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2 Effect of liquid impregnation on DBD atmospheric

3 pressure plasma treatment of cotton

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- 19
- 20 Abstract

21 This paper describes the He non-thermal plasma treatment of liquid impregnated cotton fabrics. The cotton 22 fabrics were soaked in either H_2O , D_2O or ethanol after which they were placed in an atmospheric pressure 23 parallel-plate DBD reactor. The influence of the used liquid in combination with the plasma exposure time was 24 studied using OES, FTIR, XPS and SEM. The addition of (deuterated) water was found to aid more efficiently in 25 the incorporation of polar functional groups onto the cotton surface in comparison to a pure He plasma treatment. 26 The presence of H and OH species in the discharge also caused extensive etching of the surface, which led to the 27 formation of microcraters. The impregnation with ethanol was responsible for the formation of a plasma-based 28 thin film on top of the cotton substrate, which was characterized by a high content of C-C bonds and a smooth 29 surface morphology. These results show that the soaking of cotton fabric prior to plasma exposure can help to 30 more effectively alter its surface properties compared to a dry plasma treatment.

- 31
- 32 *Keywords:* Cotton, DBD, Impregnation, Plasma treatment

33 Introduction

34 Non-thermal plasma treatments using oxygen-rich discharge gases are known to introduce a

variety of polar oxygen-containing functional groups onto the surface of cellulosic materials

including -OH, C=O and O-C=O groups, which can greatly enhance the surface wettability,

37 dyeability and adhesion of biopolymer coatings such as chitosan. Several papers have already

investigated the effect of atmospheric pressure non-thermal (or cold) plasma treatment on the

properties of cotton fabrics (Aileni et al. 2019; Haji 2017; Haji 2019; Haji et al. 2013; Haji et

al. 2016; Patino et al. 2011; Rajasekaran and Muthuraman 2019; Sun and Stylios 2004; Wang 40 et al. 2019). In particular, non-thermal plasma activation in the presence of water in liquid or 41 vapor phase is a well-known technique to incorporate functional groups onto a polymer 42 surface. In this context, underwater plasma treatments have been studied sporadically to 43 fabricate substrates with homogeneous monofunctional surfaces (Joshi et al. 2009b), to induce 44 45 direct functionalization, stimulate cross-linked layer formation and/or to graft or deposit mono-functional layers (Friedrich et al. 2008; Joshi et al. 2013; Joshi et al. 2009a; Joshi et al. 46 2008; Khlyustova et al. 2015). Despite these promising results, the presence of moisture 47 during a plasma surface treatment is often still viewed upon as an impurity (Ananth and Mark 48 2008). As a result, to date, only few studies have looked at the intentional addition of 49 liquids/vapors to discharge systems used for surface modification (Acsente et al. 2016; De 50 Geyter et al. 2013; Van Deynse et al. 2014; Van Deynse et al. 2017). Moreover, most of these 51 studies only focused on the addition of low amounts of vaporized water to the discharge gas 52 (Acsente et al. 2016; De Geyter et al. 2013; Van Deynse et al. 2014). However, more recent 53 54 studies have shown that, upon the addition of a liquid or vapor to the plasma discharge, the steady-state wettability of the plasma-exposed surface can be effectively altered, opening up 55 new possibilities for enhanced surface modification of different substrates (Acsente et al. 56 2016; De Geyter et al. 2013; Van Deynse et al. 2014; Van Deynse et al. 2017). Within our 57 research facilities, research in recent years has been focused on the impregnation of porous 58 polymeric materials (e.g. cotton, filter paper) with precursor solutions to obtain polymer-like 59 coatings by means of *in-situ* plasma polymerization (Molina et al. 2014; Molina et al. 2017b). 60 Whereas for plasma polymerization where precursors are applied in the gas phase, the coating 61 properties are modulated by the precursor flow rate, system pressure, discharge power and 62 plasma treatment time (Yasuda and Hirotsu 1978), plasma polymerization ignited onto a 63 64 liquid-impregnated substrate is usually modulated by tuning the concentration of the precursor present in the liquid (Molina et al. 2014). However, to the best of our knowledge, the direct 65 functionalization of a porous polymeric substrate by non-polymerizable liquid impregnation 66 plasma has not been studied in detail until now. For this reason, in this study, cotton fabrics 67 68 were therefore impregnated in either H₂O, D₂O or ethanol (EtOH) prior to performing plasma surface treatments. Helium (He) was chosen as discharge gas since a global model of He/H₂O 69 plasma chemistry in a parallel plate DBD reported the abundance of H_2O^+ , water cluster ions, 70 atomic and molecular metastable oxygen, OH(A) metastables and different ground-state 71 neutrals such as H,O, H₂, O₂, OH, HO₂ and H₂O₂ in He/H₂O plasmas, with varying densities 72 depending on the concentration of water vapor in the feeding gas (Liu et al. 2010). By first 73 impregnating the cotton samples in water, the hypothesis is that the local concentration of 74 water molecules at the substrate-plasma interface can be greatly enhanced, which may have a 75 positive effect on the plasma surface modification process. Moreover, by replacing normal 76 77 water with its deuterated form, better insights can be acquired on the role that H and OH species play in the surface modification processes. Finally, impregnation with EtOH is also 78 examined as it can also introduce additional OH species in the plasma, which may in turn 79 enhance the plasma surface treatment effects. Analysis of the active plasma species present in 80 the plasma gas phase was done by optical emission spectroscopy (OES), while the plasma-81 induced effects on the cotton fabrics were analyzed by X-ray photoelectron spectroscopy 82 83 (XPS), Fourier-transform infrared spectroscopy (FTIR) and scanning electron microscopy 84 (SEM).

