

Unraveling the control of reversibility for actuators based on cellulose nanofibers

Lisa Lopes da Costa, Céline Moreau, Denis Lourdin, Bernard Cathala, Ana

Villares

► To cite this version:

Lisa Lopes da Costa, Céline Moreau, Denis Lourdin, Bernard Cathala, Ana Villares. Unraveling the control of reversibility for actuators based on cellulose nanofibers. Carbohydrate Polymers, 2023, 314, pp.120951. 10.1016/j.carbpol.2023.120951. hal-04193857

HAL Id: hal-04193857 https://hal.inrae.fr/hal-04193857v1

Submitted on 1 Sep 2023

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

¹ Unraveling the control of reversibility for ² actuators based on cellulose nanofibers

3

4 Lisa Lopes da Costa,* Céline Moreau, Denis Lourdin, Bernard Cathala, Ana Villares

5 UR1268 BIA, INRAE, F-44316 Nantes, France

6 *lisa.lopes-da-costa@inrae.fr

7 Abstract

8 In this work, we have prepared cellulose-based actuators taking advantage of the pH-sensitive 9 solubility of chitosan (CH) and the mechanical strength of CNFs. Bilayer films were prepared by 10 vacuum filtration inspired by plant structures that exhibit reversible deformation under pH changes. 11 The presence of CH in one of the layers led to asymmetric swelling at low pH, thanks to the 12 electrostatic repulsion between charged amino groups of CH, and the subsequent twisting with the CH 13 layer on the outside. Reversibility was achieved by substituting pristine CNFs with carboxymethylated 14 CNFs (CMCNFs), that are charged at high pH and thus competed with the effects of amino groups. 15 Swelling and mechanical properties of layers under pH changes were studied by gravimetry and 16 dynamic mechanical analysis (DMA) to quantify the contribution of chitosan and the modified CNFs 17 on the reversibility control. This work evidenced the key role of surface charge and layer stiffness to 18 achieve reversibility. Bending was triggered by the different water uptake of each layer, and shape 19 recovery was achieved when the shrunk layer shower higher rigidity than the swollen layer.

20

21 Keywords

22 Cellulose films; chitosan; pH-responsive; programmable; carboxymethylation; dynamic mechanical

23 analysis

24 1. Introduction

25 Actuators are hierarchical structures that undergo shape changes triggered by external stimuli, such as 26 temperature, light, humidity, or pH. These materials respond to the current demand for engineering 27 multifunctional assemblies that adapt to their environment, as similarly observed in Nature (Erb et al., 2013; Ganewatta et al., 2021; Rüggeberg & Burgert, 2015; Y. Wang et al., 2020). By actuation, 28 29 biological systems such as the pinecones or wild wheat awns release their seeds in response to 30 humidity uptake and associated cell wall swelling, which guarantees plant survival (Dawson et al., 31 1997; Elbaum et al., 2007). Nature is therefore a source of inspiration for the fabrication of actuators 32 triggered by water swelling, and over the last few years, the use of natural polymers has shown a 33 growing interest in the scientific community. Among the myriad of natural polymers, cellulose 34 nanofibers (CNFs) revealed a great potential for this purpose (Chen et al., 2022; Liu et al., 2021). 35 CNFs have been used for decades in the papermaking industry, coatings, optical films, foodstuffs, 36 pharmaceutics, and cosmetics (Abdul Khalil et al., 2012; Y. Zhang et al., 2013). CNFs are obtained by 37 mechanical lamination of the cellulose fiber, and the fibrillation to the nanometer scale provides 38 attractive properties, including high aspect ratio, high strength and stiffness, low density, transparency, 39 high hydrophilicity, high surface area, and low thermal expansion (Dufresne, 2013; Eichhorn et al., 40 2010; Moon et al., 2011). Furthermore, cellulose is highly abundant and biodegradable, which makes 41 it an excellent candidate to fabricate novel materials (Huber et al., 2011). One of the main drawbacks 42 of cellulose in the fabrication of materials is the interaction with water due to the presence of a high 43 number of hydroxyl groups in the cellulose chain (Kuang et al., 2019; M. Wang et al., 2015; Wei et 44 al., 2021; Yang et al., 2021; L. Zhang et al., 2016). The water uptake is the main cause of the low 45 stability of cellulose materials, with a substantial loss of mechanical properties. Nevertheless, the 46 cellulose-water interaction can be exploited to design cellulose-based water-responsive actuators. By 47 carefully designing multilayered structures with different functionalities in each layer, water uptake 48 can be controlled to display asymmetrical expansion of the layers, and therefore to perform actuation. Indeed, the hydroxyl groups along the skeleton of cellulose enable to extend CNF versatility by 49 50 chemical modification resulting in "tailor-made" specific properties (Habibi, 2014).

51 In previous studies, we prepared pH-responsive bilayer films based on functionalized CNFs (Chemin 52 et al., 2020). The first layer contained carboxylated CNFs and the second layer was composed of 53 aminated CNFs. When the film was subjected to high pH, the deprotonation of carboxylic acid groups 54 caused electrostatic repulsion between the negative charges, which facilitated water adsorption. 55 Therefore, the different water uptake and expansion of the negatively charged carboxylated layer 56 compared to the aminated layer resulted in film bending. Then, the inverse mechanism was obtained at 57 low pH because of the protonation of the aminated layer, which adsorbed more water by a similar 58 mechanism. Despite the controlled bending at high and low pH, the actuation of these bilayer films was not reversible, and the film remained bent when it returned to the initial conditions. Therefore, in 59 60 this work, we focused on the interaction between water and the functional groups grafted on the 61 cellulose surface to understand the mechanism of actuation, and the parameters required to achieve 62 reversibility.

We succeed in fabricating a new bilayer actuator displaying reversible shape-changing under pH variation. First, CNFs were combined with chitosan (CH) to form a pH-sensitive film (CH-CNF) with promising mechanical properties by physical interlocking between CH and CNF networks without chemical cross-linkers. CH protonates in acid environments due to the formation of non-bonding pairs of electrons in amino groups, allowing its solubilization. Above pK_a of 6.5, approximatively, CH chains at their neutral form associate together and consequently aggregate (Rosca et al., 2005;

69 Vleugels et al., 2018). Hence, CH can be considered for designing responsive structures thanks to its

- 70 pH-dependent solubility properties. We studied the behavior of CH-CNF films in terms of adsorption
- of CH on CNFs, polymer amount and distribution, and pH-associated structural changes. In the second
- stage, we fabricated bilayer actuator films by combining CH-CNF layers with CNF-based layers (non-
- 73 modified or carboxymethylated) and we monitored the actuator properties in terms of swelling and
- 74 mechanical properties. This work presents polysaccharide actuators fabricated by their tailored 75 combination and provides a better understanding of the polysaccharide-water interactions that gives
- 76 more insight into the requirements to achieve reversible actuation.
- 77

