High Molecular Weight Mixed-Linkage Glucan as a Mechanical and Hydration Modulator of Bacterial Cellulose: Characterization by Advanced NMR Spectroscopy

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ABSTRACT: Bacterial cellulose (BC) consists of a complex three-dimensional organization of ultrafine fibers which provide unique material properties such as softness, biocompatibility, and water-retention ability, of key importance for biomedical applications. However, there is a poor understanding of the molecular features modulating the macroscopic properties of BC gels. We have examined chemically pure BC hydrogels and composites with arabinoxylan (BC−AX), xylloglucan (BC−XG), and high molecular weight mixed-linkage glucan (BC−MLG). Atomic force microscopy showed that MLG greatly reduced the mechanical stiffness of BC gels, while XG and AX did not exert a significant effect. A combination of advanced solid-state NMR methods allowed us to characterize the structure of BC ribbons at ultra-high resolution and to monitor local mobility and water interactions. This has enabled us to unravel the effect of AX, XG, and MLG on the short-range order, mobility, and hydration of BC fibers. Results show that BC−XG hydrogels present BC fibrils of increased surface area, which allows BC−XG gels to hold higher amounts of bound water. We report for the first time that the presence of high molecular weight MLG reduces the density of clusters of BC fibrils and dramatically increases water interactions with BC. Our data supports two key molecular features determining the reduced stiffness of BC−MLG hydrogels, that is, (i) the adsorption of MLG on the surface of BC fibrils precluding the formation of a dense network and (ii) the preorganization of bound water by MLG. Hence, we have produced and fully characterized BC−MLG hydrogels with novel properties which could be potentially employed as renewable materials for applications requiring high water retention capacity (e.g. personal hygiene products).

INTRODUCTION

Bacterial cellulose (BC) is produced by Gram-negative bacteria, such as Komagataeibacter xylinus (previously known as Acetobacter xylinum or Gluconacetobacter xylinum), and is chemically identical to plant cellulose (β-1,4-glucan) but has a different structural organization. Hence, unlike in plants, BC is secreted by bacteria as a pure polysaccharide into the surrounding environment and is hierarchically assembled into ribbons. While BC is synthesized in the absence of non-cellulosic components and consists of microfibrils with cross-sections of 4–8 nm, cellulose microfibrils of plant cell walls (PCWs) are synthesized embedded in a matrix of amorphous noncellulosic polymers and present smaller cross-sections (2–5 nm). The production (yield and properties) of BC can be modified during biosynthesis by altering the culture conditions. BC can also be modified in situ by adding carbon nanotubes, lignosulfonate, polyaniline, and water soluble polysaccharides (e.g. hemicelluloses) to the growth media during biosynthesis. The added compounds become entrapped in the forming cellulose gel, modulating its macroscopic features such as morphology, crystallinity index, water uptake capability, or electroconductivity. The ability to easily modify the gel properties of BC makes it an attractive biopolymer in the...
pharmaceutical, food, and cosmetic industries and in biomedical applications such as drug delivery.11 BC has also been used as a scaffold to study polysaccharide–polysaccharide interactions and as a model system for PCWs.1,6,10,12−15 In particular, there are strong resemblances between hemicellulose–BC and hemicellulose–PCW interactions.14,16 For instance, it has been shown that hemicellulose interactions with BC can modulate the material properties of BC hydrogels.1,8,17,18 Attempts to better understand the effect of the incorporation of individual PCW polysaccharides such as mannan19 arabinoxylan (AX),9 xylglucan (XG),1 mixed-link glucan (MLG),9 and pectins16,20 have been reported. Some studies have suggested that only XG and mannans are able to interfere with the cellulose crystallization process by intercalating between cellulose chains, thus reducing the crystallinity index and promoting the formation of the cellulose Iβ allomorph and reducing significantly gel stiffness.8,17,19 In addition, a different population of XG, which binds to the surface of cellulose ribbons, has been identified. This fraction is thought to correspond to the thin XG cross-bridges detected in the microscopy images from composite hydrogels.18 On the other hand, it has been reported that AX and MLG only affect the mechanical properties of BC composite hydrogels in a minor extent compared to XG and pectin. Based on these results, it was proposed that AX and MLG bind only to the surface of cellulose ribbons, following an entropically driven mechanism.1,8,9,15,21 However, none of the mentioned studies have investigated hydrogel network dynamics or water behavior in relation to the mechanical properties of BC hydrogels. These are important understudied characteristics of BC hydrogels due to their high water holding capacity (ca. 99 wt % H2O), which is a key aspect of many applications of BC hydrogels.1,22 For example, in commercial biomedical applications of BC hydrogels, the water holding and adsorptive capabilities of the materials are key to their applications as wound dressings,23,24 but the dynamics of water within BC hydrogels, and how to modify this behavior, have received little research interest.

Herein, by following a combined approach including advance solid-state NMR, X-ray diffraction (XRD), rheology, and microscopy, we have unraveled the molecular features responsible for the material properties of BC composites with the noncellulosic polysaccharides AX, XG, and high molecular weight MLG. To do so, we have studied hydrated 13C-labeled cellulose fibrils in the presence and absence of nonlabeled hemicelluloses and investigated water–polysaccharide interactions on the material properties of these hydrogels. In particular, we have characterized (i) the degree of short
(NMR) and long range [powder XRD (PXRD)] ordering, (ii) the crystalline allomorphism of BC, (iii) the local dynamics of the interior and partially disordered surface domains of BC ribbons at different time scales, (iv) the mobility of XG and MLG within the gel, and (v) the interactions of water with the BC particle network at site-specific resolution. To the best of our knowledge, the findings herein characterize BC gels at the highest reported resolution, and for the first time, we demonstrate that high molecular weight MLG reduces the mechanical strength of BC hydrogels and acts as a water-structuring coadjuvant for BC. The proposed methodology assessing structure, dynamics, and properties of BC hydrogels at different lengths scales and focusing on the role of hydration in directing functional applications of carbohydrate-based hydrogels forms a tool-kit which should be valuable for understanding other classes of “soft” biomaterial systems.