86 **Experimental**

87 Materials

Cotton fabrics (plain weave, woven textiles, bleached without optical brightener with a 180 88 g/m^2 weight, article 210 from EMPA) were used as substrates. Concentrated detergent for pre-89 treating textiles (Tanaterge® Advanced) was acquired from Tanatex. Cotton was first 90 conditioned by washing the fabric for 45 minutes at 60°C, under constant stirring at 250 rpm. 91 The washing solution was prepared using Tanaterge® as detergent at 2 g/L concentration and 92 93 washing was carried out at a 1/25 (w/v) load/wash solution ratio. Subsequently, the fabric was rinsed by placing it into deionized water for 20 minutes. Finally, the cotton fabric was dried 94 and stored in a conditioned room at 67% relative humidity (RH) at a temperature of 24°C. 95 Deionized water, deuterium oxide (D₂O) with $D_2 \ge 99.96\%$ provided by Eurosi-top and EtOH 96 of absolute degree provided by Merck were used as impregnation liquids. 97

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99 Impregnation and plasma treatment

A DBD reactor operating at atmospheric pressure was used in this work, similar to the one 100 reported in previous work (Jovančić et al. 2016; Molina et al. 2013a) and is schematically 101 depicted in Figure 1. A gas mass flow controller (Bronkhorst, the Netherlands) was used to 102 introduce helium gas at a fixed rate of 5 standard liters per minute (5 slm) in the reactor 103 chamber between the upper and lower electrode, which were both covered with a dielectric 104 material (borosilicate glass). The gas gap between the two dielectrics was maintained at 12 105 mm and the volume between the electrodes was approximately ~0.1 liters. A 16-kHz AC 106 signal was generated with a GF-855 function generator (Promax, Spain) connected to a linear 107 amplifier AG-1012 (T&C Power Conversion, USA). A matching network and two 108 transformers (HR-Diemen S.A., Spain) were also connected to the amplifier output to increase 109 the output voltage up to approximately 20 kV peak-to-peak. The incident power in the plasma 110 111 reactor was kept constant at 30 W.

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113

114 Fig.1 Experimental atmospheric plasma set-up scheme

115 Cotton fabrics $(3 \times 3 \text{ cm}^2, \sim 0.16 \text{ gr})$ were introduced into the plasma reactor by placing these

- on the lower dielectric plate and 100 μ l of the impregnation liquid (H₂O, D₂O or EtOH) was
- 117 placed onto the cotton substrates (see Figure 1). The liquid weight with respect to the weight

of the cotton samples used in the experiments is equal to 0.5-0.7 gr liquid/1 gr cotton. Since 118 the water absorption capacity of bleached cotton is ~ 2 gr water/1 gr cotton, plasma treatment 119 was performed immediately after liquid impregnation in order to form a thin film of liquid on 120 the cotton substrates before equilibrium of the liquid absorption capacity is reached (Hsieh et 121 al. 1996). Subsequently, a plasma treatment in helium was carried out for a fixed period of 122 123 300 s. A control sample was also prepared by performing the helium plasma treatment step on a cotton sample without any impregnation. In addition, a single cycle of consecutive cotton 124 impregnation and posterior He plasma treatment (300 s) was also repeated 3 and 6 times. 125

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127 OES analysis

OES analysis was used to characterize the plasma active species inside the He discharge during surface modification. Diagnostics of the plasma gas was carried out with a quartz optic fiber connected to a Black Comet spectrometer (Stellarnet, USA) with concave gratings. Spectra were recorded in the UV–VIS wavelength range 190–850 nm (spectral resolution of 0.5 nm) with an integration time of 4 s and a solid angle of ~0.9 steradians. No UV emission was measured below \approx 280 nm, since the DBD plasma reactor is made of borosilicate glass. The exact location of the quartz optic fiber in the DBD reactor can be seen in Figure 1.