78 2. Materials and Methods

79 2.1.Materials

80 CNFs Exilva P 01-V (10% w/v) were kindly provided by Borregaard (Sarpsborg, Norway). Chitosan (medium molecular weight, 75-85 % deacetylated) (CH), monochloroacetic acid (MCA), hydrochloric 81 82 acid (HCl), rhodamine B isothiocyanate (RBITC), and sodium hydroxide (NaOH) were purchased from Sigma Aldrich. Dimethyl sulfoxide (DMSO) was purchased from Merck, isopropanol from 83 84 VWR, and poly(allylamine hydrochloride) (PAH) from Polysciences. A Spectra/Por dialysis 85 membrane, MWCO 12-14,000 Da, was purchased from Spectrum Laboratories Inc, and 86 polyvinylidene fluoride (PVDF) membranes (pore size 0.22 µm, diameter 47 mm) from Merck. Water was purified by the Millipore Milli-Q purification system (18.2 M Ω). 87

88 2.2. Fluorescence labeling of chitosan (CH)

A CH solution was prepared at 1% (w/v) in 0.1 M HCl. Then, RBITC, dissolved in 20 mL of DMSO, was added to the CH solution. The reaction was stirred for 3 h, in dark, at room temperature. RBITClabelled chitosan (RBITC-CH) was precipitated by addition of NaOH. RBITC-CH was purified by centrifugation until no free RBITC was detected in supernatant. RBITC-CH was solubilized at 1 % (w/v) in 0.1 M HCl and dialyzed against ultra-pure water for 3 days in dark.

94 2.3.Carboxymethylation

95 Carboxymethylated CNFs (CMCNFs) were synthesized by the reaction between hydroxyl groups and 96 MCA, as previously described (Lopes da Costa et al., 2023). CNFs (500 mg) were impregnated with 97 MCA dissolved in isopropanol. Two MCA/anhydroglucose (AGU) molar ratios were performed (0.50 98 and 0.70) to achieve different degrees of substitution (DS). In parallel, NaOH was dissolved in 99 isopropanol at 60°C for 1 h. Impregnated CNFs were added to the NaOH solution and the mixture was 910 stirred at 70°C for 1 h. The resultant CNFs were purified by filtration through PVDF membranes with 91 ultra-pure water, then 0.1 M HCl, and finally ultra-pure water.

102 2.4. Film fabrication

Cellulose films were fabricated by Büchner filtration through PVDF membranes. Before filtration, all 103 104 polymer dispersions (CNFs, CH-CNF mixture, and CMCNFs) were prepared at 2.5 g·L⁻¹. For the single-layer films, the dispersion was poured into the membrane and filtrated for 30 min under 50 105 106 mbar at room temperature. For the bilayer films, after filtrating the first dispersion, the second 107 dispersion was added to the first layer and filtrated again under the same conditions. All films were, 108 then, dried between Whatman filter paper under a weight of 500 g for 48 h at room temperature. 109 Afterward, the disc-shaped films (diameter = 36 mm) were weighted and their thickness was measured with a Mitutoyo digimatic micrometer. The density (d) was calculated as represented below: 110

$$d = \frac{\mathrm{m}}{\mathrm{\pi}r^2 t} (1)$$

- 112 where *m*, *r*, and *t* are respectively the weight, radius, and thickness of the dried films.
- 113 The films were then cut into different shapes depending on the conducted analysis; rectangles (6 x 24 mm^2) for actuation tests (2.5.6) and dynamic mechanical analysis (2.5.8) and squares (24 x 24 mm^2)
- 115 for swelling ratio quantification (2.5.7).
- 116 2.5.Characterization
- 117
- 118 2.5.1. Quartz crystal microbalance with dissipation monitoring (QCM-D)

QCM-D equipment (E4, Q-Sense instrument, AB, Sweden) using a piezoelectric AT-cut quartz crystal 119 120 coated with gold electrodes on each side (QSX301, Q-Sense) was employed to assess the adsorption of CH on CNFs. The quartz crystals, cleaned in a piranha bath for 5 min, were set up into the cells and 121 ultra-pure water was injected at a flow rate of 0.1 mL·min⁻¹ at 20 °C for at least 1 h until reaching a 122 stable baseline. A PAH solution (0.1 g·L⁻¹) was first injected for 15 min before the adsorption of 123 biopolymers. Then, CNF dispersion $(0.1 \text{ g} \cdot \text{L}^{-1})$ and CH solution $(0.2 \text{ g} \cdot \text{L}^{-1}, \text{ pH 1})$ were successively 124 injected into the cells for 15 and 20 min respectively. A rinsing step with Milli-Q water or 0.1 M HCl 125 126 was done before the subsequent injection of polymer solutions.

127 Frequency $(\Delta f_n/n)$ and dissipation $(\Delta D_n/n)$ changes were simultaneously registered at 5 MHz 128 fundamental resonance frequency and its several overtones as a function of time. $(\Delta f_n/n)$ can be related 129 to the hydrated mass of the adsorbed polymers while $(\Delta D_n/n)$ provides information about 130 conformational changes and softness/rigidity of the system studied. QCM-D experiments were carried 131 out at least twice. The third overtone (15 MHz) was used in the evaluation of the QCM-D data.

- 2.5.2. Attenuated total reflection Fourier transform infrared (ATR-FTIR) spectroscopy 132 ATR-FTIR spectra were recorded with a Nicolet iS50 FTIR spectrometer (Thermo Fisher, Waltham, 133 Massachusetts, USA) with the accumulation of 200 scans between 800 and 4000 cm⁻¹ with a 134 135 resolution of 4 cm⁻¹. The single-layer films composed of CNFs, CH-CNF, or CH were analyzed. A band decomposition of CH film spectra was performed in the 1800-1500 cm⁻¹ region to determine the 136 exact positions of the peak related to C=O stretching from amide I and the two peaks associated with 137 138 N-H deformation. OriginPro 8.5 software was used for the band decomposition. The spectrum was first normalized with the peak at 1150-1085 cm⁻¹ (ether bonding) with a linear baseline made by 139 interpolation. The decomposition was done in Gaussian peaks with the minimum number of peaks and 140 141 was considered valuable when $R \ge 0.99$. For CH-CNF spectra, the characteristics of the peak at 1536 cm⁻¹ (N-H deformation) were measured to follow the CH amount in CH-CNF films (wavenumber, 142 area, full width at half maximum, and area contribution). The spectra were also normalized with the 143 peak at 1150-1085 cm⁻¹ with a linear baseline made by interpolation. 144
- 145 2.5.3. Elementary analysis

To quantify the percentage of CH in the CH-CNF films, the atomic composition (carbon, hydrogen,
and nitrogen content) of freeze-dried CNFs, single-layer CH and CH-CNF films was determined using
a Thermo Fischer Scientific Elemental Analyzer (Model Flash 2000). Each sample was measured
twice.