## MATERIALS AND METHODS

**Production of BC Composite Hydrogels.** *Gluconacetobacter xylinus* (formerly *Klebsiella xylinus*, strain ATCC 53524) was cultivated in Hestrin and Schramm medium (pH 5.0) containing 2% (w/v) glucose (Merck, UK) or 2% (w/v) 13C-labeled glucose (Cambridge Isotope Laboratories Inc., USA) and 2% (w/v) 15N-labeled ammonium chloride (Cambridge Isotope Laboratories Inc., USA). BC—hemicellulose composites were produced by adding medium viscosity (31 cSt) wheat AX (Megazyme International Ireland), tamarind seed XG (Megazyme International Ireland), or high-molecular weight (650 000 Da) MLG (Nofima, Tromsø, Norway) (Figure 1, Table 1) to the medium at a concentration of 0.5% (w/v). Incubations were performed under static conditions at 30 °C for 72 h and thereafter washed with 0.02% (w/v) sodium azide in 0.1 M NaOH (24 h), followed by two washes (24 h) with 0.02% (w/v) sodium azide. The washing steps were done at 30 °C by gentle agitation (50 rpm).

**Compositional Analysis.** Hemicellulose incorporation into the various BC composites was determined as previously described by Pettolino et al. (2012) with the following modifications. Briefly, for each composite material in duplicate, the freeze-dried sample (5 mg) was hydrolyzed at room temperature for 1 h in 12 M H2SO4 (200 μL), then diluted with Milli-Q water (2.75 mL), and incubated at 100 °C for 3 h. Thereafter, samples were cooled and neutralized with NH4OH (500 μL); the internal standard (5 mg of myo-inositol) was added to each sample and then dried under a stream of nitrogen. Hydrolyzed samples were reduced with 200 μL NaBD4 in dimethyl sulfoxide (20 mg mL−1) at 40 °C for 1.5 h. Acetic acid (20 μL) was added to destroy the reductant and then acetylated with 1-methylimidazol (25 μL) and acetic anhydride (250 μL). After 10 min, Milli-Q water (2 mL) was added, followed by dichloromethane (DCM—1 mL) to extract the alditol acetates. After mixing, samples were centrifuged to aid phase separation and the DCM phase was washed twice with Milli-Q water (2 mL), after which it was dried under a stream of nitrogen, reconstituted to 500 μL of DCM and analyzed by gas chromatography attached to a mass spectrophotometer on a high polarity BPX70 column.

**Total Moisture Content Analysis.** Moisture content of all BC hydrogel samples was measured using a TA Instruments (TA Instruments Ltd., New Castle, USA) thermogravimetric analyzer (TG) Discovery series TGA5500, using TA Instruments standard open aluminum pans (reference number 900786.901), where 3–5 mg samples were excised with minimal deformation using a surgical blade and loaded onto pretared pans. No initial weight equilibration step was used, in order to minimize the water evaporation from BC hydrogels prior to the experimental procedure. The change in sample weight was recorded at increments of 0.05 s in the temperature range of 25–200 °C, at a heating rate of 5 °C/min. The point of complete moisture loss was determined as the lowest temperature at which no further significant mass loss was observed, which for all samples was in the range of 90–150 °C. The total moisture content was determined as the mass difference (% w/w) from the start of the experiments to the point of complete moisture loss for each sample. Data presented is the average of a minimum of three replicates.

**Dynamic Oscillatory Rheology.** Dynamic rheology was performed on a TA Instruments (TA Instruments Ltd., New Castle, USA) Rheometer AR2000, with 13 mm parallel plate geometry, equipped with a Peltier device for temperature control, where hydrogel samples were excited using a 13 mm cork borer (Breckland Scientific Supplies Ltd., Stafford, UK) with minimal manipulation and excess sodium azide (0.02% w/v) solution was blotted using Kimtech precision wipes (Kimberly-Clark Professional). BC gels were loaded onto the sensor plate and left for 2 min to allow the temperature to equilibrate, minimizing the impact of loading effects. Samples were analyzed at a constant temperature of 5 °C and the interplate gap was set to 2.0 mm. All rheological data were analyzed using TA Data Analysis software package, establishing the linear viscoelastic (LVE) region, the storage and loss moduli (G’ and G″, respectively) and phase angle tangent (tan δ) of all hydrogel samples.

Strains were spread in the range of 0.01–100% at a constant frequency of 0.1 Hz to ascertain the materials’ LVE range. This was followed by frequency sweep analyses in the range of 0.01–10.0 Hz at a constant strain of 0.025% to determine the behavior of each sample’s individual storage modulus (G’), loss modulus (G″), and phase angle tangent (tan δ) as a function of angular oscillation frequency. All measurements were taken at regular time intervals, and the data presented are averages of a minimum of three runs.

