning time and used as an

229	evaluation parameter: the higher the FPI values, the better is the flame retardancy performance
230	Prior to flammability and combustion tests, all the specimens were conditioned at 23±1 °C for 48 h
231	at 50% R.H. in a climatic chamber.

¹H, ¹³C{¹H} and ³¹P{¹H} NMR spectra were recorded in D₂O solutions or in a CD₃OD/D₂O 232 233 mixture (1:1=v/v) at 500, 125 and 202 MHz, respectively; coupling constants (J) are given in hertz, 234 and the attributions are supported by heteronuclear single-quantum and multi-bond coherence 235 (HSQC-HMBC) and correlation spectroscopy (COSY) experiments; all proton NMR experiments 236 were run with a water suppression pulse sequence. 237 XPS measurements were performed by using a ESCALAB MkII spectrometer equipped with a non-238 monochromatized Al Ka source and a five channeltrons detection system. The samples were fixed 239 to the holder by metallic clip. The spectra were collected at 40 eV pass energy and the binding

energy scale was calibrated positioning the C 1s peak form adventitious carbon at BE = 285.0 eV.

All data were collected and processed by Avantage v.5 software.

242

243 **3. Results and Discussion**

- 244
- 245 3.1 NMR characterization and structure determination
- 246

The PGPTES xerogel film, covering a glass-slide, was carefully removed, chopped and suspended in a CD₃OD/D₂O mixture (1:1=v/v), in order to characterize the silylated derivative by means of homonuclear and heteronuclear ¹H, ¹³C{¹H} and ³¹P{¹H} NMR, mono- and bidimensional NMR spectroscopy. Figure 2 shows the recorded ³¹P{¹H}, ¹H, and ¹³C{¹H} NMR stacked plots, respectively, in comparison with the starting PMIDA molecule (on the bottom of the figures, in D₂O) and the suspended xerogel coating TEOS_GPTES_PMIDA (PGPTES), in a 1/1 CD₃OD/D₂O mixture.



255

Fig. 2. Stacked NMR spectra relative to a solution of starting PMIDA molecule in D_2O (bottom) and a suspension of PGPTES xerogel (top) in a CD_3OD/D_2O (1:1=v/v) mixture as solvent at 298 K: **A.** ³¹P{¹H} NMR (202 MHz); **B.** ¹H NMR, with the main proton assignment; (500 MHz); **C.** ¹³C{¹H} NMR, with the main carbon assignment (125 MHz).

261 The ³¹P{¹H} NMR spectra in Figure 2A clearly shows the expected lower frequency shift of the ³¹P 262 signal relative to the PMIDA molecule from $\delta = 6.71$ (red square) to 6.52 ppm (blue square),

evidencing that a more shielding environment is enclosing the ³¹P nucleus of the PMIDA, after the
 reaction with GPTES. Two minor ³¹P containing species are present in solution at very slight
 concentration (highlighted in Figure 2A with a star and double stars).

In agreement, the aliphatic regions of the ¹H NMR spectra in figure 2B clearly show: (i) the 266 267 corresponding upfield proton shifts for the two methylene groups CH_2 -COOH and CH_2 -P, that shift from $\delta = 4.17$ and 3.44 (m, ${}^{2}J_{PH}=12.3\text{Hz}$) ppm, to $\delta = 4.10$ and 3.36 (m, ${}^{1}J_{PH}=12.5\text{Hz}$) ppm, 268 269 respectively (red square and circle vs blue square and circle); (ii) the presence of the expected 270 protonic pattern for the GPTES open ring derivative, bringing a hydroxyl and an ester group bonded to two vicinal carbon C_e and C_f atoms ($\delta = 0.47$, CH_{2a} ; 1.51, CH_{2b} ; 3.36, $CH_{2c}+CH_{2d}+CH_{2f}$; 3.66 271 CH_{2e}), in a 2:1 concentration ratio with respect to protons belonging to the PMIDA molecule [41-272 273 43]; (iii) the presence of the methylene and the methyl proton resonances relative to free ethanol 274 moieties.

