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Additional Information

- **1** Monitoring Molecular Dynamics of Bacterial Cellulose Composites
- 2 Reinforced with Graphene Oxide by Carboxymethyl Cellulose
- 3 Addition
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18 Broadband Dielectric Relaxation Spectroscopy was performed to study the molecular

19 dynamics of dried Bacterial Cellulose/Carboxymethyl Cellulose-Graphene Oxide

- 20 (BC/CMC-GO) composites as a function of the concentration of CMC in the culture
- 21 media. At low temperature the dielectric spectra are dominated by a dipolar process
- 22 labelled as a β -relaxation, whereas electrode polarization and the contribution of dc-
- conductivity dominate the spectra at high temperatures and low frequency. The CMC
- 24 concentration affects the morphological structure of cellulose and subsequently alters its
- 25 physical properties. X-ray diffractometry measurements show that increasing the
- 26 concentration of CMC promotes a decrease of the I_{α}/I_{β} ratio. This structural change in
- BC, that involves a variation in inter- and intramolecular interactions (hydrogen-
- bonding interactions), affects steeply their molecular dynamics. So, an increase of CMC
- 29 concentration produces a significantly decrease of the β -relaxation strength and an
- 30 increase of the dc-conductivity.
- 31

32 **1. Introduction**

- 33 New materials consisting of a biopolymer matrix reinforced with carbon
- 34 derivatives (graphene and carbon nanotubes) offer interesting properties such as
- biodegradability and non-toxicity (Fan et al., 2010; Liu et al., 2010; Ye et al., 2016).
- 36 In recent years, cellulose has been employed in various nanotechnology
- applications such as filtration processes or building up matrixes for
- microelectronic purposes (Terzopoulou et al., 2015; Miyauchi et al., 2010; Han et al.,

2011;Xu et al., 2015; Feng et al., 2012). Few studies have been addressed to study
the conductivity and energy storage of these materials but all of them conclude
that there is great potential for their application in electronics development (Liu et al., 2010; Luong et al., 2011).

43 Cellulose is a promising alternative to the omnipresent oil-based polymers. It is produced by a diverse array of organisms: plants, bacteria, algae and even 44 some animal species (Dayal & Catchmark, 2016; Mohite & Patil, 2016). The 45 crystalline structure of cellulose I (native cellulose) is a mixture of two distinct 46 crystalline forms: cellulose I_{α} (triclinic) and I_{β} (monoclinic) (Atalla & Vanderhart, 47 1984). The main difference between the I_{α} and I_{β} forms is the relative 48 displacement of cellulose sheets along their "hydrogen-bonded" planes in the 49 direction of the chain's axis. The directionality of the 1-4 linkage along the length 50 of the cellulose chain affects how neighboring chains interact with each other, 51 52 and thus its physical properties. The I_{α}/I_{β} ratio have been found to vary between samples from different origins (O'Sullivan, 1997; Nishiyama, 2009). The presence 53 of dimorphism makes it difficult to interpret XRD measurements as the positions 54 of peaks from both crystalline forms are very close to each other. However, the 55 56 cellulose sample mostly contains the crystalline form, I_{α} when the intensity of the 14.5° peak is larger than that of the 16.6° peak (Lee et al., 2015; Tokoh et al., 1998). 57

Some bacteria from the genus *Gluconacetobacter* synthetize cellulose. This 58 bacterial cellulose (BC) is formed by cellulose fibers (around 110 nm in 59 diameter) that growths as a gel when cultivated in static conditions (Hirai et al., 60 61 1997). The BC gel is formed by a 3D coherent network of BC fibers that provides 62 remarkable mechanical properties, high surface area and porosity (Iguchi et al., 2000; UI-Islam et al., 2012). BC-based composites have been used for a variety of 63 applications including: membranes, strong paper, and biomaterials for biomedical 64 applications (Nakagaito et al., 2005; Kim et al., 2011a; Lin et al., 2013). 65

66 Several works have reported the potential applications of BC-based

67 composites in the electronic industry (Kafy et al., 2015; Sadasivuni et al., 2014).

68 Graphene oxide (GO) is among the materials used to provide cellulose-based

69 composites with novel properties (Shao et al, 2016). GO consists of a single-layer

of graphite oxide with hydroxyl and epoxy groups on the basal plane. These

71 groups affect its electronic, mechanical, and electrochemical properties (Gomez-

Navarro et al., 2010). The facile synthesis and substantial solubility of GO together

with its moderate conductivity and high surface area (Chen et al., 2012) make it a

promising material for electronic applications. Kafy (Kafy et al., 2015) prepared

cellulose-GO nanocomposites by incorporating graphene oxide (GO) platelets

76 modified with hexamethylene diisocyanate as grafting agent. The energy storage

77 capabilities of the nanocomposites were assessed trough the dielectric response of

78 the samples. Their results showed that the dielectric constant (ϵ') values increases

79 with the GO content.