150 2.5.4. Confocal laser scanning microscopy (CLSM)

Single-layer films containing RBITC-CH and CNF (weight ratio 1:4) were observed by confocal microscopy (Leica TCS-SP2) to visualize the distribution of CH within the film. Confocal images were acquired with a LSM-510 UV-Vis microscope equipped with 63×1.2 water immersion objectives (Zeiss, Germany). Excitation/emission wavelengths were 530/566 nm.

155 2.5.5. Conductometric titration

156 CMCNFs (20 mg) were firstly acidified with HCl (0.1 M) and titrated with NaOH (0.1 M) by a 157 TIM900 titration manager (Metrohm). The conductivity was measured simultaneously during the titration by a CDM749 conductivity cell. 158

159 The degree of substitution (DS) was calculated as (Abitbol et al., 2013; Beck et al., 2015):

160
$$DS = \frac{162(V_{eq2} - V_{eq1})C_{NaOH}}{m - M(V_{eq2} - V_{eq1})C_{NaOH}}$$
(2)

161 Where V_{eq1} and V_{eq2} are the equivalence volumes of NaOH (L), C_{NaOH} is the concentration of NaOH $(mol \cdot L^{-1})$, *m* is the dried weight of CNFs (g), *M* is the molecular weight of the carboxymethyl group, 162

163 and 162 is the molecular weight of AGU ($g \cdot mol^{-1}$).

164 2.5.6. Actuation tests

165 Actuation was monitored by immersing the bilayer films under different pH conditions (acidic, neutral, and basic). Bilayer films were cut into rectangles $(24 \times 6 \text{ mm}^2)$ and were fixed in a clamp that 166 167 was held by a stand. A glass vessel containing 0.01 M HCl (pH 2), Milli-Q water, or 0.01 M NaOH 168 (pH 12) was placed on the moveable platform. Then, a SVS-vistek EXO camera and the camera software from BIA unit were used to capture the film movement. Therefore, the moveable platform 169 170 was driven up to dip the bilayer film into the liquid in the glass vessel and start the actuation test. The test steps were the following: the bilayer films were first immersed for 10 min in 0.01 M NaOH, then 171 5 min in 0.01 HCl, and 20 min in 0.01 M NaOH. Each actuation test was repeated three times. 172

173 2.5.7. Swelling ratio quantification

Disc-shaped films were cut into squares of $24 \times 24 \text{ mm}^2$. The dry films were weighed, then immersed 174 in Milli-Q water, or aqueous solutions of 0.01 M HCl (pH 2) or 0.01 M NaOH (pH 12) for 1 min and 175 weighed again. The swelling ratio expressed as the weight of absorbed water per the weight of the dry 176 177 film (SW) was calculated following this equation:

 $(m_w - m_d)$ CIAZ 178

$$SW = \frac{(m_w - m_d)}{m_d} \tag{3}$$

- 179 where m_w and m_d are respectively the weight of the wet and dry film (mg).
- 180 Each sample was measured five times.

181 2.5.8. Dynamic mechanical analysis (DMA)

The dynamic mechanical properties as a function of pH for CNF, CH-CNF, or CMCNF films were 182 determined with a Dynamic Mechanical Analyser (DTMA MkIV, Rheometric Scientific, US). The 183 184 DMA system was flipped from bottom to top in order to measure the samples immersed in pH-185 controlled aqueous solutions at 20°C. Tests were conducted on rectangular-shaped strip samples (10×6 mm^2) in the tensile mode with a dynamic strain amplitude of 0.1%, and a constant load force of 0.1 N. 186 187 Recordings of the specimens were taken following this procedure: 10 min air equilibrium, then 10 min immersion in the liquid (HCl 0.01 M, Milli-Q water, or NaOH 0.01M). 188

189

3. Results and discussion 190

- 3.1. Structure and pH-sensitive behavior of chitosan-cellulose films (CH-CNF) 191
- 192

193 In this work, we took advantage of the pH-responsive properties of chitosan (CH) to fabricate all-194 polysaccharide actuators. CH contains amino groups that make the polymer pH-responsive. In acidic

- conditions, the charged amino groups favor the hydration and dissolution of CH chains due to the 195
- higher hydrophilicity and electrostatic repulsion between the positive charges. Above its pK_a (~ 6), 196
- 197 the neutralization of the amino groups drives CH aggregation by the association between the CH
- chains (Rosca et al., 2005; Vleugels et al., 2018). However, CH-based actuators require cross-linking 198
- 199 of the polysaccharide chains, to avoid the dissolution of CH at low pH. Therefore, in this work, CH
- 200 was combined with cellulose nanofibers (CNFs) to form a physically cross-linked network within the
- 201 film. The first step was therefore to demonstrate the irreversible association between CH and CNF.
- 202 For this purpose, the interaction between CH and uncharged CNFs were investigated by quartz crystal 203 microbalance with dissipation monitoring (QCM-D).
- 204 Fig. 1 shows the normalized frequency $(\Delta f_n/n)$ and dissipation $(\Delta D_n/n)$ signals as a function of time for 205 the successive adsorption of CNFs and CH. ($\Delta f_n/n$) shifts can be related to the change in mass of 206 adsorbed polymers (hydrated mass) and $(\Delta D_n/n)$ shifts give information on the viscoelastic properties 207 of the adsorbed polymers. The studied concentrations and injection times of the polymer solutions 208 were chosen according to our previous studies (Lombardo et al., 2021). Each polymer adsorption stage 209 was followed by a rinsing step to remove excess and loosely bound polymers. Polymer adsorption was 210 considered complete when a plateau of frequency and dissipation was reached. A layer of positively 211 charged PAH was first deposited to facilitate the adsorption of CNFs on the gold substrate. The first 212 CNF injection led to a decrease in frequency of -41 ± 4 Hz, evidencing the polymer adsorption. An 213 average frequency shift between -20 and -50 Hz has been previously reported for CNF adsorption 214 (Ahola et al., 2008; Aulin et al., 2010; de Castro et al., 2018), depending on adsorption conditions and 215 the CNF nature. In order to demonstrate the surface saturation after each adsorption step, we injected 216 twice CNF dispersions after the rinsing step to guarantee that no more CNFs were adsorbed. The 217 signals of frequency and dissipation in Fig. S1 remained unchanged after the second CNF injection, meaning that the maximum CNF adsorption was reached during the first stage. Then, when CH was 218 219 adsorbed on the CNF layer, the frequency decreased to -54 ± 4 Hz, and the dissipation increased to about $(17 \pm 2) \cdot 10^{-6}$ units (Fig. 1). During the rinsing step, the change from 0.1 M HCl to water 220 221 resulted in an increase in frequency signal and dissipation during the five first minutes. Then, the frequency remained stable at -43 ± 9 Hz while the dissipation signal decreased until attaining stability. 222 223 This type of signal can be associated with the phenomenon of dewatering occurring when CH is 224 subjected to pH changes and the binding is entropy-driven (Leray et al., 2022; Lombardo et al., 2021; 225 Lombardo & Thielemans, 2019). The subsequent injection of CNF on the CH layer resulted in a 226 frequency decrease at -70 ± 10 Hz, which demonstrated the irreversible adsorption of CH on the first
- 227 CNF layer, which allowed the adsorption of the second CNF layer.