Similar results can be drawn with the assignments of the ${}^{13}C{}^{1}H$ NMR spectra, shown in figure 275 276 2C, that undoubtedly display: (i) the presence of the pattern expected for the PMIDA_GPTES fragments, characterized by almost the same ¹³C chemical shift for the carboxylic group and for the 277 two methylene CH₂-COOH and CH₂-P, as well as the starting PMIDA molecule (i.e. $\delta = 168.6$; 278 55.9, ${}^{3}J_{PC} = 3.4$ Hz; 51.3 ppm, ${}^{1}J_{PC} = 136$ Hz vs $\delta = 168.4$; 56.1, ${}^{3}J_{PC} = 2.6$ Hz; 51.2 ppm, ${}^{1}J_{PC} = 135$ 279 280 Hz, respectively); (ii) diol, dioxane and polyethyleneoxide silvlated functionalities (not assigned 281 peaks) embedded in the final xerogel matrix [43], together with other minor phosphorous 282 containing PMIDA derivatives (star and double stars).

All these findings confirm that the GPTES epoxy ring opening reaction has successfully taken place, combined with the complete hydrolysis of the alkoxysilane end group. Even if solvent effects (due to a 50% CD_3OD presence as solvent for the PGPTES spectra) should be taken into account especially in the ¹H spectra assignments, the high and already well-known nucleophilic substitution reactivity of the carboxylic groups towards the epoxy rings [42,44] led us to conclude that, in the final xerogel, the PMIDA molecule is mainly bonded firmly and covalently through two ester bonds

to the sol-gel based 3D matrix [45,46]; minor species may be assigned most probably to the monosubstituted ester species or to the fully substituted species through the two acetic groups and the phosphorous end. As previously shown, the epoxy ring opening of GPTES gives rise to the formation of diol and silylated dioxane/polyether (PEO), through several hydrolysis and polymerization reaction steps [43].

According to the reported results, a scheme of the occurred reaction is presented in Fig. 3 showing the formation of a N-(Phosphonomethyl)imino diacetate hydroxy polysiloxane derivative through a starting 1:2 molecular species, namely N-(Phosphonomethyl)imino bis{2-hydroxy-3-[3-(triethoxysilane)propoxy]propyl acetate]}.



- 298
- 299

Fig. 3 Scheme of the reaction occurring between PMIDA and GPTES

300

303 Scanning electron microscopy was utilized to investigate the morphological features of untreated 304 and coated cotton fabrics. Fig. 4 shows SEM micrographs of cotton and the related EDX spectra 305 before and after hybrid sol-treatment. The latter results in 25.2% add-on, calculated according to 306 equation 1. In both magnifications, images of the untreated cotton sample show a flat assembly with 307 a twisted ribbon-like structure caused by spiralling of cellulose fibrils. The surface appears clearly 308 smooth with its veined natural morphology. After treatment, fibres are homogeneously covered by

³⁰¹ *3.2 SEM-EDX analysis*

309 the hybrid coating, showing free gaps between the warp and weft threads that maintain the same 310 aspect of the control cotton fabric, suggesting that the sol-gel coating on the treated sample is very 311 thin. Furthermore, semi-quantitative EDX investigation, employing a high beam voltage (i.e. 20 kV), confirms the key information concerning the element composition of samples. With the aim of 312 313 assessing the homogeneous distribution of coatings, five repeated measurements were carried out 314 on different parts of each cotton sample. Although the maps reported in the images are qualitative, 315 in the treated samples only phosphorus and silicon are present, along with carbon and oxygen 316 shown in the control fabric, since the technique cannot detect nitrogen atom. Furthermore, the data 317 listed in Tab. 1 show very similar results in all repeated tests, confirming a uniform presence of the 318 above-mentioned elements on the treated fabric sample.