**BC Gel Thickness.** BC hydrogel thickness was measured using a TA Instruments (TA Instruments Ltd., New Castle, USA) Rheometer AR2000, with 13 mm parallel plate geometry, where excited samples were loaded onto the Peltier device, followed by the lowering of the upper parallel plate until contact was established across the entirety of the upper plate, as detected by the rheometer. Sample thickness was determined as the gap distance between the upper and the Peltier device at the point of upper plate contact with the sample. Data presented is the average of a minimum of three measurements on the same gel.

**X-ray Diffraction.** XRD (Thermo ARL, Xtra fitted with Cu X-ray tube, 45 kV, 40 mA) with Cu Kα radiation (λ = 0.154 nm) was used to examine the crystal structure of the BC, XG, AX, MLG alone, and their blend. Diffraction angles (2θ) ranging from 5 to 55° were employed. All the hydrogels were analyzed in the freeze-dried form and cut into small pieces. XG, AX, and MLG were analyzed in the powder form.

**Scanning Electron Microscopy.** Hydrogels were placed into a solution of 2.5% glutaraldehyde in 0.1 M PIPES buffer (pH 7.4) and fixed overnight. After washing with 0.1 M PIPES buffer, the gel was cut into approximately 5 mm3 pieces, inserted into a critical point drying capsule, and dehydrated in a series of ethanol solutions (30, 50, 70, 80, 90, 100%). The gels were then dried in a Leica EM CPD300 Critical Point Dryer using liquid carbon dioxide as the transition fluid. The dry gel material was carefully torn across using a razor blade and tweezers and was mounted with the freshly torn surface facing upward onto silicon microtome (SEM) stubs using conductive silver paint. The samples were coated with...
gold in an Agar high resolution sputter-coater apparatus. SEM was carried out using a Zeiss Supra 55 VP FEG SEM, operating at 3 kV.  

**Atomic Force Microscopy.** The atomic force microscope used in this study was an MFP-3D BIO (Asylum Research, Goleta, CA, USA). The experimental data were captured in a so-called force–volume mode using a colloidal CP-SPNP-SiO probe with a 2 μm sphere in PBS (at a rate of 2 μm/s in the Z direction and at a scan rate of 1 Hz and a pixel density of 32 × 32). In this mode, the instrument ramps the Z piezo element of the scanner by a predetermined amount at each sample point over a selected scan area (3 μm) and records the subsequent deflection of the cantilever as it is pushed into (maximum load forces of 10, 20 or 40 nN) and then retracted away from the sample surface. This produces a matrix of 1024 force versus distance curves for each tip–sample combination. The spring constant, k, of the cantilevers was determined by fitting the thermal noise spectra,30 yielding typical values in the range 0.01–0.04 N/m. Young's modulus for the gels were determined by fitting the Hertz model of contact mechanics.30 The Hertz model has proven sufficient for modeling of atomic force microscopy (AFM) data on soft biological samples.31 Its use is favored here by the fact that it does not require prior knowledge of parameters like surface energy, which are difficult to measure accurately for the kinds of surfaces (often microheterogeneous) common in biological systems. In the current application, the force–distance plots extracted from force–volume datasets of the BC and BC–hemicellulose gels and glass, an ideally hard reference, are compared in order to determine the indentation, δ, of the tip into the gels at a given force. The indentation at a given force is defined as the depth of penetration of the tip into the sample surface. It is calculated with reference to an ideally hard surface, in this case glass, where the indentation is set at 0. The relationship between loading force F and indentation δ is then given by eq 1

\[ F = \frac{4E \sqrt{R}}{3(1 - \mu^2)} \delta^{3/2} \]  

where E is Young's modulus, determined as a fitting parameter, R is the AFM tip radius (given above as 2 μm), and μ is the Poisson ratio for the deformable gel, here taken to be 0.5.  

**Solid-State NMR.** Solid-state NMR experiments were carried out using either a Bruker AVANCE III 400 MHz spectrometer equipped with a HX4 4 mm probe or a Bruker AVANCE III 850 MHz solid-state NMR spectrometer (UK National 850 Solid-State NMR facility at Warwick) equipped with a 3.2 mm HX4 low E field H13900 probe. Experiments were acquired at MAS spinning rates of 6 and 12 kHz when using the 4 and 3.2 mm probes, respectively. Tetramethylsilane was used as an external chemical shift reference for 1H and 13C. π/2 pulses of 3.5 and 4.5 μs were used for 1H and 13C, respectively. A relaxation delay of 1 and 100 s were used for mobility-enhanced and quantitative 13C CP experiments, respectively, and 10 s for 1H–13C cross polarization (CP) experiments.  

13C–13C refocused CP-INADEQUATE experiments on 13C-labeled gels were carried out on the 850 MHz spectrometer at Warwick, at 12 kHz magic angle spinning. A relaxation delay of 2 s and a 27 spin–echo evolution time for a (τ–π–τ) spin–echo of 5 ms were employed. All the experiments were carried out at 25 °C and/or 5 °C.  

Determination of the surface-to-interior (s/i) ratio and percentage of \( I_f \) allomorph (% \( I_f \)) of BC hydrogels by 13C quantitative direct polarization (DP) and 1H–13C CP NMR experiments.  