Fig. 1. $\Delta f_n/n$ and $\Delta D_n/n$ changes for the third overtone during the adsorption of CNFs and CH on the gold surface of the QCM-D sensor as a function of time. Arrows indicate the injection of the polymers and the asterisks or points indicate the rinse steps in between with water or 0.1 M HCl, respectively.

232 Once demonstrated the association between CNFs and CH, in order to design CH-CNF films with the 233 optimal loading of CH, we prepared mixtures of CH (1 g·L⁻¹) and CNFs (2.5 g·L⁻¹) at different weight 234 ratios: 5, 10, 20, 30, and 40% wt of CH (CH05%, CH10%, CH20%, CH30%, and CH40%, 235 respectively). We prepared CH-CNF films by vacuum filtration following the protocol in section 2.4. When the CH concentration was above 50% wt, the films were stickered to the membrane hindering 236 237 their potential use. Films exhibited average thicknesses of $24 \pm 4 \mu m$ with an apparent density of 0.77 ± 0.01 g·cm⁻³. A film of only pristine CH (CH100%) was prepared by casting for comparison with the 238 CH-CNF films. Firstly, films were examined by ATR-FTIR to demonstrate that CH was not eluted 239 240 from the CNF matrix during the filtration.

ATR-FTIR spectra of films are presented in Fig. 2a-b. In Fig. 2a, the CNF film spectrum displayed 241 bands at 3500-3300 cm⁻¹ (-O-H stretching vibration), 2874-2922 cm⁻¹ (C-H stretching), 1634 cm⁻¹ 242 (adsorbed water), 1380 cm⁻¹ (CH₃ deformation), and 1150-1085 cm⁻¹ (ether bonding) (Oh et al., 2005; 243 244 Schwanninger et al., 2004). CH100% film spectrum differed from CNF due to remarkable bands at 1630 cm⁻¹ (C=O, amide I), 1555 and 1516 cm⁻¹ (N-H deformation) (Da Róz et al., 2010; Rosca et al., 245 2005) (Fig. 2a). The exact position of these three peaks was determined by deconvolution of the broad 246 band between 1800 and 1450 cm⁻¹ (Fig. S2). For CH05% and CH10%, the bands corresponding to the 247 amide groups from CH were slightly detected, probably due to a low CH concentration (Fig. 2b). 248 Starting from the CH20% spectrum, a band at 1536 cm⁻¹ indicated the presence of CH in the film, 249 250 corresponding to the merging of the two N-H bands at 1555 and 1516 cm⁻¹ (Fig. 2a-b). A peak fitting 251 between 1800 and 1450 cm⁻¹ was done to quantify the evolution of the band area at 1536 cm⁻¹. Fig. S2 and Table S1 report the peak fitting of the CH20%, CH30%, and CH40%; and the respective results. 252 253 An area of 0.62 was obtained for CH20%; then was amplified for CH30%, with a value of 0.96; and 254 finally, for CH40%, the band decreased to a similar area to CH20% (band area of 0.46) (Fig 2b). In all cases, the band associated with the C=O stretching from the amide group at 1630 cm⁻¹ was overlapped 255

with the band of adsorbed water at 1634 cm⁻¹ (Zhao et al., 2020). ATR-FTIR results suggested that, at a CH loading above 30%, excess CH was not retained by the CNFs during filtration and was eluted through the membrane. This result was consistent with the limitations encountered during film preparation at CH content above 50% *wt* by vacuum filtration.





Fig. 2. (a) ATR-FTIR spectra of CNF, CH20%, CH100% films and molecular structures of CH and
CNF. (b) ATR-FTIR spectra of CNF, CH05%, CH10%, CH20%, CH30%, CH40% and CH100% films
in the 1800 – 1450 cm⁻¹ region. The whole spectra are presented in Fig. S3.

The CH20% film was selected for the fabrication of actuators to prevent CH loss during film preparation. To get more insight into the composition of the CH20% film, the atomic percentage of carbon (%C), hydrogen (%H), and nitrogen (%N) of the CNF, CH100%, and CH20% films were measured by elementary analysis. The percentage of CH (%*CH*) in the film could be calculated as the ratio between the %N of CH20% and CH100% films:

269
$$\% CH = \frac{\% N_{CH20\% film}}{\% N_{CH100\% film}} 100$$
(4)

Table 1 shows the results of the elementary analysis. The CNF film was composed of 40.9 %C and 6.2 %H, and witnessed any presence of nitrogen in agreement with the composition of cellulose (Kulpinski et al., 2022; Missoum et al., 2012). The CH film also fitted the common values with 5.3 %N, 30.8 %C, and 5.8 %H (Gupta & Jabrail, 2006; Qian et al., 2018; Sarwar et al., 2015). In the CH20% film, 0.83 %N was detected asserting the successful CH incorporation within the film during film preparation. Therefore, a %CH of 15.6 ± 0.1 % was obtained which correlated well with the theoretical value of 20% CH.

	CNF	CH100%	CH20%
%C	40.9 ± 0.0	30.8 ± 0.2	39.8 ± 0.2
%H	6.2 ± 0.0	5.8 ± 0.7	6.3 ± 0.0
%N	0.0 ± 0.0	5.3 ± 0.0	0.8 ± 0.1

277 Table 1. Elemental analysis results of CNF, CH100%, and CH20% films.

278

279 The protonation of the amino groups of CH20% films in response to pH was monitored by ATR-280 FTIR. Fig. 3a shows the ATR-FTIR spectra of CH20% films at pH 2 and 12. The whole spectra are shown in Fig. S4. The deprotonation of the amino groups from CH was demonstrated by the shift from 281 1543 cm⁻¹ to 1574 cm⁻¹ of the band corresponding to the N-H deformation. Then, the CH20% film at 282 283 different pHs was observed by confocal laser scanning microscopy (CLSM) to evaluate the 284 distribution of CH within the film depending on pH. CH was labeled with rhodamine B isothiocyanate 285 (RBITC-CH) and was mixed with the CNF dispersion to prepare RBITC-CH mixed CNF at 20% wt (RBITC-CH20%). Then, films were prepared following the same procedure as described above. Fig. 286 287 3b-e shows CLSM images of the film after vacuum filtration and when the films are immersed in 288 different pH solutions. For the starting dry film, a homogenous distribution of CH was observed (Fig. 3b). When the film was immersed at pH 7 (Fig. 3c), the distribution profile changed compared to the 289 290 starting film, and a network of CH aggregates appeared. Then, at pH 2 (Fig. 3d), the distribution 291 pattern observed at pH 7 was amplified, and the film showed regions with a high CH concentration 292 (highly colored) and regions with low CH content. Surprisingly, when the film was at pH 12 (Fig. 3e), 293 the CH distribution seemed more homogeneous with a similar pattern to the dry film. We 294 hypothesized that under acid conditions, CH is highly swollen, as the QCM-D results suggested, so it 295 can diffuse within the CNF network creating a second network that expanded due to electrostatic repulsion between amino groups. Differently, under alkaline conditions, CH shrinks, and the layer 296 297 dewaters forming aggregates that provide a more compacted structure. This was confirmed by the 298 dissipation decrease when the CH layer was rinsed with water (Fig. 1). Therefore, CH20% film 299 exhibited strong structural changes depending on the pH.