319



Fig. 4. SEM images of untreated (CO_UT) and treated (CO_T) cotton samples, at different magnifications, coded as _1, _2, for x2.50 K, x1.00 K, respectively.

- 323
- 324

325

326

327 Table 1.

Sample		C [%]	O [%]	P [%]	Si [%]
CO_UT	Unburned	46.10±0.07	53.9 ± 0.50	/	1
	Burned*	/	/		1
CO_T	Unburned	44.19±0.03	49.32±0.73	2.63±0.58	3.86±0.87
	Burned	60.53±1.33	30.01±0.54	5.9±1.51	4.23±0.06

328 Results (wt.%) of EDX analysis of untreated and treated samples.

- 329 * no residue was collected after flammability test.
- 330
- 331 3.3 XPS analysis
- 332

In order to investigate the surface chemical composition of both coated and untreated cotton fabrics, 333 334 the samples were investigated by XPS analysis. The presence of C and O was registered on both 335 samples, while the coated sample also showed the presence of Si, P and N. The shape of the C 1s 336 signal (Fig. 5(a)) was characteristic for cotton fabric, characterized by three peaks positioned at BE 337 = 285.0 eV, 286.7 eV and 288.5 eV, with intensity ratio 1 : 1.1 : 0.2, and assigned to C–C/C–H, C– 338 OH and C=O bonds, respectively. The sol-gel coating slightly modified the shape of C 1s signal 339 (Fig. 5(b)), because it was very thin. However, the formation of the sol-gel coating was confirmed 340 by the presence of Si, P and N. After flammability tests, the shape of C 1s signal significantly 341 changed (Fig. 5(c)), where the intensity ratio of the peaks became 1: 0.3: 0.1 and a fourth peak was 342 positioned at BE = 291.4 eV, due to the presence of carbonates. This changing was not unexpected, 343 because it is typically for residual char samples [47]. Comparing the XPS quantitative analysis, 344 shown in Tab. 2, it can be noted that sol-gel coating was partially decomposed during the 345 flammability test, with a decrease of the atomic concentration of Si and an accumulation of P and C

elements on the outermost layers of the substrate. These last results may appear inconsistent with those obtained by EDX analysis (Table 1). This is due to the different depth of each analysis. In fact, XPS technique investigates the first layer of the sample (< 10 nm), while information depth of EDX is approximately 1 μ m. Furthermore, the results are comparable (e.g. Si/P ratio) even if they are given in wt% (EDX) and in at. % (XPS).

351



352

- **Fig. 5.** Comparison of C 1s spectra of (a) untreated cotton fabric (CO_UT), (b) treated cotton fabric
- 354 (CO_T) and (c) carbonaceous char (CO_C).

355

- 356 **Table 2.**
- 357 Surface chemical composition of the untreated cotton fabric (CO_UT), treated cotton fabric (CO_T)

		C (%)					O (%)	P (%)	Si (%)
	C1:	C2:	C3:	C4: -	C4':	NR ₃	-ОН	Phosphate	SiOR
	С-С/С-Н	С-ОН	C=O	СООН	CO ₃	5		Ĩ	
BE (eV)	285.0	286.6	288.1	289.2	291.4	400.5	532.8	133.7	103.3
CO_UT	29.1	32.7	7.2				31.0		
CO_T	26.4	27.6	5.8	3.7		2.3	29.1	1.7	3.5

358 and carbonaceous char (CO_C).