DP experiments using high power 1H decoupling were carried out using a 100 s relaxation delay to obtain quantitative 13C spectra, taking into account that the maximum 13C T2, times measured were 12 s (Table S7). The surface-to-interior (s/i) ratio of BC was derived by spectral deconvolution of the C4 peaks corresponding to the crystalline interior (\( A_{C4} \)) and partially disordered surface (\( A_{C4} \)) domains of cellulose (Figure 3) and using the following equation

\[ \text{s/i} = \frac{A_{C4}}{A_{C4}} \]  

For the quantification of BC allomorphs among different hydrogels, the percentage of \( I_f \) allomorph (% \( I_f \)) was obtained by deconvolution of the 1H–13C CP NMR spectra acquired at 20 T, using the following equation32

\[ \% I_f = \frac{A_{C4} + A_{C4} + A_{C4} + A_{C4}}{A_{C4} + A_{C4} + A_{C4} + A_{C4}} \]  

The experiments were run on 13C-labeled BC, BC–AX, and BC–ML gels and unlabeled BC–XG gel, the latter due to large broadening (strong 13C homonuclear dipolar coupling) of the labeled gel precluding differentiation of \( I_f \) and \( I_p \).

**Data Analysis of CP Kinetics Curves.** Different models for the analysis of CP growth over contact time, also called CP kinetics, have been developed. The simplest model of CP kinetics (1–S) was derived for homogeneous solids where the 1–S heteronuclear interactions are relatively weak and the 1–1 homonuclear dipolar interactions are strong to provide efficient spin diffusion. For a system of abundant (1) and dilute (S) 1/2 spin nuclei, provided that the rate of 1–S transfer of magnetization (\( T_{13}^{ff} \)) is faster than the relaxation rate in the rotating frame of the 1 spins (\( T_{13}^{ff} \)), then, \( T_{13}^{ff}/T_{13}^{ff} \rightarrow 0 \), the 1–S model leads to the kinetic equation

\[ I(t) = I_0 \left( 1 - \frac{T_{13}}{T_{13}} - t \right) \left( \exp \left( -t/T_{13} \right) - \exp \left( -T_{13}/T_{13} \right) \right) \]  

where \( I(t) \) is the normalized peak intensity, \( I_0 \) is the absolute amplitude, \( T_{13} \) is the relaxation time of the abundant spin in the rotating frame, and \( T_{13} \) is the CP time constant.

The fit of CP kinetics data to eq 4 allowed us to obtain \( T_{13}^{ff} \) and \( T_{13}^{ff} \) values for BC rigid domains in the absence and presence of AX, XG, and MLG. \( T_{13}^{ff} \) reports on the efficiency of 1H–13C magnetization transfer and, therefore, on the rigidity of the individual C–H vectors (the lower the \( T_{13}^{ff} \) parameter, the higher the rigidity). The fit of the experimental data was carried out with QtiPlot.  

**Mobility-Enhanced 13C DP Experiments for the Determination of BC-Disordered Domains.** An interesting application of 13C DP experiments is to enhance the presence of mobile rigid components in the spectra.30 This is achieved by employing a very short relaxation delay (1–2 s). To obtain a rough estimation of the mobility of the surface (s) relative to the interior domains (i) of BC in the different hydrogels, we have defined a mobility index (MI) alike the s/i ratio shown in eq 2 but using 13C direct polarization experiments carried out using a very short relaxation delay

\[ \text{MI} = \frac{A_{C4}}{A_{C4} + 100} \]  

where \( A_{C4} \) and \( A_{C4} \) are the areas of the C4 peaks corresponding to surface and interior BC domains, respectively, from 13C DP spectra recorded using 1 s relaxation delay. These experiments were acquired using a 20 T spectrometer operating at 12 kHz MAS rate.  

**Water Polarization Transfer CP.** The water polarization transfer CP experiment (WPT-CP) is a very powerful NMR technique to gain insights on the hydration profile of solid particles at the atomistic level.31 It starts with a \( T_1 \) filter that is optimized to remove the transverse magnetization of the immobilizing components of the system (e.g. BC fibrils), while keeping most of the \( T_1 \) magnetization of the mobile constituents (e.g. water). This \( T_1 \) magnetization is then transferred to the immobile components by spin diffusion and chemical exchange during the mixing time period. Finally, a CP building block transfer \( T_1 \) magnetization to \( ^{13}C \) for detection.  

A short contact time (500 μs) was used to reduce the contribution of spin diffusion during CP, so that the acquired signal is mostly affected by water proximity and mobility.32 It should be noted that the \( T_1 \) filter must be first optimized to reduce the less mobile peaks below 5% of their initial intensity. A \( T_1 \) filter of 2 ms was sufficient for our gels. Mixing times ranging from 1 to 324 ms were employed. The peak

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intensities at each mixing time were normalized against the spectrum giving the highest intensity peaks, that is, acquired with the longest mixing time (324 ms). By representing the normalized growth of peak area against the square root of mixing time, the spin diffusion profile of hydration at each $^1$H–$^{13}$C site is obtained. All WPT-CP experiments were acquired at 293 K.