300

Fig. 3. (a) ATR-IR spectra of CH20% film after immersion at pH 2 and 12 in the 1800 – 1450 cm⁻¹ region. The whole spectra are presented in Fig. S4. Confocal images of RBITC-CH20% film after vacuum filtration (b), and wet state for different pH values (c) 7, (d) 2, and (e) 12.

304

305 3.2. Reversible actuation of bilayer films controlled by CH and charge CNFs

We fabricated bilayer films combining a first layer of CH20% and a second layer of CNFs or 306 carboxymethylated CNFs (CMCNFs). Two molar ratios between MCA and AGU (MCA/AGU: 0.50, 307 and 0.70) were investigated to achieve different degrees of substitution (DS) based on a calibration 308 curve performed in previous studies (Lopes da Costa et al., 2023). The DS were determined by 309 310 conductometric titration following equation (2) (a representative titration curve is presented in Fig. 311 S5). CMCNFs with DS of 0.10, and 0.20 (CMCNF10, and CMCNF20) were obtained for the 312 respective MCA/AGU ratios of 0.50, and 0.70. CMCNF20 formed very stable dispersions so it was 313 mixed at 80% wt with CNFs to avoid film dispersion in aqueous media. The CH20%/CNF film 314 exhibited an average thickness of $34 \pm 2 \,\mu\text{m}$ with an apparent density of $0.71 \pm 0.05 \,\text{g}\cdot\text{cm}^{-3}$, whereas the CH20%/CMCNF films showed an average thickness of $28 \pm 5 \,\mu m$ with an apparent density of 0.75 315 $\pm 0.07 \text{ g} \cdot \text{cm}^{-3}$. 316

317 Fig. 4a-c shows the actuation performance of the CH20%/CNF and CH20%/CMCNF films under pH 318 changes. The films were first immersed for 10 min at pH 12 (NaOH 0.01M), then 5 min at pH 2 (HCl 0.01M), and back to pH 12 for 20 min. In Fig. 4a, the CH20%/CNF film bent with the CH20% layer 319 on the outside at pH 12. The presence of CH in the CH20% layer already entailed a difference in water 320 321 uptake between the CH20% and CNF layers at high pH, and subsequently a differential expansion 322 within the film. Then, the immersion of the CH20%/CNF film in the pH 2 solution resulted in 323 twisting, because of higher expansion of the CH20% layer. The protonation of the amino groups from the CH20% layer created repulsions between the positively charged groups, further enhancing the 324 difference in water uptake and expansion between CH20% and CNF layers. Back to pH 12, the 325 CH20%/CNF film did not show any change from low pH, and the film remained twisted. Now in their 326 327 basic form, the amino groups should induce a layer shrinkage, however, the shape recovery did not 328 operate.

Differently, when CH20% was combined with CMCNFs, the presence of carboxymethyl groups 329 influenced the behavior (Fig. 4b and c). The expansion of CH20% and CMCNF layers at the different 330 331 pH was illustrated in Fig. S6. CH20%/CMCNF10 and CH20%/CMCNF20 bent with the CMCNF 332 layer on the outside at pH 12. The carboxylic acid groups were deprotonated, creating the electrostatic repulsion between negative charges within the CMCNF layer, which yielded the layer expansion and 333 the film bending. Moreover, CH20%/CMCNF20 was more bent than CH20%/CMCNF10, as the DS 334 335 increased. The inverse mechanism was noted at pH 2 in which the CH20% layer is on the bending outside; in this case, due to the protonation of the amino groups. Back to pH 12, CH20%/CMCNF10 336 reached an almost straight shape while CH20%/CMCNF20 achieved complete backward movement to 337 the other side. This meant that the difference in water uptake between CMCNF10 and CH20% layers 338 was not sufficient to fully reverse bending compared to CH20%/CMCNF20. Therefore, the DS was 339

340 revealed to be a critical parameter influencing the reversible actuation.



341

Fig. 4. Photographs showing the actuation tests under pH changes of (a)CH20%/CNF, (b)
 CH20%/CMCNF10, and (c) CH20%/CMCNF20 films.

344 To give more insight into the mechanism of actuation, the swelling of each layer was studied by 345 fabricating single-layer films of CH20%, CNF, or CMCNFs. Swelling ratios (SW) of single-layer 346 films immersed at pH 2, 7, and 12 are reported in Fig. 5a. Comparing CH20% and CNF films, the difference in SW turned out to be weak. In neutral and basic conditions, CH20% and CNF films had 347 348 an average SW of 3.4 ± 0.4 and 2.4 ± 0.2 , respectively. A slightly greater difference was obtained at 349 pH 2 (SW of 4.0 ± 0.5 for CH20% and 2.0 ± 0.4 for CNF films). As the CH20% film had in any case a 350 superior SW than the CNF film, which justified that bending only occurred with the CH20% layer on 351 the outside. Moreover, this fact explained the difficulty in achieving shape recovery at pH 12 after film 352 twisting at pH 2. Regarding the CMCNF films, the water uptake depended on the number of carboxymethyl groups on the cellulose surface. CMCNF10 film showed higher water uptakes, 353 354 compared to the CNF film, but still a low dependency on pH (average SW of 3.4 ± 0.2). Differently, CMCNF20 film displayed an increasing water uptake from pH 2 to 12 ($4.5 \pm 0.5, 6.7 \pm 0.4$, and $8.0 \pm$ 355 0.8 for pH 2, 7, and 12, respectively). Hence, the introduction of carboxymethyl groups ended up 356 357 competing with the behavior of the CH20% layer. The increase in DS allowed bigger changes in SW

and thus, enabled reversible movements of the film. Therefore, when the films were again in an alkaline environment, the shape recovery can be completed only if a consequent swelling of the CMCNF layer is reached. This was the case of CH20%/CMCNF20, where both layers showed a significant difference in SW at high pH.



Fig. 5. (a) SW of single-layer CH20%, CNF, CMCNF10, and CMCNF20 films. (b) E' and (c) ε of
single-layer CH20%, CNF, and CMCNF20 films as a function of pH.