- 359
- 360 **3.4 ATR-FTIR spectroscopy**
- 361

The ATR-FTIR spectra of the untreated, treated cotton fabrics and pure xerogel prepared on glass 362 363 slides, are compared in Fig. 6 and the most important bands are collected in Table 3. The analytical 364 measurement of modified GPTES hybrid sol has been carried out, since the strong vibrational bands 365 of the pure substrate could hide the characteristic peaks of the thin film applied onto the fabric surface. Comparing the xerogel spectrum with that of PMIDA, it can be highlighted the absence of 366 a broad band at 2650-2450 cm⁻¹, characteristic of such type of compounds and attributed to P-OH 367 stretching vibration. Also the band at 929 cm^{-1} , typical of P-OH bending signal, disappears. These 368 findings, combined with the shift of the stretching vibration of P=O from 1215 cm⁻¹ to higher 369 wavenumbers (1267 cm⁻¹), demonstrate an effective interaction between phosphorous compound 370 and silica matrix. The other characteristic peaks that can be seen from xerogel spectrum are located 371 at 1736 cm^{-1} (stretching vibration of ester carbonyl), at 1635 cm^{-1} and 1540-1520 cm^{-1} (typical of 372 the absorption of NH₂ group). The region between 1200 and 750 cm^{-1} includes peaks characteristics 373 of silicon matrix: 1194 cm⁻¹ (Si-O-Si asymmetric stretching), 1045 cm⁻¹ (Si-O-Si), 1009 cm⁻¹ (Si-O 374 stretching), 907-930 cm⁻¹ (Si-OH stretching), 763-790 cm⁻¹ (Si-O-Si symmetric stretching). 375 376 Moreover, the opening of the epoxide ring was observed through the absence of its characteristic infrared bands at 1255 cm⁻¹ (ring breathing), 907 cm⁻¹ (asymmetric ring deformation) and 851 377 cm⁻¹ (symmetric ring deformation) [46, 48]. FTIR spectroscopy was also carried out in order to 378 379 confirm the successful reaction between the hybrid coating and cotton. As shown in Figure 5, the 380 spectrum of untreated cotton exhibits O–H stretching absorption between 3500 and 3000 cm⁻¹, C–H stretching absorption around 2950–2850 cm⁻¹, and C–O–C stretching absorption around 1160 cm⁻¹. 381 382 These bands are consistent with those of the typical cellulose backbone. The spectrum of the coated cotton appears quite similar to that of the untreated one. With respect to the latter, an overall slight 383

384 decrease of the intensities of characteristic hydrogen-bonded OH stretching vibrations is observed: 385 this finding may be attributed to the interaction of the coating with the cotton functional groups. In 386 the treated cotton sample, some new characteristic peaks appear, such as the broad absorption band 387 in the region 2970-2850, which is attributed to the introduction of the -CH₂ group; these peaks are 388 proportional to the quantity of carbon included in the grafted molecules and, in particular, the band of asymmetric stretching vibration of the methylene group located at 2925-2970 cm^{-1} and a band 389 attributable to symmetric stretching vibration at 2850- 2920 cm^{-1} can be detected. The absorption 390 band at 1732 cm^{-1} was assigned to the stretching vibration of carbonyl of ester group [49]. The 391 majority of the peaks typical of -PO₃ moiety are hidden by intense cellulose bands; the only one that 392 is clearly visible is at 1269 cm⁻¹, assigned to the P=O stretching vibration. The most important 393 peaks attributable to the sol-gel coating were identified at 1008, 930 and 765 cm⁻¹ assigned to Si-O-394 395 Si asymmetric stretching, Si-OH stretching, and Si-O-Si symmetric stretching, respectively. The peak expected to be at about 1040 cm⁻¹ is overlapped with a broad band between 1050 and 1014 396 cm⁻¹ attributed to the characteristic peaks of cellulose. Some changing in the intensity of IR bands 397 appearing onto the treated textile fabric in the range between 1240 and 1160 cm⁻¹ are assigned to 398 Si-O-C bonds, thus confirming the reaction between the hydrolysed silane precursor and the 399 400 cellulosic substrate. Although these peaks may be considered weak, they are of great importance 401 because they are a proof of the interaction between the substrate and the precursor [50]. Furthermore, the presence of amino group from MEA is observed at about 1629 and 1526 cm⁻¹ due 402 403 to symmetric and asymmetric N-H bending modes, respectively.