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136 Attenuated Total Reflectance FTIR (ATR-FTIR) measurements

137 ATR-FTIR spectra of untreated and plasma-modified cotton fabrics were collected using a Nicolet Avatar 360 spectrometer equipped with a Smart iTR ATR sampling accessory 138 (Thermo Scientific, USA). Spectra were obtained with an average of 32 scans using a 139 resolution of 4 cm⁻¹. An advanced ATR correction algorithm (OMNIC 7.3 from Thermo 140 Electron Corporation) was used to correct for band intensity distortion, peaks shifts and 141 polarization effects. Corrected ATR-FTIR spectra were found highly comparable to their 142 transmission equivalents (Molina et al. 2013b). The background signal was extracted by 143 means of linear backgrounds performed between 4000-3665 cm⁻¹, 3665-2987 cm⁻¹, 2987-2634 144 cm⁻¹, 2634-1800 cm⁻¹, 1800-1492 cm⁻¹, 1492-1186 cm⁻¹ and 1186-830 cm⁻¹. Spectra were 145 normalized to the C-O stretching vibration peak of cellulose located at 1031 cm⁻¹. 146

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148 **XPS analysis**

The surface chemical composition of the untreated and plasma-modified cotton fabrics was 149 analyzed using a PHI 5000 Versaprobe II XPS device equipped with an Al K_{α} X-ray source 150 (hv = 1486.6 eV) operated at 25 W. XPS analysis was conducted on samples, which have 151 been placed in the XPS introduction machine, within 1 month after plasma modification. All 152 measurements were conducted in a vacuum of at least 10⁻⁶ Pa and the photoelectrons were 153 detected with a hemispherical analyzer positioned at an angle of 45° with respect to the 154 normal of the sample surface. Survey scans and individual high resolution C1s spectra were 155 recorded with a pass energy of 187.85 eV (eV step = 0.8 eV) and 23.5 eV (eV step = 0.1 eV) 156 157 respectively. Binding energies were referenced to the C1s photoelectron peak position for C-O species at 286.5 eV (Molina et al. 2017a) or C-C species at 285.0 eV in case of the EtOH 158 impregnated sample (Gorianc et al. 2010). Surface elemental composition was determined 159 after a linear background subtraction from the area of the different photoemission peaks using 160 the corresponding sensitivity factors of the XPS machine (Castle 1984). In this paper, the 161

presented XPS results are the average of 3 different measurement points on a single cottonsample.

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167 SEM imaging

Possible plasma-induced changes in the surface morphology of the cotton samples were recorded using a Hitachi S-3500N SEM device operated at 5 kV. Prior to SEM imaging, a Quorum Q150RS gold sputter coater was used to deposit a gold coating of a thickness of approximately 20 nm on the samples to minimize charging.

172

173 **Results**

174

175 **OES**

176 OES is a very powerful method to identify active species present in a plasma. The identification of said species is of great relevance in this work, as it can provide further 177 insights into the occurring plasma/surface reactions. The UV-VIS emission spectra obtained 178 179 during the He plasma treatment of cotton and cotton impregnated with different liquids (water, D₂O and EtOH) are presented in Figure 2. The emission spectra recorded during the 180 He plasma treatment of the non-impregnated cotton sample (Figure 2a and 2b) present 181 182 different emission lines characteristic for atomic He (501.5 nm, 587.5 nm, 667.8 nm, 706.5 nm, 728.1 nm). In addition, emission bands due to N_2^+ (391.4 nm, 427.8 nm, 470 nm), N_2 183 (315.9 nm, 337.1 nm, 357.6 nm, 380.4 nm), OH radicals (OH: 308.9 nm, 287 nm) and 184 emission lines due to H atoms (H_{α} line at 656.3 nm and H_{β} line at 486.1 nm) can also be 185 observed in the OES spectra. The presence of these species can be mainly attributed to 186 residual humid air remaining in the plasma reactor. The observed plasma species are in 187 excellent agreement with earlier published OES results of helium discharges operating at 188 atmospheric pressure (Asandulesa et al. 2010; Bierstedt et al. 2015). It is also important to 189 mention that during He plasma treatment of the non-impregnated cotton sample the plasma is 190 operating in a diffuse regime independent of the used plasma treatment time. 191