Dynamic mechanical analysis (DMA) measurements were conducted on the single-layer CH20%, 365 366 CNF, and CMCNF20 films to provide more detailed information on the changes in mechanical properties of layers at the different pHs. The storage modulus (E') of the films was recorded as a 367 368 function of pH (Fig. 5b). The CNF film presented relatively the same mechanical properties for all the 369 studied pHs, with an average E' of 276 ± 4 MPa in agreement with previous studies (Lopes da Costa et al., 2023). The CH20% film displayed lower E' when immersed in aqueous solutions and an increase 370 at high pH (E' of 58 ± 2 MPa, 65 ± 6 MPa, and 155 ± 36 MPa for pH 2, 7, and 12, respectively). It has 371 372 been reported that the incorporation of CH enhanced water uptake within the film destroying 373 interfibrillar hydrogen bonding (Wu et al., 2020), which justified the lower E' values of CH20% 374 compared to CNF. The differences in E' determined the reversibility of bending upon pH changes. 375 Thus, the lower E' of the CH20% layer facilitated its expansion compared to the CNF layer, and thus 376 drove the bending of the CH20%/CNF film. However, the higher stiffness of the CNF layer prevented 377 the shape recovery. Regarding the CMCNF20 film, E' was significantly lower compared to the CNF film (average E' of 26 ± 6 MPa). The introduction of carboxymethyl groups drastically changed the 378 379 cellulose-water interaction, which resulted in a lower stiffness. Moreover, this indicates that both 380 layers of the CH20%/CMCNF20 film displayed low E'. At pH 12, the CMCNF20 layer was less rigid 381 than the CH20% layer, which facilitated the reversibility. Thus, flexibility is a key factor influencing 382 the reversibility properties of the films.

383 DMA measurements also allowed the determination of changes in the film length (ε) upon immersion 384 in the liquid (Fig. 5c). CNF layer presented low ε regardless of pH (average ε of 5.5 ± 0.3 %). The CH20% layer showed a strong decreasing ε from pH 2 to 12 (ε of 9.1 ± 0.0 %, 7.7 ± 0.4 %, and 5.5 ± 385 1.0 % for pH 2, 7, and 12, respectively). Inversely, CMCNF20 film exhibited increasing ε from pH 2 386 387 to 12 (5.5 \pm 0.0 %, 9.7 \pm 0.0 %, and 10.6 \pm 0.2 % for pH 2, 7 and 12, respectively). These changes in 388 dimension reflected the actuation behavior of the different bilayer films. Especially for the change 389 from pH 2 to 12, a strong difference in ε at pH 2 for both couples is noted (CH20% & CNF and 390 CH20% & CMCNF20) while, at pH 12, only CH20% and CMCNF20 layers exhibited a significant 391 difference in ε .

392 This work demonstrates that polysaccharides can be combined into bilayer films to achieve pH-393 responsive movement. The presence of charged groups, negative carboxymethyl or positive amino groups, increases the water uptake, mainly due to the electrostatic repulsion between charges, which 394 facilitates the separation and the interaction with water. The higher swelling also decreases the storage 395 396 modulus of the charged layers compared to unmodified CNFs. Actuation can be achieved by 397 assembling layers showing different expansions; however, reversibility requires specific mechanical properties of the layers. Therefore, actuation is not only driven by the different water uptake of each 398 399 layer, higher flexibility of the expanded layer is required to achieve reversibility. The role of the storage modulus of each layer forming the film on the actuation performance has been previously 400 401 described for fiber composites, and wood bilayer strips (Le Duigou et al., 2019; Rüggeberg & Burgert, 402 2015). Our results evidenced that, when changing the environment, the layer shrinking must provide 403 enough rigidity to drive backward movement. Therefore, by carefully selecting key polysaccharides or 404 combinations of them, we can tune the film bending and/or twisting, and the reversibility of the 405 actuation.

406 4. Conclusion

407 In this work, we demonstrated that CH, unmodified, and carboxymethylated CNFs together can give 408 rise to pH-responsive bilayer actuator films with controlled reversibility. The presence of functional 409 groups sensitive to pH and the assembly of multilayered structures permitted to generate variations of expansion within the film responsible for shape-changing. In this case, the control of reversibility was 410 achieved by the chemical pathway to obtain carboxylic acid groups at the CNF surface and their 411 412 degrees of substitution. Functionalization opens the route to control the reversibility of pH-responsive 413 actuators. Our results demonstrated that actuation was not only driven by water uptake but also by the 414 mechanical properties of the layers. We provided fundamental information in terms of CH-CNF 415 interactions, structural arrangement, swelling behavior, and mechanical properties of the different 416 layers contributing to the mechanism of actuation. Therefore, this study enables to enhance the 417 understanding of polysaccharides for the development of programmable materials in various 418 applications such as soft robotics, biomedical, food, and electronic applications.

419 Acknowledgments

The authors gratefully thank the financial support of Région Pays de la Loire and Transform Division from the French National Research Institute for Agriculture, Food and Environment. The authors acknowledge the Bioresources, Imaging, Biochemistry, and Structure (BIBS) platform of INRAE for the access to microscopy facilities (Bruno Novales). Anne-Laure Reguerre is acknowledged for the excellent technical support for image acquisition and treatment. Denis Loquet (Nantes University) is acknowledged for his technical support in elementary analysis. We thank Borregaard for kindly providing the cellulose nanofibers.

427 Availability of data

428 The datasets generated during the current study are available from the corresponding author on 429 reasonable request.

430 Conflict of interest

431 The authors declare that they have no known competing financial interests or personal relationships 432 that could have appeared to influence the work reported in this paper.