405

- 406 Fig. 6. AT-FT IR of hybrid coating onto glass slide (PGPTES), untreated (CO_UT) and treated
- 407 cotton (CO_T) samples.

- 409 **Table 3**
- 410 Major vibrational frequencies of the sol-gel based films.

Experimental wa	venumbers (cm ⁻¹)	Literature wavenumbers	Vibrational mode	
Glass substrate	Fabric substrate	(cm ⁻¹)		
3300	3500-3000	3500-3000	ν (O-H)	
		[51]		
2850	2950-2850	2980–2800	ν (C-H)	
		[51]		

	ACCEPT	ED MANUSCRIPT	
1736	1732	1735	v (C=O)
		[52]	
1635, 1540-1520	1629, 1526	1600, 1575	v (N-H)
		[53]	
_	1424	1429	ω (C-H)
		[51]	
1266	1269	1267	ν Ρ=Ο
		[54]	
	1244	1240	Si-O-C
		[55]	
1194	1198	1200	v _{as} (Si-O-Si)
		[51]	
	1159	1160	v_{as} C-O-C
		[51]	
1009	1008	1001	v (Si-O-Si)
		[51]	
907-930	930	952	v (Si-OH)
		[51]	
790-763	765	749–786	v _s (Si-O-Si)
		[51]	

411

412 **3.5 Thermal behaviour**

413 The thermal and thermo-oxidative stability of the untreated and sol-gel treated cotton fabrics has

414 been assessed by thermogravimetric analyses performed in nitrogen and air, respectively.

415 The TG and dTG thermograms of sol-gel treated and untreated cotton samples are shown in Fig. 7;

416 Table 4 collects the corresponding data in terms of T_{onset} , T_{max} (corresponding to the peak(s) in dTG

curves) and related residues and final residue at 700°C. For both the environments, the weight loss 417 418 up to 100°C, due to absorbed moisture in all samples, was not considered for this degradation study. 419 In nitrogen, the TG curves of untreated cotton sample show the onset degradation temperature at 420 about 310°C, together with its maximum mass loss rate at about 360°C, due to depolymerization by 421 trans-glycosylation reactions. The coated fabric shows an anticipation in the onset degradation 422 temperature, reaching the maximum mass loss rate at 275°C: this finding is attributed to the earlier 423 degradation of the phosphorus-containing compound that catalyses the decomposition of cotton 424 towards the formation of a carbonaceous residue (char). This latter increases the thermal stability of 425 the fabric: in fact, at the end of the test, the char residue achieves 38%, significantly higher than 426 pure cotton, for which the residue is below 6%.

In air, the thermo-oxidation of cotton takes place in a similar way: the only difference is for the appearance of a second degradation step at high temperatures (T_{max2} : 460°C). This phenomenon can be attributed to the oxidation of the char formed during the first step and of all the hydrocarbon species still present [56]. Once again, the hybrid coating is responsible for the decrease of both T_{onset} and T_{max1} as well as for the increase of the residues at T_{max1} , T_{max2} and 700°C: these findings confirm the protective effect exerted by the formed stable char.

433 Compared to untreated cotton control, the effectiveness of the silica-based coating as a flame 434 retardant for cotton, directly attributable to the considerably higher energy and ionic character of the 435 Si-O bond (443.7 kJ/mol) relative to the C-C bond (345.7 kJ/mol) [57], is indicated by a substantial 436 lowering of the decomposition temperature, due to the earlier degradation of the phosphorus 437 contained in the modified precursor, which is able to catalyse the dehydration of the treated cotton 438 to form the intumescent char. As confirmed by TGA results, the higher the residue in the silica-439 phosphorylated cotton, the lower is the amount of volatile products obtained upon decomposition of 440 the samples during the test.