Fig.2 UV-VIS emission spectra obtained during the He plasma treatment of cotton and cotton impregnated with
 different liquids (H₂O, D₂O and EtOH) at 30 (a) and 300 s (b)

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Figure 2 does however reveals small differences in the OES spectra as a function of plasma 197 treatment time for the He plasma treatment: at long plasma treatment time (300 s), an 198 emission line due to atomic oxygen (777.4 nm) appears which might suggest that possible 199 water evaporation from the non-impregnated cotton substrates occurs at longer plasma 200 treatment times. At the short plasma treatment time (30 s), the emission spectrum during the 201 He plasma treatment of cotton impregnated with water is mainly composed of emission 202 lines/bands corresponding to He, H and OH, the latter 2 mainly originating from the 203 dissociation of H₂O used for the impregnation of the cotton samples. Furthermore, in contrast 204 to the OES spectrum obtained during the plasma treatment of pure cotton, only a single, very 205 small band attributed to N2⁺ species could be distinguished. However, at long plasma 206 207 treatment times (300 s) (Figure 2b), the emission spectrum during the He plasma treatment of cotton impregnated with water again resembles that of pure He, indicating that within 300 s, 208 most of the impregnated water had been evaporated from the cotton substrate. 209

Emission spectra recorded during the plasma treatment of cotton impregnated with D₂O 210 resemble these of cotton impregnated with H₂O. Emission bands due to N₂ and N₂⁺ species 211 are however more pronounced in case of the D₂O impregnated sample when compared to the 212 H₂O impregnated cotton. In addition, a small additional OD[•] emission band at 287 nm could 213 also be observed resulting from the dissociation of D₂O used to impregnate the cotton sample. 214 215 It is also important to mention that when cotton is impregnated with H_2O or D_2O , two different plasma regimes can be observed as a function of plasma treatment time which can be 216 clearly differentiated in the OES spectra by examining the intensity of the He line at 706.5 217 nm. As the presence of filaments decreases the overall plasma light emission and thus also the 218 219 intensity of the He line at 706.5 nm compared to the diffuse plasma regime, the following can be concluded: at short treatment times (30 s) in case of the H₂O and D₂O impregnated 220 221 samples, the plasma is operating in the filamentary regime due to the presence of high amounts of H₂O or D₂O. In contrast, at longer treatment times (300 s), most of the liquids are 222 evaporated and the plasma again operates in the diffuse mode similar as in case of the He 223 treatment of the non-impregnated sample. 224

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During the He plasma treatment of cotton impregnated with EtOH, new emission bands 226 227 attributed to carbon species (CN, CH and C₂) could be observed at short plasma treatment times (30 s). These emission bands correspond to the CN violet system (360-390 nm), CH 228 (430, 390 and 314 nm) and the C₂ Swan system $(a^3\prod_u -d^3\prod_g)$ consisting of various vibrational 229 transitions (450–470 nm (Δv =-1), 480–520 nm (Δv =0) and 520–570 nm (Δv =1)). This is in 230 accordance with literature and could be attributed to the dissociation of EtOH used for 231 impregnation (Ikegami et al. 2004). In case of EtOH impregnation, emission bands due to 232 carbon and nitrogen species resulting from ethanol dissociation could still be clearly 233 234 distinguished after 300 s, whereas the atomic oxygen line was no longer recorded. These results provide a strong indication that the ethanol reforming had not yet been completed after 235 236 300 s (Yanguas-Gil et al. 2004).

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238 ATR-FTIR

Besides investigating the plasma gas phase, the effects of impregnation with different liquids (H_2O , D_2O and EtOH) and posterior He plasma treatment (300 s) on the chemical composition of the cotton substrates were also investigated by means of ATR-FTIR.



Fig.3 ATR-FTIR spectra obtained on non-impregnated (a) and H_2O (b), D_2O (c) and EtOH (d) impregnated cotton samples after 1, 3 and 6 cycles of consecutive impregnation/300 s He plasma treatment. UT stands for the untreated cotton sample. Enlarged carboxylate region (1800-1500 cm⁻¹) obtained for cotton samples impregnated with H_2O (e) and EtOH (f)