433 References

- Abdul Khalil, H. P. S., Bhat, A. H., & Ireana Yusra, A. F. (2012). Green composites from sustainable
 cellulose nanofibrils: A review. *Carbohydrate Polymers*, 87(2), 963–979.
 https://doi.org/10.1016/j.carbpol.2011.08.078
- Abitbol, T., Kloser, E., & Gray, D. G. (2013). Estimation of the surface sulfur content of cellulose
 nanocrystals prepared by sulfuric acid hydrolysis. *Cellulose*, 20(2), 785–794.
 https://doi.org/10.1007/s10570-013-9871-0
- Ahola, S., Salmi, J., Johansson, L. S., Laine, J., & Österberg, M. (2008). Model films from native cellulose nanofibrils. Preparation, swelling, and surface interactions. *Biomacromolecules*, 9(4), 1273–1282. https://doi.org/10.1021/bm701317k
- Aulin, C., Johansson, E., Wågberg, L., & Lindström, T. (2010). Self-organized films from cellulose i
 nanofibrils using the layer-by-layer technique. *Biomacromolecules*, 11(4), 872–882.
 https://doi.org/10.1021/bm100075e
- Beck, S., Méthot, M., & Bouchard, J. (2015). General procedure for determining cellulose nanocrystal
 sulfate half-ester content by conductometric titration. *Cellulose*, 22(1), 101–116.
 https://doi.org/10.1007/s10570-014-0513-y
- Cai, Z., & Kim, J. (2009). Cellulose–chitosan interpenetrating polymer network for electro-active
 paper actuator. *Journal of Applied Polymer Science*, *114*(1), 288–297.
 https://doi.org/10.1002/app.30456
- Chemin, M., Beaumal, B., Cathala, B., & Villares, A. (2020). Ph-responsive properties of asymmetric
 nanopapers of nanofibrillated cellulose. *Nanomaterials*, 10(7), 1–14.
 https://doi.org/10.3390/nano10071380
- Chen, W., Sun, B., Biehl, P., & Zhang, K. (2022). Cellulose-Based Soft Actuators. *Macromolecular Materials and Engineering*, 2200072. https://doi.org/10.1002/mame.202200072
- Christina, J. Y., Håkan, D., Björn, E., Norgren, L. M., Yang, J., Dahlström, Á. C., Edlund, Á. H.,
 Lindman, Á. B., & Norgren, Á. M. (2019). pH-responsive cellulose-chitosan nanocomposite
 films with slow release of chitosan. *Cellulose*, 26, 3763–3776. https://doi.org/10.1007/s10570019-02357-5
- 461 Da Róz, A. L., Leite, F. L., Pereiro, L. V., Nascente, P. A. P., Zucolotto, V., Oliveira, O. N., &
 462 Carvalho, A. J. F. (2010). Adsorption of chitosan on spin-coated cellulose films. *Carbohydrate*463 *Polymers*, 80(1), 65–70. https://doi.org/10.1016/j.carbpol.2009.10.062
- 464 Dawson, C., Vincent, J., & Rocca, A. (1997). How pine cone open. *Nature*, *390*(18–25), 668.
- de Castro, D. O., Tabary, N., Martel, B., Gandini, A., Belgacem, N., & Bras, J. (2018). Controlled
 release of carvacrol and curcumin: bio-based food packaging by synergism action of TEMPOoxidized cellulose nanocrystals and cyclodextrin. *Cellulose*, 25(2), 1249–1263.
 https://doi.org/10.1007/s10570-017-1646-6
- 469 Duan, J., Liang, X., Zhu, K., Guo, J., & Zhang, L. (2017). Bilayer hydrogel actuators with tight
 470 interfacial adhesion fully constructed from natural polysaccharides. *Soft Matter*, *13*(2), 345–354.
 471 https://doi.org/10.1039/c6sm02089e
- 472 Dufresne, A. (2013). Nanocellulose: a new ageless bionanomaterial. *Materials Today*, *16*(6), 220–227.
 473 https://doi.org/10.1016/j.mattod.2013.06.004
- Eichhorn, S. J., Dufresne, A., Aranguren, M., Marcovich, N. E., Capadona, J. R., Rowan, S. J., Weder,
 C., Thielemans, W., Roman, M., Renneckar, S., Gindl, W., Veigel, S., Keckes, J., Yano, H., Abe,
 K., Nogi, M., Nakagaito, A. N., Mangalam, A., Simonsen, J., ... Peijs, T. (2010). Review:
 Current international research into cellulose nanofibres and nanocomposites. *Journal of*

- 478 *Materials Science*, 45(1), 1–33. https://doi.org/10.1007/s10853-009-3874-0
- Elbaum, R., Zaltzman, L., Burgert, I., & Fratzl, P. (2007). The role of wheat awns in the seed dispersal unit. *Science*, *316*(5826), 884–886. https://doi.org/10.1126/science.1140097
- 481 Erb, R. M., Sander, J. S., Grisch, R., & Studart, A. R. (2013). Self-shaping composites with
 482 programmable bioinspired microstructures. *Nature Communications*, 4, 1–8.
 483 https://doi.org/10.1038/ncomms2666
- 484 Ganewatta, M. S., Wang, Z., & Tang, C. (2021). Chemical syntheses of bioinspired and biomimetic
 485 polymers toward biobased materials. *Nature Reviews Chemistry*, 5(11), 753–772.
 486 https://doi.org/10.1038/s41570-021-00325-x
- 487 Gupta, K. C., & Jabrail, F. H. (2006). Effects of degree of deacetylation and cross-linking on physical
 488 characteristics, swelling and release behavior of chitosan microspheres. *Carbohydrate Polymers*,
 489 66(1), 43–54. https://doi.org/10.1016/j.carbpol.2006.02.019
- Habibi, Y. (2014). Key advances in the chemical modification of nanocelluloses. *Chemical Society Reviews*, 43(5), 1519-1542. Royal Society of Chemistry. https://doi.org/10.1039/c3cs60204d
- Huber, T., Müssig, J., Curnow, O., Pang, S., Bickerton, S., & Staiger, M. P. (2011). A critical review
 of all-cellulose composites. *Journal of Materials Science*, 47(3), 1171–1186.
 https://doi.org/10.1007/S10853-011-5774-3
- Inphonlek, S., Sunintaboon, P., Léonard, M., & Durand, A. (2020). Chitosan/carboxymethylcellulose stabilized poly(lactide-co-glycolide) particles as bio-based drug delivery carriers. *Carbohydrate Polymers*, 242, 116417. https://doi.org/10.1016/j.carbpol.2020.116417
- Kuang, Y., Chen, C., Cheng, J., Pastel, G., Li, T., Song, J., Jiang, F., Li, Y., Zhang, Y., Jang, S. H.,
 Chen, G., Li, T., & Hu, L. (2019). Selectively aligned cellulose nanofibers towards highperformance soft actuators. *Extreme Mechanics Letters*, 29, 100463.
 https://doi.org/10.1016/j.eml.2019.100463
- Kulpinski, P., Jeszka, J. K., Malolepszy, A., & Stobinski, L. (2022). Regenerated Cellulose/Graphene
 Composite Fibers with Electroconductive Properties. *Autex Research Journal*, 22(2), 177–183.
 https://doi.org/10.2478/aut-2020-0027
- Le Duigou, A., Keryvin, V., Beaugrand, J., Pernes, M., Scarpa, F., & Castro, M. (2019). Humidity
 responsive actuation of bioinspired hygromorph biocomposites (HBC) for adaptive structures.
 Composites Part A, *116*, 36–45. https://doi.org/10.1016/j.compositesa.2018.10.018
- Leray, N., Talantikite, M., Villares, A., & Cathala, B. (2022). Xyloglucan-cellulose nanocrystal-*chitosan double network hydrogels for soft actuators*, 29, 119753.
 https://doi.org/10.1016/j.carbpol.2022.119753
- Li, Q., & Renneckar, S. (2011). Supramolecular structure characterization of molecularly thin
 cellulose I nanoparticles. *Biomacromolecules*, 12(3), 650–659.
 https://doi.org/10.1021/bm101315y
- Liu, Y., Shang, S., Mo, S., Wang, P., Yin, B., & Wei, J. (2021). Soft actuators built from cellulose
 paper: A review on actuation, material, fabrication, and applications. *Journal of Science: Advanced Materials and Devices*, 6(3), 321–337. https://doi.org/10.1016/j.jsamd.2021.06.004
- Lombardo, S., Cathala, B., & Villares, A. (2021). Adsorption of biopolymers onto nanocelluloses for
 the fabrication of hollow microcapsules. *Nordic Pulp and Paper Research Journal*, *36*(4), 651–
 661. https://doi.org/10.1515/npprj-2021-0040
- Lombardo, S., & Thielemans, W. (2019). Thermodynamics of adsorption on nanocellulose surfaces.
 Cellulose, 26(1), 249–279. https://doi.org/10.1007/s10570-018-02239-2
- 522 Lopes da Costa, L., Moreau, C., Lourdin, D., Cathala, B., & Villares, A. (2023). Shape-recovery in