**RESULTS AND DISCUSSION**

**Production and Molecular Analysis of BC Composites in the Presence of AX, XG, and MLG.** Pure BC, as well as composite gels containing AX, XG, and MLG were successfully produced using aerobic cultures of *K. xylinus*. The thickness of the gels produced was variable (Table S1); however, the water content was similar for all gels (98–99% w/w) (Tables 2 and S2). In particular, the incorporation of XG and MLG reduced the water content was similar for all gels (98–99% w/w) (Tables 2 and S2). In particular, the incorporation of XG and MLG reduced

**Table 2. Composition and Thicknesses for the BC Gel and BC Composite Gels**

<table>
<thead>
<tr>
<th></th>
<th>hemicellulose incorporation (% w/w)</th>
<th>moisture content (% w/w ± S.D.)</th>
<th>temperature at point of complete water loss (°C ± S.D.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BC</td>
<td>n.a.</td>
<td>98.4 ± 0.4</td>
<td>117.0 ± 9.9</td>
</tr>
<tr>
<td>BC–AX</td>
<td>10.2</td>
<td>98.4 ± 0.8</td>
<td>109.0 ± 15.5</td>
</tr>
<tr>
<td>BC–XG</td>
<td>14.0</td>
<td>98.7 ± 0.5</td>
<td>141.7 ± 5.8</td>
</tr>
<tr>
<td>BC–MLG</td>
<td>n.a.*</td>
<td>98.7 ± 0.3</td>
<td>135.7 ± 6.0</td>
</tr>
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</table>

“Standard deviations are from triplicate measurements. Not applicable (n.a.).” Incorporation data could not be determined as the BC–MLG gels only contain glucose.

S2). In particular, the incorporation of XG and MLG reduced gel thickness 2- to 3-fold (Table S1), which is in agreement with other studies.8 Monosaccharide analysis revealed that there was a higher level of incorporation of XG into the BC gels than AX (Tables 2 and S3), in line with previous work.35 In this study, MLG was not quantified as the monosaccharide analysis used cannot differentiate glucose from MLG and BC. However, CPSP NMR experiments provided clear evidence of MLG incorporation into the gel composite (Figure S5b).

**Microstructure and Mechanical Properties of BC Composites with AX, XG, and MLG.** *XRD and SEM.* PXRD analysis was used to investigate the effect of additive polysaccharides (AX, XG, and MLG) on BC crystallinity and structure at the long-range length scale. BC and BC–AX gels yielded well-defined diffraction peaks (Figure S2), indicating that ordered BC fibrils were formed. On the other hand, XG gave rise to the formation of the most disordered BC fibrils, followed by BC–MLG gels (increased peak broadening in the PXRD profile, Figure S2). No diffraction patterns were observed for the hemicellulose components, indicating they are disordered polymers (Figure S2).

SEM images of pure BC gels exhibited the mesh-like network structure of cellulose microfibrils typical of BC (Figure S3). A similar mesh-like structure can be seen for each of the composite hydrogels. There were no obvious differences observed between the composites and the pure BC. Previous authors have shown evidence of filamentous cross-links between cellulose microfibrils in composite gels produced with XG.8 These could not be clearly seen in the present study. AX aggregates were not observed for the BC AX gels. The ca. 2 μm oval structures observed in some images are residual bacterial cells.

**Rheological and AFM Assessment of Gel Rigidity.** The micromechanics of the individual cellulose fibers and interactions with hemicelluloses affect the rheology of the bulk gel.8 The visual counting of 2D AFM maps (Figures S4–S7) was carried out for three different probe forces applied (10, 20 and 40 nN). The results (Figure 2) show a very similar strength for BC, BC–AX, and BC–XG (within the standard deviation), but a significantly ($p < 0.05$) lower stiffness for BC–MLG. This is in contrast with the oscillatory shear rheological data, which shows that the $G'$ values for the gels decrease in the order BC > BC–AX > BC–MLG > BC–XG (Figure S8). However, the rheological behavior of cellulose hydrogels is known to be highly dependent on the gel thickness, and the $G'$ values measured in this study correlate closely to gel thickness (Table S1). The micromechanics of the gel fibers measured by AFM (Figure 2) shows that for individual fibers or bundles of fibers, there is no significant difference in the mechanical properties of the BC, BC–AX, and BC–XG gels ($p > 0.05$). This finding is in agreement with recent AFM studies using “drag-and-dip” lateral force AFM experiments to probe adhesive forces and micromechanics of BC composites, which produced results suggesting that BC, BC–AX, and BC–XG have very similar micromechanical properties.36 Our observation that MLG significantly reduces the stiffness of cellulose fibers has, to the best of the authors’ knowledge, not been previously reported.9 More generally, the discrepancy between AFM-based and rheological data on the BC–MLG gels may simply reflect the fact that the AFM indentation experiments probe only the region at and immediately beneath the gel surface, whereas the rheological experiments reflect the behavior of a much deeper section (as shown by their dependence on the gel thickness).

**Local Structure and Dynamics of BC Composites with AX, XG, and MLG: Solid-State NMR Characterization.** Ordered and Disordered Domains in BC Hydrogels. Among the accumulated literature on the structural characterization of cellulose-based materials, different models to describe a cellulose fibril have been proposed, and different names are used to refer to their ordered and disordered regions.37 For BC and PCWs, it is common to describe their fibrils as particles containing amorphous (or partially disordered) surface domains and crystalline (or ordered) core or interior domains.34,38,39 Further, Wang and Hong proposed a more detailed model for PCW fibrils,38 which considers three regions in a BC microfibril, that is, the partially disordered surface (s), a surface-bound para-crystalline domain (b), and...
the crystalline core (c), where b and c form together the interior region (i). Thus, s and i represent the partially disordered and ordered domains, respectively. Herein, we have used this model for our NMR data analysis. It should be noted that, due to limitations of resolution, we could only differentiate some b and c peaks in our 2D NMR experiments. Hence, we mostly refer to s and i domains below, except when otherwise indicated.