Fig. 7. TG and dTG curves of CO_UT and CO_T samples, in: A. nitrogen and B. air atmospheres. 443



445 Table 4.

Atmosphere: Nitrogen						
	T _{onset} (°C)	T _{max1} *	Residue	T _{max2} *	Residue	Residue
Sample	(s)	(° C)	@T (%)	(° C)	@T _{max2}	@700°C
	(3)			(C)	(%)	(%)
CO_UT	310	360	40	/	/	5.8
CO_T	275	310	70	/	/	38
			Atmosphere: A	ir		
CO_UT	300	340	46	460	3.2	/
CO_T	260	300	73	520	30	12

446 TG data for CO_UT and CO_T in nitrogen and air.

* from dTG curves 447

449 **3.6** Cone calorimetry tests

450

The forced-combustion behaviour of cotton samples was investigated through cone calorimetry tests, measuring the time to ignition (TTI), the peak heat release rate (pkHRR), the total heat release (THR) and the final residue (%), as well as CO and CO₂ yields (%).

454 The curves of HRR vs. time for control and silica-coated fabric are shown in Fig. 8B. From Tab. 5, 455 it can be seen that if compared to untreated cotton, the coated fabric shows a reduction in both heat 456 release rate peak (pkHRR) and in the total heat release (THR). In fact, with respect to the uncoated sample, in the treated fabric pkHRR decreases from 154 to 135 kW/m², corresponding to an 457 458 approximately 12% reduction. The THR also drops significantly, going from 3.9 down to 2.3 MJ/m^2 . In the case of coated sample, it is worthy to note that after ignition, the HRR value increases 459 460 rapidly in a very short time (from 25 to 40 s) and then decreases sharply. This behaviour is 461 attributable to the formation of an intumescent char instantly after ignition by the phosphorus-462 containing silica coating, indicating that more cotton fabric participates in the carbonization 463 process, due to the presence of the deposited coating. Consequently, less degradation products that 464 serve as "fuel" go into the gas phase, hence lowering the pkHRR and THR values. In addition, in 465 order to evaluate the fire performance index (FPI), the ratio between final residue and pkHRR 466 values for treated and untreated samples was calculated. The FPI value of the control cotton is 467 0.023 %/s, whereas that for silica coated cotton is 0.76 %/s. The higher FPI value for the coated 468 cotton sample justifies the higher amount of final residue and the shorter time to ignition with 469 respect to the untreated counterpart. Based upon the aforementioned fire behaviour, it is believed 470 that the formation of intumescent char layer is responsible for suppressing fire propagation at the 471 selected heat flux, or greatly inhibiting the amount of flammable gases available for combustion. 472 Finally, through CO and CO₂ analysis it is possible to provide useful information on the mechanism 473 of decomposition of cotton fabrics, since low CO₂/CO ratio means low conversion of CO to CO₂, 474 thus suggesting inefficiency of combustion. When cotton fabrics were treated with PGPTES, the

475 CO production changed a little, whereas the CO₂ production was decreased, leading to a remarkable 476 decrease of CO₂/CO ratio (19.5). This suggests that the designed flame retardant mainly acts in 477 condensed phase. It should be remarked that the char residues from cone calorimeter tests, reported 478 in Figure 8A, are consistent with high-temperature residues obtained by TG analysis. The poor 479 residue from the neat sample was completely broken, hence confirming the low charring ability of 480 cellulose. On the other hand, the residue of the coated sample shows a coherent and dense char, 481 maintaining a compact structure and its original texture. This is a direct consequence of the silica 482 precursor formulation containing phosphorus and nitrogen. In fact, both species can synergistically 483 act [28,51], catalysing the formation of the char layer that limits the heat and mass transfer, thus 484 reducing the formation of flammable gases. At the same time, this char was made more stable by 485 the beneficial presence of silica that does not allow the fire spread.

