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Published in:
Cellulose

DOI:
10.1007/s10570-019-02600-z

Published: 15/08/2019

Document Version
Publisher's PDF, also known as Version of record

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Please cite the original version:
Weißl, M., Hobisch, M. A., Johansson, L. S., Hettrich, K., Kontturi, E., Volkert, B., & Spirk, S. (2019). Cellulose carbamate derived cellulose thin films: preparation, characterization and blending with cellulose xanthate. Cellulose, 26, 7399–7410. https://doi.org/10.1007/s10570-019-02600-z

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Cellulose carbamate derived cellulose thin films: preparation, characterization and blending with cellulose xanthate

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Received: 28 March 2019 / Accepted: 26 June 2019 / Published online: 4 July 2019 © The Author(s) 2019

Abstract  Cellulose carbamate (CC) was employed as a water-soluble precursor in the manufacturing of cellulose based thin films using the spin coating technique. An intriguing observation was that during spin coating of CC from alkaline aqueous solutions, regeneration to cellulose was accomplished without the addition of any further chemicals. After rinsing, homogeneous thin films with tunable layer thickness in a range between 20 and 80 nm were obtained. Further, CC was blended with cellulose xanthate in different ratios (3:1, 1:1, 1:3) and after regeneration the properties of the resulting all-cellulose blend thin films were investigated. We could observe some slight indications of phase separation by means of atomic force microscopy. The layer thickness of the blend thin films was nearly independent of the ratio of the components, with values between 50 and 60 nm for the chosen conditions. The water uptake capability (80–90% relative to the film mass) determined by $\text{H}_2\text{O}/\text{D}_2\text{O}$ exchange in a quartz crystal microbalance was independent of the blend ratio.

Keywords  Cellulose carbamate · Cellulose xanthate · All cellulose blend films · Cellulose thin film · Cellulose swelling

Introduction

Despite being rather old, the Viscose process still is the most important and frequently used technology for the production of regenerated wood based fibers with annual production volumes exceeding 3.5 million tons, mainly for the textile industry (Wang et al. 2016; Hämmerle 2011). However, there are several environmental drawbacks of this technology. For instance, the necessity to use CS$_2$ to form the cellulose precursor material (cellulose xanthate, CX), as well as the development of volatile sulfur containing compounds (e.g. H$_2$S, COS) during the regeneration procedure requires complex recovery technologies, which manifest into higher prices of the final fiber products. These disadvantages have led to the development of new fiber spinning technologies such as the Lyocell...
and the Ioncell processes that both avoid the use of cellulose derivatives by direct dissolution into NMMO/water and ionic liquids (Rosenau et al. 2001; Sixta et al. 2015). However, costs are still higher for fibers derived from these processes since also here the recovery of the solvents requires substantial energy input. Another technology that has raised attention in recent years is the Carbacell process. The Carbacell process relies on cellulose carbamate (CC), which is easily obtained by reacting cellulose with urea. CC is soluble in cold alkali and can be subjected to wet spinning processes similar to those in viscose plants (Fink et al. 2014). As regeneration bath, a diluted acid or a sodium carbonate solution can be used (Wendler et al. 2012). The similarities in the solvent, the spinning process and the regeneration bath also allows for fiber spinning of solutions containing CX and CC, with the goal to reduce the usage of CS₂ in the fiber manufacturing process.

In recent literature, the processing of regenerate cellulose fibers from spinning dopes, blended with CC and CX was described and mechanical properties of the obtained fibers were studied. But fundamental questions concerning the miscibility of the two cellulose derivatives or possibly occurring phase separation during various process steps, could not be answered in these macroscopic studies. In addition, development of dense core shell structured fibers during wet spinning and subsequent fiber stretching prohibits a detailed study of the cellulose interfaces in those blend materials (Protz et al. 2018).