Figure 3 shows the $^1$H–$^{13}$C CP/MAS NMR spectrum of $^{13}$C-labelled BC gel, acquired at room temperature, 20.0 T and 12 kHz MAS spinning rate. (c) 2D $^{13}$C–$^{13}$C CP-RINADEQUATE spectrum of $^{13}$C-labelled BC gel, acquired at room temperature, 20.0 T and 12 kHz MAS spinning rate. A 2τ spin-echo evolution time for a (τ–π–τ) spin-echo of 5 ms was employed.

Table 3. Surface-to-Interior Ratio (s/i) and Percentage of $I_\alpha$ Allomorph (% $I_\alpha$) Obtained by Spectral Deconvolution of iC4 and sC4 Peaks from Quantitative $^{13}$C–$^1$H CP NMR Spectra (9.4 T) and $^1$H–$^{13}$C CP Spectra (20 T) of $^{13}$C-Labeled BC Gels

<table>
<thead>
<tr>
<th></th>
<th>BC</th>
<th>BC–XG</th>
<th>BC–AX</th>
<th>BC–MLG</th>
</tr>
</thead>
<tbody>
<tr>
<td>s/i</td>
<td>1:4:1</td>
<td>1:2:9</td>
<td>1:3:8</td>
<td>1:3:8</td>
</tr>
<tr>
<td>% $I_\alpha$</td>
<td>64%</td>
<td>&lt;64%*</td>
<td>60%</td>
<td>56%</td>
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*The % $I_\alpha$ of BC–XG had to be qualitatively estimated from the CP spectrum on the unlabelled gel due to much greater broadening of the $^{13}$C-labeled gel spectrum precluding the differentiation of $I_\alpha$ and $I_\beta$ peaks (Figure S13).

has been suggested to indicate that AX and MLG are physically adsorbed onto the surface of BC, and are not involved in the fiber assembly. The increased stiffness of BC–XG reported by AFM correlates with the increased surface of BC fibers, possibly due to the interaction of XG with BC chains during assembly.

On the other hand, previous works have suggested that XG interactions with BC lead to the conversion of BC $I_\alpha$ domains into $I_\beta$. Hence, the percentage of $I_\alpha$ allomorph (% $I_\alpha$) was calculated from deconvolution of the iC4 peak (eq 3) on $^1$H–$^{13}$C CP spectra of $^{13}$C-labelled BC, BC–AX, and BC–MLG gels acquired at 20 T. For $^{13}$C-labeled BC–XG gel, however, the large spectral broadening due to strong $^{13}$C homonuclear dipolar coupling precluded the differentiation of $I_\alpha$ and $I_\beta$ peaks (Figure S13). Thus, we acquired the CP spectrum for the unlabelled gel (Figure S13). While global spectral deconvolution (GSD) worked in the pure BC, as shown previously, while no significant differences were observed for BC–AX and BC–MLG (Table 3, Figure S12). This finding is supported by previous studies,
a large reduction in the $I_\beta$ peak and an increase of $I_\alpha$ for BC–XG (Figure S13). Notably, the conversion of $I_\beta$ into the lower energy $I_\alpha$ state has been previously reported for BC–XG. The ability of XG to alter BC allomorphism has been related to direct association with cellulose chains close to the point of deposition before mature fibrillar ribbons are formed. On the other hand, a decrease in $I_\beta$ was observed for BC–AX (60%) and BC–MLG (56%) with respect to BC (64%). However, visual comparison of the spectra (Figure S13) suggests that these differences are not large enough to confidently make a statement on the effect of AX and MLG on BC allomorphism.

**Molecular Level Interactions between BC and XG.** $^1$H–$^{13}$C CP NMR spectra were acquired at ultra-high field for all $^{13}$C-labeled BC hydrogels. BC, BC–AX, and BC–MLG composites (Figure 4) showed similar linewidths, which is in agreement with AX and MLG not interfering significantly with the ordering of BC (Table 3) and with previously proposed models of physical adsorption of AX and MLG to the surface of BC particles. Importantly, the $^1$H–$^{13}$C CP and CPSP spectra of unlabeled BC–MLG showed peaks corresponding to C6 of $\beta$-(1–3) chains of MLG at 50.5, 60.2, and 60.9 ppm (Figure 5b, A6(1), A6(2), and A6(3) peaks, respectively). Spectral deconvolution of the A6(3) peak in both CP and CPSP spectra indicated that the CP contribution accounts for about 40% of the total BC–XG (red line) shows much broader peaks than the other 3 hydrogels, indicating intimate interactions between BC and XG.

Figure 4. Comparison of the $^1$H–$^{13}$C CP MAS NMR spectra of the $^{13}$C-labeled hydrogels acquired at 20.0 T, 12 kHz MAS, 1 ms contact time, and 2 s relaxation delay. The spectrum of BC–XG (red line) shows much broader peaks than the other 3 hydrogels, indicating intimate interactions between BC and XG.

Figure 5. Comparison of $^1$H–$^{13}$C CP (blue line) and CPSP (red line) NMR spectra of unlabeled BC composite hydrogels with (a) XG and (b) MLG. The CP and CPSP spectra of $^{13}$C-labeled BC hydrogels are shown as green and black lines, respectively, for comparison. The assignment of mobile XG and MLG peaks are indicated with arrows. All spectra were acquired at a $^{13}$C frequency of 213.8 MHz. For the purpose of clarity, the anomeric carbons are not shown in the spectra due to the very broad distribution of mobile C1 peaks of BC present in the CPSP spectrum.