First of all, the ATR-FTIR spectra of untreated and He plasma-treated cotton samples are 250 presented in Figure 3A. The FTIR spectrum of untreated cotton shows multiple bands 251 characteristic for cellulose (Chung et al. 2004; Garside and Wyeth 2003): a very broad OH 252 stretching peak in the region 3750-2995 cm⁻¹, a broad peak in the region 3000-2800 cm⁻¹ 253 attributed to CH stretching vibrations, a smaller but rather broad peak at ≈ 1635 cm⁻¹ which is 254 usually assigned to the OH bending mode of adsorbed water, a peak at 1430 cm⁻¹ due to CH 255 and C-OH bending vibrations, a broad peak in the region 1400-1300 cm⁻¹ due to CH and CH₂ 256 bending vibrations, a peak in the region 1160-1108 cm⁻¹ due to C-O-C stretching and a peak 257 in the region 1030-998 cm⁻¹ due to C-O stretching vibrations. Figure 3A reveals no significant 258

differences in the ATR-FTIR spectra of non-impregnated cotton samples after performing multiple consecutive He plasma treatments. This is in excellent agreement with earlier published results showing no differences in ATR-FTIR spectra of polymer films as a result of plasma surface treatment (Morent et al. 2008). Indeed, as plasma active species are only able to interact with the top few nanometers of a material, the penetration depth of ATR-FTIR is too large to detect the surface chemical changes induced by plasma treatment (Morent et al. 2008).

The ATR-FTIR spectra of cotton samples undergoing consecutive water impregnation and He 267 268 plasma treatments are shown in Figure 3B. In contrast to the non-impregnated cotton sample, differences in ATR-FTIR spectra could be observed between the untreated and plasma-269 270 modified samples: with an increasing number of impregnation/plasma treatment cycles, the broad peak at 1635 cm⁻¹ gradually increases. A more detailed investigation of this 271 272 wavenumber region (see Figure 3E) indicates that a new band attributed to carboxylate (O-C=O⁻) stretching vibrations appears at 1720 cm⁻¹ which gains in intensity as a function of 273 treatment cycles. In addition, figure 3E also reveals that the peak at 1635 cm⁻¹ also increases 274 as a function of impregnation/plasma treatment cycles. As the broad OH band in the region 275 276 3750-2295 cm⁻¹ decreases with plasma treatment cycles, this increase at 1635 cm⁻¹ could thus not be attributed to higher amounts of adsorbed water, but is most likely the result of the 277 generation of carbonyl groups (C=O) as a consequence of the plasma-induced cotton 278 oxidation process. This seems to be corroborated by the ATR-FTIR spectrum corresponding 279 to untreated cotton immersed in D_2O for 24 hours and dried at room temperature (sample UT 280 D₂O). The UT D₂O ATR-FTIR spectrum highly resembles the ATR-FTIR spectrum of 281 untreated cotton (see Figure 3B and E), but a new band in the region 2650-2250 cm⁻¹ 282 283 associated to the OD stretching peak of D₂O can be observed in the former sample (see Figure 3B). It could therefore be concluded that the intensity of the peak at 1635 cm^{-1} should not only 284 be attributed to the OH bending mode of adsorbed water but the possible signal formation of 285 286 carbonyl groups should also be taking into account. These carbonyl groups could be formed in 287 the untreated cotton samples as a consequence of the bleaching process carried out to the cotton prior to its use in this work. 288

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290 Similar trends are observed when D₂O is used as impregnation liquid (Figure 3C). The ATR-FTIR spectra also show an increasing peak at 1635 cm⁻¹ and at 1720 cm⁻¹, proving the 291 formation of carbonyl and carboxylate groups on the plasma-modified cotton samples. Similar 292 as for the water impregnated samples, both peaks also increase with increasing treatment 293 cycles, suggesting a higher incorporation of carbonyl and carboxylate groups with increasing 294 295 impregnation/plasma treatment cycles. However, because of the different position of the OH stretching peak of H₂O (3750-2995 cm⁻¹) and the OD stretching peak of D₂O (2650-2250 cm⁻¹) 296 ¹), a new band corresponding to the absorbed D_2O in the cotton substrate also appears. 297 Additionally, the H-O-H bending band of water (1638 cm⁻¹) also shifts towards lower 298 wavenumbers for D-O-D (1205 cm⁻¹) (Belhadj et al. 2015; Liu et al. 2016). Nonetheless, the 299 intensity of this band is typically much lower compared to the OD stretching band, making it 300 difficult to clearly differentiate the D-O-D peak in the presented ATR-FTIR spectra. 301 302 Additionally, it also seems that there is a correlation between the decrease in the band associated to the OH stretching peak (3750-2995 cm⁻¹) of cellulose and the increase in the 303 carbonyl and carboxylate groups as a consequence of a plasma-induced oxidation process and 304 an interchange of OH groups present in the cotton substrates with OD groups. Therefore, 305 when replacing H₂O by D₂O, it is again suggested that the increased peak at 1635 cm⁻¹ cannot 306 be attributed solely to adsorbed D₂O and should thus be partially attributed to carbonyl groups 307 formed because of cellulose oxidation (C=O). Comparing the ATR-FTIR spectra of He 308

plasma-treated cotton with and without liquid impregnation, it seems that the presence of H_2O or D_2O favors the oxidation of cellulose because of the diffusion and formation of oxygen and nitrogen reactive species (ROS, RNS) in the adsorbed liquids (H_2O_2 , OH[•], $O_2^{-•}$, ONOO⁻ and NO[•]) (Brisset and Pawlat 2016; Gorbanev et al. 2016; Ikawa et al. 2010; Tani et al. 2012).