Thus, the processing of cellulose thin films based on CC, CX and blends of these solutions delivers opportunities to study the influence of the CC on the properties of CX based materials. Cellulose thin films provide a confined two-dimensional network with a well-defined surface morphology and chemistry (Kontturi et al. 2006). Literature gives many examples, where thin films have been employed to study the influence of certain conditions on the behavior of cellulose and its derivatives, or to observe the interaction of cellulose with biopolymers or proteins (Ehmann et al. 2015; Mohan et al. 2017; Niigelhell et al. 2016; Niinivaara et al. 2016). In a previous work, some of us published a method for the processing of cellulose thin films based on the spin coating and following HCl vapor exposure of CX films (Weiβl et al. 2018). This made us interested in the fundamental suitability of CC solutions to form thin film structures. As the processing of the CC can be assigned as green and sustainable, the solubility in diluted sodium hydroxide allows to avoid organic solvents and the regeneration in acidic media should also be possible in an acidic vapor phase. Further, CX and CC are soluble in the same solvent, offering opportunities to blend the solutions and spin coat cellulose thin films from two different sources, thereby creating all-cellulose blend thin films (Protz et al. 2018). The blending of these two promising and well known cellulose derivatives does not only allow to study cellulose–cellulose interfaces in a well-defined environment, but can be used to explore the influence of CC on CX during the formation and regeneration to cellulose. Through the confined two-dimensional structure, this blend film processing could show interactions and effects which are not visible in the wet spinning of blended solutions from CC and CX.

Experimental

Materials

Cellulose carbamate (CC) with a DS of 0.3 (N content 2.51% according to elemental analysis), a DP of 450 and a dry mass of 92% was provided from Fraunhofer Institute for Applied Polymer Research (Potsdam-Golm, Germany) and used without any further treatment. Cellulose xanthate (CX) stock solution with a cellulose content of 10 wt%, a DS of 0.5 and a DP of 550 was provided from Lenzing AG (Lenzing, Austria). Sodium hydroxide, hydrochloric acid (37%), sulfuric acid (95%) and hydrogen peroxide (30%) were purchased from VWR chemicals. De-ionized (DI) water was produced in house with an Elga PURELAB Prima (Bucks, United Kingdom) water treatment system. Chromafil Filters Xtra PVDF-45/25 0.45 µm were used as obtained. Single side polished silicon wafers from Siegert Wafers (Aachen, Germany, wafer thickness: 675 ± 25 µm, 1 cm × 2 cm) and gold coated glass slides from BioNavis (Tampere, Finland, gold layer thickness: 50 nm, 1 cm × 2 cm) were used as substrates for the cellulose thin films. QCMD sensors (QSX301) were purchased from nanoscience Instruments (phoenix, USA). For spin coating, a Polos 150.
Cellulose carbamate film processing

For the spin coating process, CC (1.0, 1.5, 2.0, 2.5 wt%) was dissolved in 8% NaOH solution at −10 °C under vigorous stirring for 24 h. The solutions were filtered through a 0.45 μm filter and 80 μl cm−2 were transferred on the employed substrates. The slides used as substrates for the films were cleaned by dipping them into piranha acid (H2SO4:H2O2 = 3:1 (v/v)) for 30 min (10 min for gold slides) followed by intense rinsing with MilliQ water in advance. The spin coating itself was performed on a Polos Spin 150i (from SPS Europe, Ingolstadt, Germany) with a speed of 4000 rpm and an acceleration of 2500 rpm/s for 60 s as already described in previous work (Weißl et al. 2018, 2019). After spin coating, the films were stored for at least 1 h under ambient conditions before they were rinsed with 10 ml water and dried under nitrogen flow. Afterwards, the films were treated in acid vapor atmosphere (37% HCl) for a period of 20 min (Figure S1, SI) with subsequent rinsing (10 ml water) and a final drying process at 105 °C for 60 min.

CC/CX blend film processing

A 2.0 wt% CX solution was prepared by diluting the provided stock solution with 8% NaOH under rigorous shaking by hand for a period of 2 min and subsequent filtration through a 0.45 μm filter. This CX solution was mixed with a 2.0 wt% CC solution in different ratios (1:0, 3:1, 1:1, 1:3, 0:1) for a period of 10 min under sonication. This solution was deposited on a substrate and subjected to spin coating (v = 4000 rpm, a = 2500 rpm/s, t = 60 s), followed by HCl vapor exposure (20 min) with subsequent rinsing (10 ml water) and a final drying process at 105 °C for 60 min.

Optical microscopy

Light microscopy images were taken by an Olympus BX60 equipped with an Olympus E-520 camera. Prior to investigating the samples, carrier substrates were fixed on a glass layer.