80 The incorporation of GO in a cellulose matrix is usually achieved by

- dissolving or disrupting the cellulose network to add GO platelets into a cellulose
- solution (Kim et al., 2011b; Tian et al. 2014; Liu et al., 2015). In the work presented
- 83 here, BC/GO composites were prepared using a bottom-up technique that
- 84 preserves the 3D network structure of BC. This technique requires the
- 85 modification of the BC culture medium in order to allow cellulose fibers to grow
- in the presence of a second phase. The viscosity of the medium must be tuned to
- 87 prevent the precipitation of the second phase. Several polysaccharides including
- 88 pectin, sodium alginate, and corn steep liqueur have been used to this end (Dayal
- 89 & Catchmark, 2016; Gu & Catchmark, 2012).
- For this investigation, we have used carboxymethylcellulose (CMC), a watersoluble semi-synthetic derivative of cellulose (Yadav et al., 2013), to increase the
- 92 viscosity of the culture medium and prevent the precipitation of GO platelets. GO
- 93 platelets are hydrophilic and can be dispersed in the culture medium. Several
- 94 studies have suggested that the hydrophilic functional groups on the surface of
- 95 GO may form hydrogen bonds with the -OH groups on BC (Si et al., 2014; Feng
- 96 et al., 2012; Shao et al., 2015). Previous studies show that CMC can be
- 97 successfully used to suspend hydroxyapatite (HA) and starch to prepare
- 98 BC/starch and BC/HA composites (Grande et al., 2009b; Grande et al., 2009a).
- 99 CMC has highly hydrophilic properties and is non-toxic, renewable,
- 100 biocompatible and biodegradable (Kamarudin & Isa 2013). Several studies report
- 101 that the addition of CMC into the BC culture media promotes a reduction on the
- diameter of BC fibers (Chen et al., 2011; Huang et al., 2010; Hirai et al., 1997).

103 Dielectric Relaxational Spectroscopy (DRS) is one of the most versatile 104 techniques used to examine the dynamics of polymers and small molecules (Kremer et al., 2003; Riande et al., 2004). A detailed dielectric analysis of various 105 106 cellulose and derived materials has been reported by Einfeldt (Einfeldt et al., 2001a; 2002 and 2004). Due to the presence of polar molecular structures within 107 these materials, the dielectric spectrum shows several relaxation processes. 108 Studying dried samples, two processes with secondary relaxation character 109 appear in the sub- T_g region labelled as β - and γ -relaxations. The β -relaxation is 110 dominant and related to the local backbone or segmental motion of the chain. The 111 γ -relaxation is associated with the motion of the side groups of the repeating unit. 112 The alleged primary or α -relaxation, directly connected to the glass transition, is 113 absent from cellulosic material with low moisture content. Moreover, for wet 114 cellulosic materials a relaxation labelled as β_{wet} is observed in the room 115 116 temperature range (Einfeldt et al., 2001b; Gelin et al., 2007). This process may be associated with the collective motion of solvent-swollen cellulose regions. There 117 is an additional process in the high temperature range, corresponding to the dc-118 119 conductivity of the sample. This relaxation is unrelated to polymer motion and is 120 characteristic of all disordered solids with ionic conductivity.

Because of the potential interest of BC derivative materials, we have focusedthis paper on the analysis of the CMC content effect on the molecular dynamics

- 123 of BC/CMC/GO composite samples. The BC, used in this study, was produced
- 124 by the bacterium *Gluconacetobacter Xylinus*. Values of CMC were varied to
- evaluate their influence on thermal stability and on dynamic molecular behavior.
- 126 The data will provide useful information for further development of materials
- 127 derived from polysaccharides and biopolymers to be used in the electronics
- 128 industry.

129 2. Experimental

130 *2.1. Materials*

131 All chemicals were of analytical grade and used without purification.

- 132 Carboxymethyl-cellulose sodium salt (CMC) was procured from Acros Organics
- 133 (average MW 25,0000, DS = 1.2). Graphene oxide (GO) platelets of 0.7-1.2 nm
- in thickness were purchased from Cheep Tubes Inc. (cat. no SKU: 060101, USA).

135 2.2. BC production

The static culture medium for the growth of BC consisted of 1.0% (w/v) Dglucose, 1.5% (w/v) peptone, 0.8% (w/v) yeast extract and 0.3% (w/v) glacial acetic acid, adjusted to 3.5 pH with hydrochloric acid. The culture medium was autoclaved at 121°C for 20 min and then allowed to reach room temperature. Ethanol was added after sterilization to improve the growth of the cellulose gel. The strain *Gluconacetobacter Xylinus* was inoculated and cultivated at 30°C for 21 days at room temperature.

143 2.3 BC/GO/CMC composite preparation

The viscosity of the culture medium was increased using carboxymethyl 144 145 cellulose (CMC), in order to avoid the precipitation of GO platelets. CMC was added to the culture medium at concentrations of 1.0%, 2.0% and 3.0% (w/v) and 146 147 stirring was carried out until CMC was completely dissolved. GO was added in concentrations of 0.01% (w/v) (Table 1). BC/CMC/GO gels were removed, 148 washed with water, boiled in 1.0 M NaOH at 70 °C for 90 min and rinsed in 149 deionized water. Samples were pressed in a hydraulic plate press at 105 °C to 150 151 produce sheets.