486



Fig. 8. A. Residues of CO_UT and CO_T from cone calorimetry tests performed at 35 kW/m²; B.
Heat release (HRR) curves of CO_UT and CO_T.

- 491
- 492

493

494 **Table 5.**

495	Combustion data of CO	UT and CO T from con	e calorimetry tests	performed at 35 kW/m ²
170	comoustion adda of co_		e calonineary costs	

	TTI	Flame out	pkHRR	THR	Residue	CO	CO ₂	
Sample	(s)	(s)	(kW/m ²)	(MJ /m ²)	(%)	(%)	(%)	02/00
CO_UT	43±2	75	154±4	3.9±0.1	1	0.0014	0.20	143
CO_T	34±2	48	135±3	2.3±0.1	26	0.0087	0.17	20

496

497

498 **3.7 Flammability tests**

499

500 In order to evaluate flame retardant performance, the uncoated and coated fabrics were subjected to 501 horizontal and vertical flame spread tests. The images of the burned samples, together with the 502 flammability data (after flame time, afterglow time and residue (%)), for both vertical and 503 horizontal configurations, are presented in Table 6. At the end of the tests, the residues the uncoated 504 fabrics have been completely destroyed, leaving only few amounts of ash, whereas for the treated 505 sample there was only smoulder progression of fire that stopped before the scribed line for 506 measurement. More in detail, in horizontal configuration, immediately after ignition, on the 507 untreated sample a vigorous flame appears for about 23 s, followed by 139 s of afterglow that did 508 not leave any residue. Conversely, the treated fabric sample shows no visible after-flame time, and 509 very short after-glow time (8 s) and achieves self-extinction a few seconds after the flame 510 application. As a result, at the end of combustion, the residue is 99.5% and the Flammability 511 Performance Index (FPI) is 12.44%/s. Also in vertical configuration, the untreated cotton fabric is 512 fully consumed during the test; conversely, the treated sample completely stops the flame 513 propagation almost as soon as the flame is removed, showing neither afterflame, nor afterglow and

514 obtaining self-extinguishing classification. Once again, the results confirm the good behaviour of 515 the phosphorous and nitrogen containing sol-gel coating onto cotton fibres, as a dehydrating and 516 char-forming layer, which hinders heat, fuel and oxygen transmission by creating a ceramic barrier 517 onto the treated fabric surface from further burning.

518

519 **Table 6**

Flammability behaviour and data related to the horizontal and vertical tests of CO_UT and CO_Tsamples.





522

523 In order to understand the char composition and to investigate the mechanism of flame retardancy, 524 achieved with the hybrid coating, after the vertical flame spread tests, SEM analysis and ATR-FTIR 525 spectra of burned coated cotton fabrics were carried out. SEM micrographs (Figure 9) show that the 526 charred region in the treated sample still maintains the shape of the original fibres and its weave 527 structure, displaying only minor shrinkage. Fibres look very rough and show many bubbles on their 528 surface: these findings are attributed to swelling and expansion of the coating due to its intumescent effect. It's possible to observe big bubbles on the surfaces of charred layer of FR cotton sample. 529 530 These bubble char not only inhibits the release of flammable gases from cellulose degradation but 531 also prevents the heat source to convey heat to the cellulosic substrate and insulates the oxygen 532 source [52]. The intumescent layer protects the fibres from further burning, preserving the woven 533 structure and fibre integrity, which is responsible for the self-extinguishing phenomenon of cotton. 534 Besides, the results of the elemental analysis carried out on the same residue still indicate the 535 presence of phosphorous and silicon, homogeneously distributed, notwithstanding the presence of 536 carbon and oxygen, as for the control sample. This good performance was mainly produced by three 537 components of developed molecule: first, the phosphorous group able to release P-based acids that 538 catalyse the dehydration of cotton to form char. This char is able to prevent heat, fuel, and oxygen 539 from being transferred from the flame to the fabric. Second, the deoxyribose units acted both as a

540 carbon source and blowing agents, with which a (poly)saccharide dehydrates to form char and 541 releases water upon heating. Third, the nitrogen-containing base released ammonia, which can 542 further induce char development and produce non-combustible gases such as N_2 and CO_2 .