Elemental analysis

Elemental analysis was carried out by a vario MICRO analyser from Elementar (Langensebold, Germany), where the samples are burned within a jet injection of oxygen. The gaseous components were purified and separated on a TPD column, before quantification with a thermal conductivity detector. Helium was used as a carrier gas.

X-ray photoelectron spectroscopy (XPS)

Surface chemistry was evaluated with XPS, using a Kratos AXIS Ultra photoelectron spectrometer with monochromated Al Kα irradiation at low power (100 W) and under neutralization. Prior to the experiment samples were pre-evacuated overnight, in order to stabilize experimental conditions. Samples were measured together with the lab-defined in situ reference of 100% cellulose (Whatman). Both wide energy resolution scans, with 1 eV step and 80 eV CAE, and high resolution scans of Cls and O1s regions with 0.1 eV step and 20 eV CAE were recorded on 2–3 locations. Nominal analysis spot size is 400 × 800 μm. All binding energies were charge-corrected using the cellulose Cls main component of C–O at 286.7 eV, after fitting the high resolution carbon data into four Gaussian components with equal half widths, using CasaXPS software (Johansson and Campbell 2004).

Profilometry

The layer thickness was determined with a Bruker DekTak XT surface profiler. The scan length was set to 1000 μm over the duration of 3 s with the hills and valleys scanning profile. The diamond stylus had a radius of 12.5 μm and the employed force was 3 mg. The sample was scraped before measuring and the obtained profile was then used to determine the thickness. Each layer thickness has been determined by averaging 9 measurements on three different slides.

Attenuated total reflection infrared spectroscopy

The infrared spectra were recorded with an ALPHA FT-IR spectrometer (Bruker; Billerica, MA, U.S.A.). For the measurement, an attenuated total reflection (ATR) attachment was used with 48 scans at a resolution of 4 cm−1 and a scan range between 4000 and 400 cm−1. The samples were prepared on Au-coated glass slides (SPR102-AU). The data were analyzed with OPUS 6.0 software.
Atomic force microscopy

The surface was investigated with a Tosca™ 400 atomic force microscope (AFM, Anton Paar, Austria) in tapping mode using Al-coated cantilevers (ARROW-NCR, NanoWorld AG, Switzerland) with a resonance frequency of 285 kHz and a force constant of 42 N m\(^{-1}\). All measurements were acquired at room temperature under ambient conditions. All calculations and image processing was performed with Tosca™ analysis (V7.4.8341, Anton Paar, Austria).

Contact angle and surface free energy determination

For the calculation of the surface free energy (SFE) Milli-Q water (\(> 18 \text{ MΩ cm}^{-1}\)) and diiodomethane were employed as test liquids. The drop shape analysis was done in the sessile drop modus at 25 °C with a DSA100 system (Krüss GmbH, Hamburg, Germany) equipped with a T1E CCD video camera (25 fps). The dispense rate of the three \(\mu\text{L}\) droplets was adjusted to 256 \(\mu\text{L/min}\) and the time before the image was captured was set to 1 s. Each sample was measured at least six times on at least two samples. The contact angle (CA) calculations (software: DSA1 v 1.90) were performed with Young–Laplace equation and the surface free energy calculation according to the Owens–Wendt–Rabel & Kaelble method (Owens and Wendt 1969).

Quartz crystal microbalance with dissipation monitoring

A QCM-D from Q-Sense (Gothenburg, Sweden) with simultaneous detection of the resonance frequency \(\Delta f\) and energy dissipation \(\Delta D\) of an oscillating piezoelectric crystal was employed. Dissipation describes the frictional losses which cause a damping of the oscillation in correlation to the viscoelastic properties to the materials absorbed on the crystals. Rigid layers are assumed to fully couple to the oscillation of the crystal, where the change in frequency \(\Delta f\) is given by the Sauerbrey equation

\[
\Delta m = C \frac{\Delta f}{n}
\]

\(C\) is the Sauerbrey constant (\(-17.7 \text{ ng Hz}^{-1} \text{ cm}^{-2}\) for a 5 MHz crystal), \(\Delta m\) is the changing mass of the crystal due to the adsorbed layer and \(n\) is the overtone number (Sauerbrey 1959). The baseline resonance frequencies for the crystals were determined before depositing the cellulose layers and subsequent determination of the water content in the layers was done with a H\(_2\)O/D\(_2\)O exchange experiment, as described by Kittle et al. (2011) and Mohan et al. (2012).