152

153 **Table 1.** Sample code, GO and CMC content, char residues and temperature of 154 maximum degradation, $T_{max deg}$.

155			-		
156	Code	GO-content (% w/v)	CMC-content (% w/	v) Char residues (%) $T_{max deg}$ (°C)
157	BC	0	0	13	377.7
158	1CMC/C	GO 0.01	1	19	363.8
159	2CMC/C	GO 0.01	2	20	362.7
160	3CMC/C	GO 0.01	3	20	352.5
161					

162 2.3. Morphological Characterization

163 Both scanning electron microscopy (SEM) and Atomic Force Microscopy (AFM) were employed to characterize the morphology of the samples. SEM was 164 carried out in a Hitachi S-4800 microscope at an accelerating voltage of 20 kV 165 166 and at working distance of 14 mm. Small pieces of sample was placed in the sample holder (2 inch). The samples were metalized with an Au-Pd coating 167 before observation. Images of the surfaces of BC and BC/CMC/GO samples were 168 carried out using AFM (easyScan 2, Nanosurf AG, Switzerland) in the tapping 169 mode. A cantilever with a nominal spring constant of 48 N/m, resonance 170

171 frequency of 190 kHz and a tip radius lower than 10 nm was used.

172 2.4. Thermal Gravimetric Analysis (TGA) Characterization

TGA was performed using a Perkin-Elmer TGA4000 thermogravimetric
analyzer. Samples weighing approximately 10-11 mg were analyzed at a heating
rate of 10°C/min from 30° to 700°C under a nitrogen flow of 20 mL/min.

176 2.5. Dielectric Relaxation Spectroscopy (DRS) Characterization

177 The isothermal relaxation spectra of BC/CMC/GO samples were collected with a Novocontrol Broadband Dielectric Spectrometer (Hundsagen, Germany) 178 consisting of an Alpha Analyzer to carry out measurements from 5×10^{-2} to 3×10^{6} 179 Hz. The measurements were performed in an inert N₂ atmosphere from 150°C to 180 -150°C. The temperature was controlled with a nitrogen jet (QUATRO from 181 Novocontrol), with a temperature error of ± 0.1 K during each sweep in frequency. 182 183 Molded disc shaped samples of about 0.12 mm thickness with diameters 10 and 184 20 mm were used. The thickness of each sample was measured with a micrometer screw. In order to avoid an increase of conductivity due to water, the samples 185 were initially dried at 40°C in a vacuum oven until constant weight was reached. 186 In order to minimize the effect of moisture that may have been introduced during 187 transport, the samples were heated at 150 °C and measurements were carried out 188 from high to low temperatures. The experimental uncertainty was less than 5% in 189 all cases. 190

191 From the several models that have been employed to characterize the192 isothermal relaxation spectra, the more extensively used is the empirical

193 Havriliak-Negami (Havriliak et al., 1966 and 1967) model (HN):

194
$$\varepsilon^{*}(\omega) = \varepsilon_{\infty} + \frac{\varepsilon_{0} - \varepsilon_{\infty}}{\left[1 + \left(j\omega\tau_{HN}\right)^{a_{HN}}\right]^{b_{HN}}}$$
(1)

195 where ε_0 is the relaxed permittivity, ε_∞ is the unrelaxed permittivity, $\Delta \varepsilon = \varepsilon_0 - \varepsilon_\infty$ is 196 the strength relaxation, the parameters a_{HN} and b_{HN} [0< (1- a_{HN}), (1- a_{HN}) b_{HN} <1] 197 define the symmetrical and asymmetrical broadening of the loss peak

- 198 respectively, and τ_{HN} is the characteristic relaxation time. The temperature
- 199 dependence of the last parameter follows the Arrhenius (ARRH) or Vogel-
- 200 Fulcher-Tammann-Hesse equations (VFTH) depending of the relaxation nature.
- 201 According to the Onsager–Fröhlich–Kirkwood (OFK) equation (Fröhlich,
- 202 1959), the $\Delta \varepsilon_{\beta}$ is related to the reorientation of N_p dipoles per unit volume over all spatial orientations as follows:

204
$$\Delta \varepsilon_{\beta} = \frac{\varepsilon_0 \left(2 + \varepsilon_{\infty}\right)^2}{\left(2\varepsilon_0 + \varepsilon_{\infty}\right)} \frac{4\pi N_p \mu_0^2}{9k_B T} g_{\beta}$$
(2)

where μ_0^2 is the uncorrelated mean-square dipolar moment, k_B is the Boltzmann constant, N_p is the number of dipoles participating in the process and g_β is a factor that accounts for the intramolecular and intermolecular correlations. The term $N_p \mu_0^2 g_\beta$ depends on the dipolar reorientation of *N* identical molecules per volume unit, in which each individual *i* has an instantaneous conformation associated with a dipole moment.