- 523 organic solvents of water-responsive cellulose nanofiber actuators. *Cellulose, submitted*.
- Missoum, K., Belgacem, M. N., Barnes, J. P., Brochier-Salon, M. C., & Bras, J. (2012).
 Nanofibrillated cellulose surface grafting in ionic liquid. *Soft Matter*, 8(32), 8338–8349.
 https://doi.org/10.1039/c2sm25691f
- Moon, R. J., Martini, A., Nairn, J., Simonsen, J., & Youngblood, J. (2011). Cellulose nanomaterials
 review: structure, properties and nanocomposites. *Chemical Society Reviews*, 40(7), 3941–3994.
 https://doi.org/10.1039/c0cs00108b
- Oh, S. Y., Dong, I. Y., Shin, Y., Hwan, C. K., Hak, Y. K., Yong, S. C., Won, H. P., & Ji, H. Y. (2005).
 Crystalline structure analysis of cellulose treated with sodium hydroxide and carbon dioxide by
 means of X-ray diffraction and FTIR spectroscopy. *Carbohydrate Research*, 340(15), 2376–2391. https://doi.org/10.1016/j.carres.2005.08.007
- Qian, J., Wang, X., Shu, J., Su, C., Gong, J., Xu, Z., Jin, J., & Shi, J. (2018). A novel complex of
 chitosan-sodium carbonate and its properties. *Marine Drugs*, 16(11), 1–12.
 https://doi.org/10.3390/md16110416
- Rosca, C., Popa, M. I., Lisa, G., & Chitanu, G. C. (2005). Interaction of chitosan with natural or
 synthetic anionic polyelectrolytes. 1. The chitosan-carboxymethylcellulose complex.
 Carbohydrate Polymers, 62(1), 35–41. https://doi.org/10.1016/j.carbpol.2005.07.004
- Rüggeberg, M., & Burgert, I. (2015). Bio-Inspired wooden actuators for large scale applications. *PLoS ONE*, 10(4), 1–16. https://doi.org/10.1371/journal.pone.0120718
- Sarwar, A., Katas, H., Samsudin, S. N., & Zin, N. M. (2015). Regioselective sequential modification
 of chitosan via azide-alkyne click reaction: Synthesis, characterization, and antimicrobial activity
 of chitosan derivatives and nanoparticles. *PLoS ONE*, 10(4).
 https://doi.org/10.1371/journal.pone.0123084
- Schwanninger, M., Rodrigues, J. C., Pereira, H., & Hinterstoisser, B. (2004). Effects of short-time
 vibratory ball milling on the shape of FT-IR spectra of wood and cellulose. *Vibrational Spectroscopy*, *36*(1), 23–40. https://doi.org/10.1016/j.vibspec.2004.02.003
- Vleugels, L. F. W., Ricois, S., Voets, I. K., & Tuinier, R. (2018). Determination of the 'apparent pKa' of selected food hydrocolloids using ortho-toluidine blue. *Food Hydrocolloids*, *81*, 273–283. https://doi.org/10.1016/j.foodhyd.2018.02.049
- Wang, M., Tian, X., Ras, R. H. A., & Ikkala, O. (2015). Sensitive Humidity-Driven Reversible and
 Bidirectional Bending of Nanocellulose Thin Films as Bio-Inspired Actuation. Advanced
 Materials Interfaces, 2(7), 1500080. https://doi.org/10.1002/admi.201500080
- Wang, N., Chen, Y., & Kim, J. (2007). Electroactive paper actuator made with chitosan-cellulose
 films: Effect of acetic acid. *Macromolecular Materials and Engineering*, 292(6), 748–753.
 https://doi.org/10.1002/mame.200700020
- Wang, Y., Huang, W., Huang, W., Wang, Y., Mu, X., Ling, S., Yu, H., Chen, W., Guo, C., Watson,
 M. C., Yu, Y., Black, L. D., Li, M., Omenetto, F. G., Li, C., & Kaplan, D. L. (2020). Stimuliresponsive composite biopolymer actuators with selective spatial deformation behavior. *Proceedings of the National Academy of Sciences of the United States of America*, 117(25),
 14602–14608. https://doi.org/10.1073/pnas.2002996117
- Wei, J., Jia, S., Guan, J., Ma, C., & Shao, Z. (2021). Robust and Highly Sensitive Cellulose NanofiberBased Humidity Actuators. ACS Applied Materials and Interfaces, 13(45), 54417–54427.
 https://doi.org/10.1021/acsami.1c17894
- Wu, M., Sukyai, P., Lv, D., Zhang, F., Wang, P., Liu, C., & Li, B. (2020). Water and humidityinduced shape memory cellulose nanopaper with quick response, excellent wet strength and
 folding resistance. *Chemical Engineering Journal*, 392, 123673.

- 569 https://doi.org/10.1016/j.cej.2019.123673
- Yang, L., Cui, J., Zhang, L., Xu, X., Chen, X., & Sun, D. (2021). A Moisture-Driven Actuator Based
 on Polydopamine-Modified MXene/Bacterial Cellulose Nanofiber Composite Film. *Advanced Functional Materials*, *31*(27), 1–11. https://doi.org/10.1002/adfm.202101378
- Zhang, L., Chizhik, S., Wen, Y., & Naumov, P. (2016). Directed Motility of Hygroresponsive
 Biomimetic Actuators. *Advanced Functional Materials*, 26(7), 1040–1053.
 https://doi.org/10.1002/adfm.201503922
- Zhang, Y., Nypelö, T., Salas, C., Arboleda, J., Hoeger, I. C., & Rojas, O. J. (2013). Cellulose
 nanofibrils: From strong materials to bioactive surfaces. *Journal of Renewable Materials*, 1(3),
 195–211. https://doi.org/10.7569/jrm.2013.634115
- Zhao, K., Wang, W., Teng, A., Zhang, K., Ma, Y., Duan, S., Li, S., & Guo, Y. (2020). Using cellulose
 nanofibers to reinforce polysaccharide films: Blending vs layer-by-layer casting. *Carbohydrate Polymers*, 227, 115264. https://doi.org/10.1016/j.carbpol.2019.115264
- Zhu, X., Yang, C., Jian, Y., Deng, H., Du, Y., & Shi, X. (2022). Ion-responsive chitosan hydrogel
 actuator inspired by carrotwood seed pod. *Carbohydrate Polymers*, 276, 118759.
 https://doi.org/10.1016/j.carbpol.2021.118759

585