\textbf{Determination of the water equilibrium content}

By exploiting differences in viscosity and density of H\(_2\)O/D\(_2\)O, the amount of water uptake by the accessible parts of the thin films can be determined. Based on the Kanazawa (Keiji and Gordon 1985) equation it follows:

\[
\frac{\Delta f_{\text{H}_2\text{O}}}{n} = \frac{\Delta f_{\text{film}}}{n} - \frac{\Delta f_{\text{bare}}}{n} \frac{\rho_{\text{D}_2\text{O}}}{\rho_{\text{H}_2\text{O}}} - 1
\]

(2)

where \(\Delta f\) (film) and \(\Delta f\) (bare) describe the frequency shifts caused by an exchange of H\(_2\)O by D\(_2\)O on a bare substrate and on substrates coated with a blend film respectively. The density \(\rho\) is 0.9982 g/cm\(^3\) for H\(_2\)O and 1.1050 g/cm\(^3\) for D\(_2\)O, \(n\) is the overtone number and \(\Delta f\) (H\(_2\)O) is the frequency shift, induced by the water absorbed in the cellulose films. With the Sauerbrey equation, \(\Delta f\) (H\(_2\)O) can directly be transferred in a total water content.

\[
\Gamma_{\text{water}} = C \frac{\Delta f_{\text{H}_2\text{O}}}{n}
\]

(3)

After recording the resonance frequencies before and after coating, the films were equilibrated with MilliQ water (flow rate: 0.1 ml/min) for a period of 30 min. Afterwards, the water was exchanged by D\(_2\)O and the cells were flushed with D\(_2\)O until no further change in frequency was observed (i.e. for 15 min). The last step was to change back to water again until equilibration was reached (15 min). The resulting frequency shifts before and after injection of D\(_2\)O were used to determine \(\Gamma_{\text{water}}\) according to (3). The H\(_2\)O/D\(_2\)O exchange was also done with uncoated crystals that served as reference. For each experiment, at least three parallel measurements have been performed. The determined mass of the exchanged
water was then correlated to the film mass to normalize the results to the thickness of the films.

Results and discussion

Formation and characterization of pure CC thin films

The spin coating of CC solutions with concentrations between 1.0 and 2.5 wt% yielded films with full and regular coverage of the substrates as determined by optical microscopy. Below a concentration of 1.0 wt%, the coverage of the slide was incomplete and inhomogeneous; above 2.5 wt% the solution could not be filtered anymore and the resulting films showed a high surface roughness and irregularities caused by aggregates (data not shown).

After spin coating was finished, needle-shaped salt crystals formed a top layer and covered the whole substrate within seconds (Figure S2, SI). After rinsing the films with de-ionized water, the crystalline top layer was washed away and a smooth cellulose layer with a defined morphology remained on the substrate. The following acid vapor deposition as well as the drying process did not cause a visible change in the surface morphology of the films. The layer thickness for films prepared from different CC solutions is depicted in Fig. 1a. It is noticeable that the evolution of the layer thickness during the processing is similar for the different concentrations. Prior to rinsing the films with water, a thickness of 200 to 400 nm was determined. After rinsing and simultaneous removal of the salt crystallites with water, the thickness decreased significantly to 22, 42, 56 and 75 nm for concentrations of 1.0, 1.5, 2.0 and 2.5 wt%, respectively. Subsequent HCl vapor treatment or the drying step at 105 °C did not show strong impact on the film thickness.

Figure 1b summarizes the IR spectra of the thin films during processing and the corresponding reference materials. The CC used as starting material features characteristic broad bands in a region of 3600–3100 cm\(^{-1}\) (OH vibration) and around 2900 cm\(^{-1}\) (CH stretching) followed by a series of small bands between 1430 to 1150 cm\(^{-1}\) (C–O–H bending at 1430 cm\(^{-1}\), C–H deformation at 1372 cm\(^{-1}\), OH in plane deformation at 1330 and at 1200 cm\(^{-1}\)) and strong overlapping bands between 1160 and 890 cm\(^{-1}\) (asym. C–O–C vibration at 1155 cm\(^{-1}\), sym. C–O vibration at 1060 cm\(^{-1}\), C–O stretching at 1035 cm\(^{-1}\) and C–O–C valence vibration at 899 cm\(^{-1}\)) (Široký et al. 2010). In addition to the cellulose induced bands, two significant bands at 1620 and 1700 cm\(^{-1}\) indicate the presence of carbamates (Nada et al. 2000; Xiong et al. 2017).