211 2.6. X-ray Diffraction (XRD) Characterization

Wide angle XRD measurements were carried with a Bruker AXS D5005 diffractometer. The samples were scanned at 4°/min using Cu K α radiation (λ =0.15418 nm) at a filament voltage of 40 kV and a current of 20 mA. The diffraction scans were collected at room temperature over the 2 θ range of 5°-80° with a 2 θ step of 0.01°.

217 Peak separations of the profiles were carried out using the non-linear least 218 squares fitting program, where a pseudo-Voigt function was used. The *d*-spacings were calculated using the Bragg's equation. The crystallinity index (CrI) 219 quantifies the relative amount of crystalline material in cellulose (Park et al. 2010). 220 The CrI value varies significantly, depending on the evaluation method 221 (Ahvenainen et al., 2016). Segal et al. (Segal et al., 1962) estimate the CrI by means 222 of a (I_{crys} - I_{am})/ I_{crys} relation, where I_{crys} is the intensity of the peak at 20 around 223 22.5° and I_{am} is the intensity of the baseline at 20 around 18°. This method 224 usually provides exceedingly high CrI values (Park et al. 2010; Terinte et al., 2011; 225 Bansal et al., 2010). CrI can also be estimated from the ratio of the separated 226 crystalline peak area to the total reflection area, including the background 227 228 (Hermans et al., 1949; Wada et al., 2001).

229 To calculate crystal size, the Scherrer equation (Alexander, 1979) has been used

230
$$L = \frac{K \cdot \lambda}{FWHM \cdot \cos \theta}$$
(3)

where L is the crystallite size perpendicular to the plane, K is 0.94, λ is the X-ray

232 wavelength, *FWHM* is the full-width at haft-maximum in radians and θ is the

233 Bragg angle.

234 **3. Results and discussion**

235 3.1. Morphological Characterization

The morphology of pure BC has been examined to compare with that of 236 BC/CMC/GO composites with an aim to understand the effects of GO 237 reinforcement and CMC content on the formation of the BC network. A SEM 238 micrograph of a pure BC surface is shown in Fig. 1a. The image reveals typical 239 fibrils around 100 nm in diameter, which coincides with data for pure BC 240 241 reported previously (Grande et al., 2008) (117.76 nm). Fig. 1b shows the surface of a BC/CMC/GO composite (code 1CMC/GO), which differs significantly from 242 the pure BC surface. The packing density of cellulose microfibrils varies 243 significantly with the CMC addition. The diameter of the fibrils and pore size in 244 245 the composite are smaller than those found in pure BC. White arrows indicate the presence of GO agglomerates in the BC/CMC/GO composite. The formation of 246 agglomerates may be due to low dispersion, probably caused by the solution-247 248 casting method and the high surface area of GO sheets (Ebrahimzadeh et al., 249 2016).

AFM images of pure BC and BC/CMC/GO composites (code 3CMC/GO) are shown in Fig. 1c and 1d. The average diameter of pure BC fibrils is 120.89±10.69 nm, in agreement with the measurements reported by Grande (Grande et al.,

253 2009a) and Bohn (Bohn et al., 2000). In contrast, the average diameter of fibrils
254 grown in the presence of CMC (3%) and GO (0.01%) was 86.72±7.93 nm. The

reduction of the BC fibrils diameter is attributed to the presence of CMC in the

culture medium. This result is in agreement with previous report for

BC/CMC/HAp nanocomposites (Grande et al., 2009a; Huang et al., 2010; Hirai
et al., 1997).

The average pore size in pure BC network meshes was 224.4±68.35 nm. However, for BC/CMC/GO composites it was not possible to determine consistent mesh pore sizes (Fig. 1e) since the pores in several cases appeared to be closed. The presence of GO in the composites is indicated with arrows in Fig. 1f. AFM images and SEM micrographs confirmed the presence of GO in the composites.

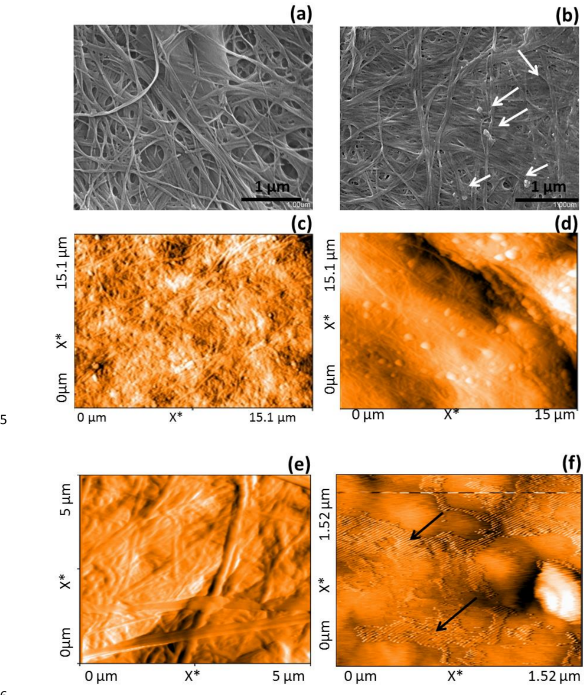




Fig. 1. SEM micrograph of: (a) surface of pure BC and (b) surface of BC/CMC/GO
nanocomposite code 1CMC/GO. Arrows indicate agglomerates of GO within the BC
matrix. AFM images depicting the topography of a representative pure BC film (c),