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314 The ATR-FTIR spectra corresponding to consecutive ethanol (EtOH) impregnation and He plasma treatments of cotton substrates (up to 6 impregnation/plasma treatments) are depicted 315 in Figure 3D. In this particular case, a small increase in absorbance of the region 316 corresponding to CH stretching modes (3000-2800 cm⁻¹) is observed as a function of 317 impregnation/plasma treatment cycles, suggesting the incorporation of hydrocarbon groups, 318 which is in agreement with the obtained OES spectra showing the presence of excited carbon 319 species in the plasma. Additionally, an increase in the band at 1720 cm⁻¹ is observed whereas 320 the band at 1635 cm⁻¹ only marginally increases (see Figure 3F), suggesting that also in this 321 case, plasma-assisted oxidation processes occur resulting in the incorporation of mainly 322 carboxylate groups. 323

325 **XPS**

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Since the analysis depth of ATR-FTIR is typically between 600 and 1000 nm, crucial 326 information is lost from the uppermost polymer layers, which is particularly important in case 327 of plasma surface modification, as non-thermal plasmas are known to primarily influence the 328 first 50 nm of a material (Holländer et al. 1999; Tatoulian et al. 1995; Wang et al. 2008). In 329 contrast, XPS analyses only the first 3-10 nm of a material surface, making it a very suitable 330 technique to track plasma-induced surface changes (Castle 1984; Cools et al. 2015). 331 Therefore, the changes in the surface chemical composition of the cotton substrates have also 332 333 been studied by XPS in this work.

334

The atomic surface chemical composition of the He plasma-treated cotton samples and the 335 336 cotton samples first impregnated with different liquids (water, D₂O and EtOH) prior to plasma modification are presented in Table 1. Consecutive liquid impregnation and He plasma 337 treatment steps of cotton substrates did not significantly change the surface elemental 338 composition and for this reason, only results of first impregnation and He plasma treatment 339 were mentioned in table 1. This latter reveals that the He plasma treatment of cotton substrates 340 does not significantly change the atomic surface chemical composition with respect to the 341 342 untreated one. This result is quite surprising as plasma-induced surface oxidation was expected, since OH radicals and atomic oxygen were observed in the OES spectra during He 343 plasma treatment (Figure 2). It is also important to mention that the obtained O/C ratio for the 344 untreated cellulose sample is considerably lower than the expected O/C ratio for pure 345 cellulose (O/C = 0.83), which can be attributed to surface non-cellulosic components, which 346 naturally occur on cotton (Kolarova et al. 2013) and which have not been removed from the 347 samples by the performed washing step. Table 1 also reveals that if the cotton sample was 348 349 first impregnated with H₂O, no significant difference in oxygen content could again be observed. On the other hand, a small increase in oxygen content was observable on the D₂O 350 impregnated sample. Moreover, when cotton was impregnated with EtOH, a strong increase 351 in carbon content combined with a large decrease in oxygen content was observed, which was 352 highly unexpected. 353

Table 1. Elemental composition (%) at the surface determined by XPS on cotton treated by He plasma and cotton
 impregnated with different liquids and subsequently treated by He plasma (300 s).

	C (%)	O (%)	O/C
Untreated	61.5±2.4	38.5±2.4	0.62
He plasma	61.8 ± 1.8	38.2±1.8	0.61
H ₂ O/He plasma	60.5±0.9	39.5±0.9	0.65
D ₂ O/He plasma	57.8±0.9	42.2±0.9	0.72
EtOH/He plasma	85.9±1.2	14.1 ± 1.2	0.16

- High-resolution spectra of the C1s envelopes were also recorded to obtain additional insights into the chemical changes induced on the cotton substrates and the fitted C1s peaks are depicted in Figure 4 for a single cycle of impregnation and plasma treatment.