The strong association between BC and XG could lead to broader BC peaks via (i) shorter $T_2$ times of BC and/or, most likely, (ii) increased disordered of BC fibrils as shown above (Table 3). One XG peak corresponding to C6 was observed in both CP and CPSP spectra (B6 peak at 60.3 ppm in Figure 5a). As the CP contribution accounts for about 45% of the total B6 peak area (CPSP spectrum), we can roughly estimate that about 45% of the total XG is bound to BC.

**Dynamic Regimes in BC Composite Hydrogels.** Global and local dynamics are key features to understand the structural and binding properties of biomolecules. However, the broad distribution of time scales of molecular motions that might occur in complex systems require to tackle them using several approaches specific for different time scales. We have employed (i) the $^1$H–$^{13}$C CP kinetics approach to probe for slow motions within the millisecond-to-second time scale, and (ii) $^{13}$C{$^1$H} DP with short recycle delay and CPSP experiments to monitor the fast motions (faster than the millisecond time scale) of the surface domains of BC and the hemicelluloses, respectively.

CP kinetics analysis demonstrated the presence of very rigid domains in BC, BC–AX, and BC–XG (very short $T_{\text{CH}}$ and very long $^1$H $T_{1,\text{H}}$ Tables 4 and S4). Interestingly, this observation not only applies for the interior but also for the...
surface regions. We could speculate that this population of highly rigid surface domains might represent the areas of fibril–fibril interactions, where the conformational dynamics of the surface chains of BC is restricted. 1H T1ρ times of BC could not be determined accurately from CP kinetics build-up curves due to extremely slow 1H T1ρ relaxation compared to the longer CP contact time probed (6 ms), but we can reliably claim that they are at least longer than 100 ms (Tables 4 and S4, Figures S15–S22). For BC–MLG, the experimental T1H value was about 6–7 times longer for all BC peaks (Tables 4 and S4), hence proving the presence of less rigid BC interior and surface domains in the presence of MLG (partial averaging of dipolar interactions). This is in agreement with the AFM results, which showed that BC–MLG composite gels are structurally weaker than BC-only gels (Figures 2 and S4–S7).

13C[1H] DP NMR experiments using a very short relaxation delay (1 s) allowed for the characterization of very mobile BC surface domains present in all gels. 33 The surface-to-interior mobility of BC increased in the presence of XG (MI from 79 to 92%), decreased for BC–AX (MI from 79 to 67%), and no significant change was observed for BC–MLG (Table 4, Figure S14). This is in agreement with the proposed intercalating interactions between BC and XG leading to a larger partially disordered surface (Table 3). 1 The MI reduction observed for BC–AX might be due to coating of BC ribbons’ surfaces by amorphous AX chains, as previously proposed. 7

While 13C-labeling of BC is a very powerful approach for the detailed characterization of BC hydrogels, it has the disadvantage of hiding the spectral features of other nonlabeled components (e.g. hemicelluloses) present in composite hydrogels. Aiming to characterize XG and MLG in our gels, we carried out CP and CPSP experiments for unlabeled BC–XG and BC–MLG hydrogels at ultra-high field (20 T). It should be noted that MLG and XG hemicellulose components are more mobile compared to BC fibers (Figure 5; ic4 and ic6 peaks of BC compared to MLG and XG peaks in the CPSP spectra). The dynamics of BC particles would lie within the ms-to-s timescale, as shown by CP kinetics (Tables 4 and S4). On the other hand, the B6 peak (60.3 ppm, corresponding to C6 of XG β-1,4 chains, was observed with increased intensity in the CPSP spectrum of BC–XG gel (Figure 5a). This peak was also visible in the CP spectrum (Figure 5a), we can roughly estimate an intermediate mobility regime (μs-to-ms time scale) for XG in BC–XG gels. XG is known to establish strong molecular association with nascent cellulose fibrils prior to the formation of the final ribbon-shaped fiber but also to be located at the surface of BC fibrils and act as a physical cross-linker. 10,13 This population of surface XG might be responsible of the mobility-enhanced intensity of the B6 peak (Figure 5a).

In addition, three peaks attributable to mobile domains were identified in the CPSP spectrum of unlabeled BC–MLG gel within the expected chemical shift region for C6 of MLG β-1,3 chains. 33 In particular, (i) a very mobile fraction at 59.6 ppm [Figure 5b, peak A6(1); no CP intensity, μs-to-ns time scale], (ii) a mobile population at 60.2 ppm [Figure 5b, peak A6(2); very small CP intensity, μs-to-ns time scale], and (iii) a population of intermediate mobility at 60.9 ppm [Figure 5b, peak A6(3); high CP intensity, ms-to-μs time scale]. To our knowledge, these peaks have never been individually assigned; however, due to their differences in mobility, we could speculate that they might correspond to different populations of BC-bound and free MLG. Another two very mobile peaks of MLG can be observed at 79.2 (C5) and 68.9 ppm (C4) in the CPSP spectrum only, in agreement with NMR assignment of fungal cell walls. 33 Also, despite complete overlapping with BC peaks, the mobile peak observed at 75.1 ppm matches the expected chemical shift for C2 of β-1,3 chains of MLG. 33