BC/CMC-GO (sample 3CMC/GO) composite film (d) and (e). Arrows on (f) show GO
agglomerates within BC sheets.

272 *3.2. Thermal Gravimetric Analysis Characterization*

- The thermal stability of the BC/CMC/GO samples was analyzed by TGA. Fig.
- 274 2a shows the TGA spectra obtained for the BC/CMC/GO samples and for CMC.
- Table 1 collects values of the final weight and the temperature of maximum

degradation, $T_{max deg}$, of the samples. Two processes are observed: one at low 276 temperature related to the dehydration process and another at high temperature 277 derived from thermal degradation. The weight loss associated with the first 278 279 process (around 100°C) increases with the increase of CMC concentration. Thus, the CMC sample depicts a 16% weight loss, while a reduction to 5% and 6.5% is 280 281 observed for BC and for the composite samples, respectively. The changes 282 observed as a result of the CMC content appear to be linked to the hydrogen bonds present in the cellulose structure (Watanabe et al., 2006 and 2007; Poletto et 283 al., 2013). In addition, the formation of hydrogen bonds between water molecules 284 and hydroxyl groups of BC and CMC may lead to an increased retention of such 285 286 water molecules in the BC composites. A second weight loss stage is observed with an onset temperature between 290° and 310°C. This temperature decreases 287 with addition of GO and CMC. Pure CMC has an initial decomposition 288 temperature around 290°C and pure BC around 310°C, so BC/CMC/GO 289 composites depict a thermogravimetric behavior between the raw components 290 291 according to the addition rule (pure GO starts weight loss around 600°C) (Goméz et al. 2013). Fig. 2b shows the DTG curves. Each sample presents a well-defined 292 maximum degradation speed. The $T_{max deg}$ decreases as the CMC content 293 294 increases. Additions of GO and CMC components change the final weight loss 295 from 13% to 19-20% accordingly.

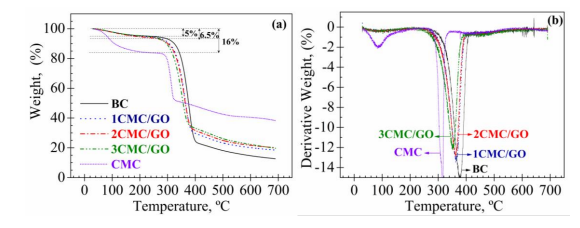




Fig. 2. Curves of weight as function temperature (a) and derivative weight % asfunction of temperature (b) from the TGA tests of BC composites.

299 3.3. Dielectric Relaxation Spectroscopy Characterization

The dielectric spectra of the dried BC/CMC/GO samples show two zones related 300 301 to the microscopic fluctuations of the molecular dipoles (dipolar relaxations) and charges transport across the samples (conductive processes). An isochronal 302 303 representation of the data has been chosen for the sake of clarity. Fig. 3a displays the temperature dependence of the dielectric permittivity (ϵ') and loss permittivity 304 (ε'') of the dried BC/CMC/GO samples at 10^2 Hz. The ε' spectrum presents two 305 zones. At low temperatures a step associated with the dipolar processes is 306 307 displayed, while at higher temperatures a significant increase of the permittivity related to the electrode polarization (EP) effect is observed. Additionally, the ε'' 308

309 spectrum shows different relaxation zones. At a low temperature, an ambiguous 310 process (γ -relaxation) merges as a shoulder of the main process (β -relaxation). Unfortunately, the γ -relaxation falls on the low temperature limit for 311 measurements, making the subsequent analysis difficult. By increasing the order 312 of temperature, the loss isochrones present a dominant absorption labelled as β -313 314 relaxation, which is observed for all samples studied. The intensity of the β relaxation observed between -120°C and -50°C decreases significantly by 315 increasing CMC concentration. As for other dried cellulose materials, the primary 316 α -relaxation directly connected to glass transition was not observed (Einfeldt et 317 al., 2001a; 2001b; 2004). One possible explanation of this fact could be the low 318 amorphous content due to the high crystallinity of the BC. In the second zone, the 319 conductive processes dominate the dielectric response. The processes include the 320 321 migration of mobile charge carriers across the medium (ohmic conduction or 322 electronic conduction) and the trapping of charges at interfaces and boundaries (non-ohmic conduction or polarization effects). 323

For the convenience of further analysis, the dielectric loss was also plotted in the frequency domain. One dominant relaxation (β -peak) between -150° to 0°C appears. For the sake of clarity, the dielectric loss of one isotherm (-90°C) of the five studied samples is shown in Fig. 3b. From this plot, one may imply that the position of the β -peak is scarcely affected by the CMC concentration. However, as CMC concentration increases, the β -peak tends to (i) slightly increase in broadness (see inset Fig. 3b) and (ii) significantly decrease in intensity.