361 Fig.4 XPS high-resolution C1s spectra of untreated cotton, cotton treated with He plasma and cotton

362 impregnated with different liquids and subsequently treated by He plasma

As shown in Figure 4, the C1s spectrum of the untreated cotton sample is deconvoluted into 4 363 distinct peaks, the largest peak being the one at 286.5 ± 0.1 eV, corresponding to aliphatic C-364 O-/C-OH functionalities, as expected for cellulose. Two shoulder bands at higher binding 365 energies are also observed that can be attributed to O-C-O/C=O functionalities (287.7 \pm 0.1 366 eV), which are present in the chemical structure of cellulose and O-C=O functionalities (289.1 367 368 \pm 0.1 eV) most likely originating from the bleaching process carried out on the cotton samples (Molina et al. 2017a; Patino et al. 2011). Moreover, a peak at 285.0 ± 0.1 eV needs to be 369 included as well to obtain a good fit of the C1s peak: this peak probably originates from some 370 non-cellulosic components such as waxes, proteins or pectin, which typically cover natural 371 cotton fibers (Topalovic et al. 2007). Figure 4 also reveals that after performing the different 372 plasma modification procedures, similar XPS fitting peaks as the ones used for the untreated 373 cotton sample can still be applied to deconvolute the high resolution C1s spectra of the 374 plasma-modified samples, although the area below each individual peak seems to vary 375 depending on the performed plasma modification step. From these fitted spectra, relative 376 concentrations of the different carbon bonds present at the surface of the cellulose samples 377 can be determined and these quantitative results are presented in Table 2 for all samples under 378 study using a single cycle of impregnation and plasma treatment. 379

Table 2. Relative concentrations of carbon bonds on untreated cotton, cotton treated with He plasma and cotton
 impregnated with different liquids and subsequently treated by He plasma.

Sample	C-C	C-O-/	C=O/	0-C=0
		С-ОН	0-C-0	
Untreated	41.3 ± 0.3	35.0 ± 0.1	14.6 ± 0.3	9.1 ± 0.5
1x He	39.1 ± 0.5	35.9 ± 0.5	15.6 ± 0.2	9.4 ± 0.5
1x H2O	30.0 ± 2.0	42.9 ± 2.1	16.3 ± 0.7	10.8 ± 0.8
1x D ₂ O	25.3 ± 1.6	45.6 ± 1.6	17.5 ± 0.4	11.6 ± 2.8
1x EtOH	60.4 ± 2.6	23.6 ± 0.5	11.6 ± 1.9	4.4 ± 1.0

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This table clearly reveals that the He plasma treated cotton has a surface chemical 383 composition which closely resembles the surface chemical composition of the untreated 384 cotton sample, which is consistent with the results shown in Table 1. As such, it can thus be 385 386 concluded that the performed helium plasma treatment does not induce a significant surface oxidation, although reactive oxygen plasma species are present in the discharge. Most likely, a 387 combination of functionalization and etching simultaneously occurs leading to an unaffected 388 total oxygen content after He plasma treatment. Table 2 also shows that when performing the 389 390 plasma treatment on the H₂O impregnated sample, a more pronounced increase in oxygencontaining surface functional groups is obtained: a considerable increase in the amount of C-391 O, C=O/O-C-O and O-C=O groups can be seen upon plasma treatment. This increase in C-O, 392 C=O/O-C-O and O-C=O groups is even more pronounced when the cotton sample is first 393

impregnated in D₂O, which is in agreement with the O/C ratio results given in Table 1. The XPS results in this section thus confirm the incorporation of C=O and O-C=O groups on the cotton surface, which was also seen in the ATR-FTIR spectra shown in the previous section and additionally also reveal the incorporation of C-O functional groups on the cotton surface. Taking into account the considerably large penetration depth of FTIR ($\approx 0.6-1 \mu m$), it can also be concluded that the C=O and O-C=O functional groups are not only incorporated at the surface of the cotton samples, but also in the subsurface region.

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When examining the samples impregnated with ethanol prior to plasma treatment, a completely different trend can be observed: a strong decrease in the oxygen-containing functional groups occurs in combination with a large increase in the amount of C-C groups. This peculiar result suggests the formation of a thin oxygenated hydrocarbon coating on the cotton sample, which will be further confirmed by SEM imaging performed in the following section.