After spin coating, the spectrum of the precipitated CC showed particular differences in comparison to the CC source material. Strong overlapping bands at 1480, 1420 and 880 cm\(^{-1}\) are characteristic for sodium carbonate and clearly dominate the spectrum, whereas bands caused by cellulose and sodium hydroxide are

![Fig. 1](image-url)
much less pronounced (Neufeld et al. 2002). The intensive sodium carbonate band and the described growth of a crystalline structures after spin coating suggest the expected formation of a sodium carbonate top layer (Weißl et al. 2018). After rinsing the substrates, a spectrum matching all the cellulose bands as described by Široky et al. was obtained (Široký et al. 2010). Further treatment in an HCl vapor phase or drying at 105 °C did not affect the cellulose structure according to data from infrared spectroscopy. The detailed spectra are shown in Figure S3 to S6, SI. The feature at 1647 to 1650 cm\(^{-1}\) can be assigned to adsorbed water on cellulose as described in literature. This band is present in all the spectra even in those that do not contain any cellulose carbamate (i.e. cellulose from CX).

To further confirm that regeneration of the CC already occurs during spin coating, XPS was employed. The XPS survey spectra did not reveal the presence of nitrogen and sodium in the films after rinsing (Figure S7, SI). Figure 2 summarizes carbon and oxygen high resolution spectra of spin coated CC after rinsing the films with water and after additional HCl vapor treatment.

Both, the C1s and the O1s environments match the bonding energies previously described for cellulose materials. The C1s peaks showed three single environments namely C–C (H) at 284.5, C–OH at 286.5 and C–O–C at 288 eV, respectively, whereas the O1s peak features only one environment at 533.5 eV. A shift in the described binding energies or new bands suggesting changes in the cellulose structure, could not be detected within all post spin coating treatments (Shchukarev et al. 2002).

The conversion to cellulose by spin coating is an intriguing observation. To the best of our knowledge,
there are not any reactions that are induced by spin coating in literature. During spin coating, the basicity of the system is steadily increasing due to water evaporation (i.e. higher hydroxide concentration). It is known that primary carbamates in general may be subject to decomposition under extreme alkaline conditions via an addition–elimination mechanism (Scheme 1). In such reactions, the nucleophile (OH–) attacks the carbon at the carbamate group under elimination of NH₃ (Moidoveanu and David 2002). These reactions are very often kinetically driven, i.e. steric hindrance plays a role and determines under which conditions these reactions actually take place. In case of cellulose, the carbamate group is sterically protected and anchored on a polymer, thereby slowing down the reaction (Fu et al. 2014; Nada et al. 2000). There are reports to regenerate cellulose carbamate fibers in diluted hot alkaline solutions, corroborating the kinetic nature of the decomposition reaction (Klemm et al. 2005; Kunze and Fink 2005). During spin coating of CC solutions, the conversion to cellulose probably involves the diffusion of the labile carbamic acid to the cellulose-air interface, where it is decomposed to form sodium carbonate under elimination of ammonia. This would also explain the formation of the crystalline sodium carbonate layer shortly after spin coating has been accomplished and is supported by the ATR IR spectra that show all the typical bands associated with sodium carbonate.

AFM was employed to gain further information about the surface morphology and structure of the CC films. Compared to cellulose thin films from different sources like trimethylsilyl cellulose (TMSC) (Niegelhell et al. 2016; Schaub et al. 1993) or cellulose xanthate (CX) (Weiβl et al. 2018), the surface of CC thin films features more aggregates (10–13 nm) concomitant with slightly higher roughness. The AFM images (Fig. 3) suggest that concentrations as high as 1.0 wt% are suitable to completely cover the substrates. The morphology of films was regular and smooth with a RMS roughness between 5 nm (lowest concentration) and 12 nm (highest concentration). Compared to the increase in the root mean square (RMS) roughness with increasing CC concentration, no trend of increasing or decreasing surface roughness or any other change in surface morphology could be identified within the single processing steps (Figure S8, SI).