Probing Water–BC Interactions by WPT-CP NMR. 13C–13C WPT-CP represents a very powerful technique for monitoring interactions of water to cellulose particles at atomic resolution, as it has been previously shown for PCWs. 34 Also, WPT-CP build-up curves can indirectly report on the architecture of cellulose composites and thus provide structural insights on their function. 34 Recently, Kang et al. used 2D water-edited 13C–13C correlation CP experiments to show that polysaccharides containing β-(1–3), β-(1–4), and β-(1–6) glucans are better hydrated than α-(1–3) glucans and chitin in fungal cell walls. 35 However, they could not identify either the influence of cell walls polysaccharides on the hydration properties of BC or the specific polysaccharides presenting increased hydration. By carrying out WPT-CP experiments at different mixing times on pure 13C-labelled BC gels and composites with either XG, AX, or MLG, we have been able to characterize to which extent individual PCW polysaccharides affect the hydration of BC. It should be highlighted that, strictly speaking, WPT-CP curves report only on water interactions to rigid components as it relies on CP. Hence, the interactions of water molecules to the very mobile components present in BC gels (Table 4, MI; Figure 5) would be mostly invisible in WPT-CP experiments. The WPT-CP build-up curves acquired for the four 13C-labeled BC gels showed that BC fibrils hold a higher population of bound water molecules (faster WPT-CP growth) in the presence of XG and MLG but not AX (Figures 6 and S23–S28). The latter did not exert a significant effect on BC hydration in comparison to the control sample (BC only gel). This is hence in qualitative agreement with the data reported for fungal cell walls, which further validates the use of BC hydrogels as ideal mimics of PCWs. BC–XG and BC–MLG gels featured the shortest lag phase followed by a very fast build-up. This indicates closer proximity of water to the surface of BC fibrils and more efficient spin diffusion transfer at the BC–water interface (Figures 6 and S23–S28). The increased hydration of BC–XG gels is in agreement with our thermogravimetric analyses, showing a higher temperature of complete water loss for this gel (Tables 2 and S2), and correlates with the larger surface area of BC fibers (Table 3, s/ i) and greater mobility of BC surface domains (Table 4, MI). Hence, the effect of XG on the surface properties of BC fibrils.
is possibly responsible for the increased population of bound water contained in BC−XG gels.

For BC−MLG gels, the growth of the WPT-CP curves is similar to BC−XG (Figure 6), which correlates with a similar temperature of complete water loss (Tables 2 and S2). Unlike XG, it has been previously shown that MLG does not bind BC chains during the fibril assembly but adsorbs onto the already formed fibrils.9,21 Also, the presence of MLG did not impact either the surface ratio (Table 3) or mobility of the BC surface (Table 4). Thus, the increased binding of water to BC in BC−MLG gels is probably due to the greater hydrophilic nature of MLG increasing the fraction of structured water readily available for binding BC fibers. In this regard, our WPT-CP data further support the previously suggested mechanism of physical adsorption of the very hydrophilic MLG onto the surface of BC particles, where MLG would act as a “coating agent” of BC particles and as a source of prestructured water ready to interact with BC, thus reducing the entropic penalty of water−BC interactions.9

Table 5 shows a qualitative summary of the macroscopic and molecular scale features determined for the three BC composites in comparison with the BC-only hydrogel. BC−AX and BC presented a very similar macroscopic behavior, and this correlates with minor changes at the molecular scale. On the other hand, the reduced stiffness of BC−MLG composites and the differences in the macroscopic order and population of bound water observed for BC−XG and BC−MLG gels demonstrate the different mechanisms which can influence the gel structure. Reduced cellulose order does not directly influence cellulose gel fiber stiffness, which is instead related to BC ribbon mobility within the gel.

## CONCLUSIONS

The combined analysis of AFM, XRD, TGA, and solid-state NMR data has provided insights into the complex structural features governing the properties of cellulose hydrogels forming composites with different polysaccharides. The use of mobility-enhanced NMR methods together with water polarization transfer NMR experiments allowed us to differentiate core, surface-bound, and surface domains present in BC.

Our results indicate that BC ribbons are formed with increased long-range disorder when produced in the presence of XG-containing media, giving hydrogels with similar mechanical properties to BC-only gels (Table 5). At the molecular level, this is related to specific BC−XG interactions (strong peak broadening), greater BC surface area and surface mobility, and increased BC-bound water (Table 5). We propose that the association between BC and XG during fibril assembly increases gel disorder, giving rise to a looser fibrillar packing. Although it is tempting to correlate reduced network particle density to weaker gels, the increased population of bound water contained in BC−XG gels might compensate for the effect of looser fibril network on gel strength.

For the first time, we report that high molecular weight MLG increases long-range disorder in BC, producing weaker BC hydrogels. However, MLG did not affect the surface area of BC ribbons but it increased their mobility and boosted water interactions with the fibrillar network. This is in agreement with the previously proposed mechanism of MLG physical adsorption onto the surface of BC, reducing the entropic penalty of BC−water interactions via MLG-mediated preorganization of bound water. The use of high molecular weight MLG represents an unexplored opportunity to generate hydrogels with tailored properties by acting as a hydration and mechanical modulator of BC-based materials, opening the possibility to generate biocompatible BC-based hydrogels with tuneable hydration and water holding properties for different applications.

## ASSOCIATED CONTENT

* Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.biomac.9b01070.

Gel thickness, TGA, PXRD, SEM, AFM, rheology, and NMR spectra and data analysis (PDF)

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**Figure 6.** Normalized water spin diffusion build-up curves of the anomeric carbon for BC (blue dots), BC−XG (red dots), BC−AX (grey dots), and BC−MLG (green dots).
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