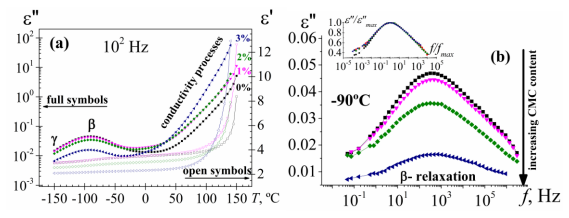
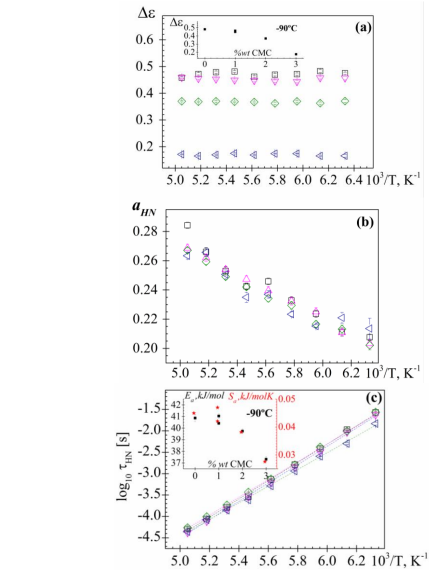


Fig. 3. (a)Temperature dependence of the dielectric permittivity at 10² Hz and (b)
frequency dependence of loss permittivity at -90°C for the BC/CMC/GO analysed
samples. Inset: Normalized plot. (BC: square, 1CMC/GO: triangle down, 2CMC/GO:
diamond, 3CMC/GO triangle left).

331

In order to characterize the β -process observed in the spectra of all analyzed samples, the empirical HN (Havriliak et al., 1966 and 1967) model has been used. The HN fitting parameters were determined at several temperatures from a multiple nonlinear regression analysis of the experimental data. Due to the symmetry of the secondary absorptions, the value of the b_{HN} parameter was fixed to 1 in the fitting procedure.

- The temperature dependence of $\Delta \varepsilon_{\beta}$, displayed in Fig. 4a, shows that in the
- 343 range of temperature analyzed, this parameter is nearly constant. Nevertheless, a
- 344 significant change was observed in the $\Delta \varepsilon_{\beta}$ between samples in which BC was
- incubated in the presence or absence of CMC. By plotting the CMC content
- dependence of $\Delta \epsilon_{\beta}$ obtained at -90°C, an inverse relationship between both
- parameters is observed (inset Fig. 4a), probably due to the fact that CMC
- 348 presence promotes a structural change in the BC as has been reported previously
- 349 (Watanabe et al., 1998; Yamamoto et al. 1989 and 1996). This structural change,
- 350 involves a variation in inter- and intramolecular interactions (hydrogen-bonding
- interactions), and thus, subsequent variation in the molecular mobility of the $\frac{1}{2}$
- composite. Hence, the term $N_p \mu_0^2 g_\beta$ in the Onsager–Fröhlich–Kirkwood (OFK)
- equation (Fröhlich, 1959) that depends on the dipolar reorientation of *N* identical
- 354 molecules per volume unit, probably is reduced by increasing the CMC content.



- **Fig. 4.** Temperature and CMC content dependence of (a) $\Delta \varepsilon_{\beta}$ (inset: CMC)
- content dependence of $\Delta \varepsilon_{\beta}$ at -90°C), (b) a_{HN} and (c) $\tau_{HN\beta}$ (inset: CMC content

355

356

360	dependence of E_a (kJ/mol) [square] and S_a (kJ/(mol·K) [star] at -90°C.) for
361	BC/CMC/GO samples. BC: square, 1CMC/GO: triangle down, 2CMC/GO:
362	diamond, 3CMC/GO triangle left.

The temperature dependence of the width parameter is plotted in Fig. 4b. For all samples, the a_{HN} shape parameter increases linearly with temperature from 0.21 to 0.28. The low value obtained for this parameter indicates that the β relaxation is a very distributed process.