- 408
- 409 **SEM**

410 Possible surface damage induced by the performed plasma treatments was also investigated

- 411 by SEM and the obtained SEM images are shown in Figure 5.
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415 Fig.5 SEM images of cotton treated with He plasma and cotton impregnated with different liquids and

416 subsequently treated by He plasma

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418 A single He plasma treatment is found to have a minimal impact on the surface morphology 419 of the cotton sample, as a similar physical appearance compared to the untreated cotton was recorded (not shown for reader clarity). In contrast, a more profound impact on the surface 420 421 integrity is observed with increasing repetitions of He plasma treatments. After 6 He plasma treatments, the cotton surface exhibits some surface damage in the form of microcraters due to 422 plasma-induced etching processes (Karahan and Özdoğan 2008). Similar morphological 423 424 changes have been reported when performing corona plasma treatments of cotton (Mihailović et al. 2011). 425

426

427 On the other hand, when cotton is first impregnated either with H_2O or D_2O and subsequently

- 428 exposed to the He plasma, profound topographic changes could be observed even after one 429 treatment cycle. In addition, by increasing the number of impregnation/plasma treatments, the
- treatment cycle. In addition, by increasing the number of impregnation/plasma treatments, theobserved surface damage significantly increases and after 6 impregnation/plasma treatment

cycles, structures similar to particles appear on the cotton fibers, proving the excessive 431 damage to the cotton fibers because of the high plasma exposure times. It is suggested that the 432 433 presence of H₂O or D₂O on the cotton surface may induce many streamers/filaments during plasma treatment which could in turn be responsible for the excessive damage promoted on 434 the cotton surface when compared to the He treatment without impregnation. This hypothesis 435 436 was further confirmed by the OES results, which revealed that the He plasma on the H₂O and D₂O impregnated cotton samples operated in the filamentary mode at treatment times below 437 438 300 s.

439

When EtOH is used as impregnation liquid prior to the He plasma treatment, it could be 440 observed that the cotton surface is covered with a flat coating, which is in agreement with the 441 conclusions drawn from XPS analysis. In the case of ethanol impregnation, a plasma 442 deposition process thus takes place instead of a plasma activation process. The film-forming 443 behavior of EtOH-enriched plasmas has already been described before (Van Deynse et al. 444 2017), thereby further confirming the results obtained in this work. It is also important to 445 mention that the presence of this film does not produce any significant visual color change of 446 the cotton substrates. 447

448

449 **Conclusions**

450 In this study, cotton fibers were impregnated with H₂O, D₂O or EtOH prior to an atmospheric pressure He plasma treatment in a parallel-plate DBD set-up. OES analysis showed an 451 increase in H atoms and OH radicals during the plasma treatment of H₂O impregnated cotton 452 samples, as was to be expected. Replacing water with its deuterated counterpart resulted in 453 higher concentrations of N metastable species, which showed the indirect influence of free H 454 species in the plasma. Ethanol plasma induced the formation of a variety of CN, C and CO 455 species, even after longer treatment times, indicating that the presence of carbon induced a 456 completely different plasma chemistry. Chemical analysis of the cotton samples revealed that 457 the impregnation with (deuterated) water resulted in a more efficient plasma-induced 458 oxidation of the cotton compared to the dry plasma treatment, while the presence of ethanol 459 led to the formation of a thin oxygenated hydrocarbon film on top of the cotton sample. SEM 460 analysis showed the profound etching of the cotton fibers as a result of the presence of the 461 (deuterated) water, while for the EtOH impregnated samples, a smooth morphology could be 462 distinguished, further indicating the presence of the thin film. Overall, it can be concluded that 463 the impregnation of cotton fabrics with a variety of liquids can induce a more efficient surface 464 modification compared to a plasma treatment as such. 465

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608 List of Figure Legends

609 Fig.1 Experimental atmospheric plasma set-up scheme

- Fig.2 UV-VIS emission spectra obtained during the He plasma treatment of cotton and cotton impregnated with
 different liquids (H₂O, D₂O and EtOH) at 30 (a) and 300 s (b)
- **612 Fig.3** ATR-FTIR spectra obtained on non-impregnated (a) and H_2O (b), D_2O (c) and EtOH (d) impregnated **613** cotton samples after 1, 3 and 6 cycles of consecutive impregnation/300 s He plasma treatment. UT stands for the **614** untreated cotton sample. Enlarged carboxylate region (1800-1500 cm⁻¹) obtained for cotton samples impregnated **615** with H_2O (e) and EtOH (f)
- Fig.4 XPS high-resolution C1s spectra of untreated cotton, cotton treated with He plasma and cotton
 impregnated with different liquids and subsequently treated by He plasma
- 618 Fig.5 SEM images of cotton treated with He plasma and cotton impregnated with different liquids and
- 619 subsequently treated by He plasma
- 620

621 List of Table Titles

- Table 1. Elemental composition (%) at the surface determined by XPS on cotton treated by He plasma and cotton
- 623 impregnated with different liquids and subsequently treated by He plasma (300 s).
- Table 2. Relative concentrations of carbon bonds on untreated cotton, cotton treated with He plasma and cottonimpregnated with different liquids and subsequently treated by He plasma.