543

Fig. 9. A. SEM images of the residues of CO_T at different magnifications, coded as _1, _2, for
x2.50 K, x1.00 K, respectively, B. EDX images of the treated sample; C. ATR-FT IR of treated
cotton fabric pyrolyzed at 600°.

547

To investigate the chemical composition of the burned sol-gel coated fabric after the vertical flame spread test, the sample was examined by ATR-FTIR: its spectrum is shown in Figure 9C and detailed band assignments are provided alongside the spectrum. The presence of aromatic-type structure was confirmed by the intense band at around 1590 cm⁻¹ and a shoulder at 1200 cm⁻¹, ascribed to the presence of polynuclear aromatic structures (-C=C- stretching) [53] and the vibration

of CH groups, out of plane bending, respectively. The shoulder peak at 1200 cm⁻¹ may also 553 554 comprise the signal assigned to the functionality of a phosphorus-nitrogen structure. Furthermore, the peaks in the region 1000–650 cm^{-1} may be assigned to the out of plane deformation vibrations 555 of the ring [61]. Finally, the peaks at 1070 cm⁻¹ are ascribable to the Si–O–Si stretching vibration 556 557 that confirms the formation of a silica matrix in the char. These characteristic absorptions are in 558 accordance with depolymerization of cellulose and formation of char [62]. From both analyses it is 559 clear that the char of the treated sample seems to consist of phosphorous, carbon and silicon-rich 560 compounds. The carbonaceous char that originates on the surface of the burning fabric during 561 combustion is thus covered by silicates and phosphonates, hence creating an excellent physical barrier, which protects the substrate from heat and oxygen, and slowing down the escape of 562 563 flammable volatiles generated during cellulose degradation.

564

565 **4. Conclusions**

566

A novel sol-gel based flame retardant containing phosphorus, nitrogen, and silicon was synthesized 567 successfully, and its chemical structure was fully characterized by Fourier transform infrared 568 spectrometry and nuclear magnetic resonance spectrometry (¹H NMR and ¹³C/³¹P NMR). After its 569 570 application on cotton fabric, the combustion behaviour of the treated sample was investigated, 571 confirming that the so obtained coating acts as an efficient flame retardant in the condensed phase. 572 In particular, the presence of PGPTES lowers the decomposition temperature and favours the 573 formation of char after pyrolysis; in addition, in forced combustion tests, the coating is responsible 574 for the increase of TTI and the reduction of both HRR value and CO/CO₂ ratio for treated cotton 575 sample. These results demonstrated the formation of a compact and thermostable char residue that 576 effectively improved the thermal stability of cotton fabrics by hindering the formation of volatile species and favouring the creation of a stable char. It can be concluded that the hybrid GPTES-577 modified precursor can be potentially used as a new flame retardant replacing halogen-based 578

579 finishes in the field of flame retardant textile materials. Combining the above-mentioned 580 advantages, it emerges as a promising candidate for the design of a next-generation of hybrid 581 materials with breakthrough flame retardant performances. Further research will be developed in 582 order to investigate the washing fastness of the proposed sol-gel coating and its influence on the 583 mechanical properties of treated cotton fabrics.

584

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HIGHLIGHTS

- NMR analysis confirms the functionalization of GPTES sol-gel precursor with PMIDA.
- PMIDA acts as nitrogen and phosphorous source in the P/N flame retardant synergism.
- The concurrent presence of Si, P and N enhances cellulose dehydration mechanism.
- The final residue for treated cotton sample, after cone calorimetry test, is 26%.
- Self-extinguishing properties are achieved by the treated cellulose-based fabric.