CC films spin coated from a 1.5 wt% solution displayed static water contact angles between 12° and 27°, depending on the post treatment. At higher CC concentrations used for spin coating, the development of the contact angle followed the same direction but the range between the different treatments was much smaller (from 19° to 26° for 2.0 wt% CC, and from 29° to 32° for 2.5 wt%).

The SFE of films spin coated from different concentrations was determined from the H₂O and diiodomethane contact angles following the Owens, Wendt, Reynolds and Kaeble method (Owens and Wendt 1969). The obtained SFE (66–77 mN/m) are in the range reported for cellulose thin films (Fig. 4). The slight variations in the SFE values are caused by the changes in the surface roughness of the films since the chemistry of the films is the same. Such deviations in SFEs were also observed for cellulose thin films.
derived from other precursors (Mohan et al. 2011; Weißl et al. 2018, 2019).

Formation and characterization of CC/CX blend films

CC and CX are both soluble in 8% NaOH solutions which was exploited to blend solutions in 1:3, 1:1 and 3:1 ratios (v/v). After spin coating the blend solutions, and subsequent regeneration to all-cellulose blend thin films, their properties were determined and correlated to cellulose thin films derived from CX and CC, respectively. CX does not regenerate during spin coating as demonstrated in previous reports; therefore an HCl vapor treatment was performed to ensure conversion of the CX domains to cellulose (Weißl et al. 2018, 2019). The domains consisting of CC were not affected by this procedure as discussed above (Figs. 1, 2, 3).

Taking a closer look into the ATR-IR spectra of the blend films and reference materials (Fig. 5a), the region between 850 and 950 cm$^{-1}$ attracts attention. For the pure CX, the bands can be assigned to vibrations related to sodium sulfide (1420, 920 cm$^{-1}$) and sodium trithiocarbonate (1670, 1427, 925 and 885 cm$^{-1}$). After HCl vapor exposure and rinsing with water, the blend films and the source materials were both fully regenerated and infrared spectroscopy only revealed bands related to cellulose (Figure S9, SI). The IR spectra in combination with XPS of the pure CC and the CX shown in a previous publication confirm the full regeneration of the different cellulose derivatives to yield all-cellulose blend films (Weißl et al. 2019).
The determination of the layer thickness was executed by stylus profilometry. The reference materials had a thickness of 57 and 59 nm for CC and CX, respectively. The thickness of the blend films were all in the range of 52 nm after regeneration and rinsing, regardless of the blend ratio (Figure S10, SI).

Determination of the static \( \text{H}_2\text{O} \) contact angles is documented in Fig. 5b. The pure CC film had a contact angle of \( 24^\circ \) after HCl vapor exposure and with increasing CX content in the blend films, the contact angle was continuously increasing, until a value of \( 37^\circ \) — similar to the pure CX based film — was reached. The increase in the contact angle is unlikely caused by any chemical difference in the films but connected to a change in the surface morphology for the CC and the CX based layers. If the Wenzel equation is used for correcting the contact angles for the roughness of the films, differences between the films are rather small (33°–38° for water; Figure S11a, SI). The surface free energy followed the linear behavior, shown in the contact angle measurements, ranging from 72 mN/m for pure CC to 64 mN/m for pure CX based films (Figure S11b, SI).

AFM is employed to check the miscibility of CC and CX on a microscale. The evaluation of the surface morphology is summarized in Fig. 6 and indeed, slight
differences in surface structure and in the roughness were observable. The detailed mechanism on the formation of the structures visible in the phase images remained unclear.

The CC based films showed large bulky substructures, causing pronounced hills and valleys. These structures were further reflected in the RMS roughness of the thin films, where pure CC films had $R_q$ values of around 10 nm. With increasing CX content in solution for preparing the blend films, the substructures became smaller and the surface showed a more regular shape. The RMS roughness decreased to 3.5 nm for the blend films prepared from the 1:3 CC:CX ratio. The cellulose film prepared from pure CX had the smallest RMS roughness among all investigated samples.

Finally, the water uptake and the swelling behavior of the blend films were investigated by a H$_2$O/D$_2$O exchange monitored in a QCM-D device (Kittle et al. 2011; Mohan et al. 2012). In such exchange experiments, the different densities of H$_2$O and D$_2$O are exploited to determine $I_{\text{water}}$ according to Eqs. (2) and (3). Since thicker films incorporate more water than thinner ones, the results need to be normalized to the film thickness or mass if one aims at comparing water contents of films with varying thicknesses.