The temperature dependence of τ_{HN} is shown in Fig. 4c. It can be seen that the 367 process for all samples follows ARRH behavior, $\tau_{HN} = \tau_0 \exp[E_a/RT]$. Their 368 activation energies (E_a) and pre-exponential factors (τ_0) are summarized in Table 369 2. Both quantities represent two different effects: (i) energetic effect (E_a) and (ii) 370 entropic effect, $S_a \approx -R \cdot \ln(\tau_0/\tau_D)$, where S_a is the excess entropy of the 371 reorientation process of the dipole system taking place during the dielectric 372 polarization and $\tau_D = h/k_B T_0 = 1.7 \times 10^{-13} s$ is the Debye relaxation time 373 (Starkweather, 1991). With rising CMC concentration, both E_a and S_a values 374 375 slightly decrease (inset Fig. 4c). The activation energies were found to lie between 41 and 37 kJ/mol and the pre-exponential factor between 10^{-15} and 10^{-14} 376 s. The E_a values are similar to those reported for cellulose (46 kJ/mol) and for BC 377 (45 kJ/mol) (Einfeldt et al., 2001a; Grande et al., 2008). 378

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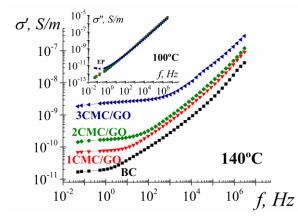
Table 2. Characteristic parameters of Arrhenius fit corresponding to the β and conductive processes.

383	β-process			Conductive process		
384	sample	$\log_{10} \tau_0$	E_a (kJ/mol)	$\ln \sigma_0$	E_a (kJ/mol)	
385	BC	-15.1±0.2	40.9±0.3	8.1±0.8	113.6±2.6	
386	1CMC/GO	-15.2±0.1	41.0±0.2	11.4±0.4	119.9±1.2	
387	2CMC/GO	-14.8±0.1	39.7±0.2	12.8±0.80	118.3±2.5	
388	3CMC/GO	-14.2±0.1	37.3±0.2	13.1±0.6	115.0±1.9	

389

390 With the goal of characterizing the conductive behavior, it is advisable to plot the obtained dielectric data in terms of the complex conductivity related to the 391 complex dielectric permittivity as $\sigma^*(\omega) = i \cdot \omega \cdot e_0 \cdot \varepsilon^*(\omega)$, where e_0 is the vacuum 392 permittivity. As usual, in the frequency domain, the isotherms corresponding to 393 low temperatures of the real component of the conductivity (σ') show a nearly 394 linear dependence with frequency (Jonscher, 1977). However, isotherms 395 corresponding to higher temperatures, display a plateau in the low frequency 396 397 region, reflecting frequency independent conductivity, i.e., dc conductivity (σ_{dc}),

- attributed to resistive conduction through the polymer bulk. The σ_{dc} increases by
- increasing CMC concentration. So, this result indicates that the BC structural
- 400 changes, promoted by the presence of CMC, enhance the conduction through the401 polymer bulk. Fig. 5 clearly conveys this effect, displaying the conductivity
- 402 spectra of the BC/CMC/GO samples analyzed at 140°C. The presence of GO and
- 403 CMC clearly produces a rise in conductivity (Fig. 5). In addition, the EP,
- associated with the accumulation of charges at the electrode-sample interface, is
- 405 enhanced by increased CMC concentration (inset of Fig. 5). This effect can be
- 406 related to the increase of ion charge produced by increasing the CMC sodium salt
- 407 concentration. The temperature dependence of the σ_{dc} , obtained from
- 408 extrapolations at low frequencies, can be described by an ARRH relationship.
- 409 The fit parameters (E_a and σ_0) obtained are summarized in Table 2. A slight
- 410 reduction of the E_a with CMC concentration was once again observed.



412 Fig. 5. Frequency dependence of the σ' for BC/CMC/GO samples at 140°C. Inset: 413 Frequency dependence of the σ'' at 100°C.

414 *3.4. X-ray Diffraction*

The structural order of BC/CMC/GO samples is expected to influence the thermal and dielectric properties. For this reason, it is pertinent to carry out a XRD study of our samples in order to establish correlation between structural, thermal and dielectric properties.

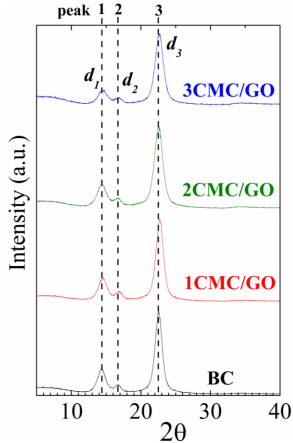
The XRD patterns obtained for BC/CMC/GO composites and pure BC are shown in Fig. 6. The sharp peaks indicate that the BC was semicrystalline, as has previously been reported (Klemm et al., 2005). All patterns show three diffraction peaks at 2 θ near 14.4° (d_1), 16.7° (d_2) and 22.5° (d_3). These peaks can be

- 423 attributed to the (100), (010) and (110) planes of cellulose I_{α} or the (1 $\overline{10}$), (110)
- 424 and (200) of cellulose I_{β} (French, 2014). Analysis of the three diffraction peaks

425 was carried out to determine the *d*-spacing, the crystallite sizes (L), I_{α}/I_{β} , and 426 arrestallite index (CrI) of each pack of the complex studied, as summarized in