Figure 7a shows the frequency shifts of a bare substrate and the cellulose covered substrates during the H$_2$O/D$_2$O exchange recorded by QCM-D. It can be clearly seen that the frequency after the D$_2$O injection decreased and reached the initial frequency after exchanging back to H$_2$O, i.e. there is not any hysteresis.

The black dashed line represents the bare substrate and shows a $\Delta f_3$ shift of ca. $-71$ Hz between H$_2$O and D$_2$O. If a cellulose layer was present on the substrates, the $\Delta f_3$ were significantly higher and in the range of $-115$ up to $-120$ Hz, where the blend films (1:3, 1:1; 3:1) and the pure CC based films all had comparable shifts. Only the pure CX based showed a larger frequency shift (ca. $-130$ Hz), which correlated to the higher layer thickness.

According to the frequency shifts, the total amount of water $I_{\text{water}}$ was calculated and yielded $6.3 \pm 0.2 \mu\text{g/cm}^2$ for the CC based cellulose films (Table 1). The blend films systems exchanged $6.7 \pm 0.2$ (3:1), $6.7 \pm 0.1$ (1:1) and $6.7 \pm 0.2$ $\mu$g water/cm$^2$ (1:3). With $7.4 \pm 0.1$ $\mu$g/cm$^2$ the total water uptake was higher in the CX based regenerated cellulose film. These values scaled with the mass of the cellulose films, therefore the film thickness needs to be taken into account for determining the degree of water uptake. The thickness was 50 nm for CC based layers, around 52 nm for the blend films and 57 nm for the CX based films. With the layer thickness and the density of cellulose (1.5 g/cm$^3$) (Holbery and Houston 2006), the dry mass of the cellulose films was determined. The dry mass was $7.5 \pm 0.2 \mu$g/cm$^2$ for the CC based cellulose films, $7.3 \pm 0.2$ (3:1),...
7.7 ± 0.1 (1:1) and 7.9 ± 0.1 μg/cm² (1:3) for the blend films and 8.7 ± 0.2 μg/cm² for the CX based films. The swelling behavior is presented in Fig. 7b and it can be concluded that there is no trend observable within the thin films derived from different cellulose sources. Only the 3:1 blend (CC:CX) featured a slightly higher water uptake (92%), all the other films featured a water uptake capacity between 84 and 87%.

**Conclusion**

Here, we introduced a new system for manufacturing cellulose thin films based on ecofriendly CC. Since CC is water soluble, the use of organic solvents is omitted compared to the other often employed cellulose derivative, TMSC. In addition, CC can be synthesized in large scale via environmentally friendly procedures. The regeneration process itself does not require any additional treatment but is induced by increasing the NaOH concentration during the spin-coating via evaporation of the water, as confirmed by IR and XPS spectroscopy. Such an in situ reaction in a polymer film during spin coating has no precedent in literature. Exposure to acid vapors or elevated temperature does not cause changes in the film morphology or chemistry of the films. Further, the film thickness can be easily adjusted within a range of 20 to 80 nm by varying the CC concentration. Only the increased surface roughness, caused by the high sodium hydroxide content in the solvent, could be a drawback for the processing of thin and well-defined cellulose layers based on CC, depending on their purpose.

Blending of CX with CC is a promising method to partly substitute the frequently employed CX as source for regenerated cellulose products. Further, these all-cellulose blend films deliver an interesting possibility to investigate cellulose interfaces. Within this study, it could be shown that the blending and the regeneration of the cellulose derivatives do not lead to extensive phase separation regardless of the mixing ratio.

**Acknowledgments** Open access funding provided by Graz University of Technology. Lenzing AG and Fraunhofer Institute for Applied Polymer Research IAP are gratefully thanked for providing the cellulose xanthate stock solution and cellulose carbamate powder. Professor Gregor Trimmel and the PCCL are thanked for experimental support. EK is grateful for the support by the FinnCERES Materials Bioeconomy Ecosystem. We also thank the Dr. T. Mohan (University of Maribor) for experimental support in the QCM-D experiments.

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