- 426 crystallite index (CrI) of each peak of the samples studied, as summarized in427 Table 3.
- 428 The intensity of the d_1 -spacing is higher than the d_2 -spacing, which is 429 characteristic of typical cellulose samples that contain mostly I_{α} phase (Lee et al.,

- 430 2015; Tokoh et al., 1998). Nevertheless, by increasing the CMC content, the
- 431 difference between the intensity of both peaks is reduced. On the other hand, the
- difference between the Bragg angle of peak 1 (d_1) and peak 2 (d_2) , decreases with
- an increased CMC content. This fact suggests that increasing the CMC content
- 434 decreases the cellulose I_{α} phase fraction (Watanabe, et al., 1998; Yamamoto et
- al., 1989 and 1996). Wada et al. (Wada et al., 2003) have proposed a method to
- estimate the I_{β} fraction of the sample from the *d*-spacing (d_1 , d_2). According to
- 437 our results, the mass fraction of cellulose I_{α} is greatly decreased with an
- 438 increasing concentration of CMC. So, the positions of the XRD peaks change
- 439 with CMC concentration due to the varying I_{α}/I_{β} ratio in the samples.
- 440 The CrI values evaluated Segal (Segal et al., 1962) procedure decrease as the 441 CMC concentration increases (Table 3). The $\Delta \varepsilon_{\beta}$ and the CrI values obtained for
- BC samples are in agreement with those reported by Einfeldt et al. (Einfeldt et al., 2004).



445 **Fig. 6**. X-ray diffractometry profiles of BC/CMC/GO samples.

The shape and size of crystallites can affect the obtained CrI values. Cellulose crystallites with a square cross sectional shape should have nearly equal intensities of the d_1 (peak 1) and d_2 (peak 2) spacings (Moon et al., 2011). The slightly stronger d_1 reflection in Fig. 6 indicates that cellulose crystallites are of

450 rectangular shape.

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anaryzeu.					
		BC	1CMC/GO	2CMC/GO	3CMC/GO
<i>d</i> -spacing (nm)	d_1	0.617	0.618	0.615	0.610
	d_2	0.533	0.534	0.533	0.530
	d_3	0.394	0.395	0.395	0.393
L(nm)		6.46	6.13	5.46	5.65
$\left(I_{\alpha}/I_{\beta}\right)^{a}$		_	_	0.98/0.02	0.74/0.26
CrI ^b (%)		96.2	95.8	95.6 94.	8

452 **Table 3.** *d*-spacing, crystallite sizes (*L*), I_{α}/I_{β} , and crystallite index (CrI) of samples 453 analyzed.

461 ^a Wada et al. 2003: $f_{\beta X-Ray} = -70.542 \cdot d_1 + 37.583 \cdot d_2 + 23.360$; ^bCrI Segal

462 The crystal size, L, estimated from peak 3 (d_3) , decreases from 6.46 nm to 5.65 nm as the concentration of CMC increases (Table 3). This harmoniously decrease 463 agrees with the decrease in average microfibril size and the change of packing 464 density obtained from SEM and AFM observations. At the same time it agrees 465 466 with the thermal stability range of the samples obtained by TGA. Thus, according to Poletto et al. (Poletto et al., 2013) higher CrI and L is related to a higher 467 thermal stability, but for the same CrI, the order of the different crystallites can 468 affect the thermal stability (Poletto et al., 2014). Additionally, we should consider 469 470 that further degradation of the CMC component may favor the degradation of the BC component. 471

472 **4.** Conclusions

473 This work presents a combined analysis of the structure and properties of cellulose bacterial composites reinforced by graphene oxide. As a result, the 474 475 crucial role of CMC addition to induce substantial structural changes in Cellulose is evidenced. The fraction of the two allomorphs of I structure (I_{α}/I_{β}) is greatly 476 decreased with increasing concentrations of CMC sodium salt (CMC) in the 477 478 incubation medium. The crystalline size and crystallinity sample index decrease 479 with an increasing concentration of CMC. The sample CrI values, estimated from the ratio of the area of the crystalline peaks to the total area, experiences a 480 reduction of a 13% with respect BC alone. These changes in the sample's CrI 481 482 value and morphology are in good agreement with the results obtained from SEM 483 and AFM.

The proportion of both phases and how they are distributed in the microfibrils determines electrical behavior and therefore, their potential applications. As expected, the presence of GO produces a rise in the conductivity of the BC composites. Additionally, this increase is higher with increasing the CMC content. The dynamic mobility is dramatically affected by the number of hydroxyl groups and their ability to make hydrogen bonds between cellulose 490 chains. Both the β -relaxation and conductivity process are thermally activated 491 processes, and both are steeply dependent on the CMC concentration.

Given the results obtained, we believe that the composites described herein could find various applications in portable and bendable electronics. According to our results the sample with a 3% of CMC has the higher conductivity, but it would be appropriate to perform a future study of samples with higher GO and CMC contents, in order to find the composition that optimizes the electric

- 497 properties.
- 498

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- 504

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