α-Cellulose-Based Films: Effect of Sodium Lignosulfonate (SLS) Incorporation on Physicochemical and Antibacterial Performance

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Abstract

A homogeneous $\alpha$-cellulose film was prepared by regeneration method from ZnCl$_2$/CaCl$_2$/cellulose mixed system and was further combined with sodium lignosulfonate (SLS) by crosslinking through interaction hydrogen bonds and “bridge linkages”. The physicochemical and antibacterial performance of films were all investigated and results showed that modified films exhibited stronger tensile strength, higher thermal stability, lower hydrophilic effect, better UV shielding as compared with those of pure cellulose film, and especially, better antibacterial ability derived from the presence of phenolic and sulfonate groups in SLS. This study proposed a simple and sustainable method for fabricating a multifunctional and environmentally friendly composite film by using two main lignocellulose resources as raw materials.

Highlights

- A novel lignin-containing cellulose film was fabricated via crosslinking.
- Superior tensile strength was obtained due to dense crosslinking structure.
- Excellent antibacterial effect was derived from inherent structure of SLS.

1. Introduction

Cellulose, as one of the three main ingredients in lignocellulosic biomass, is a famous biopolymer and has great potential for preparing film-based materials (Khalili et al., 2018; Koli et al., 2018; Zhang et al., 2017). The properties of bio-compatibility, flexibility, chemical modification, and universal applicability make cellulose-based films promising to be utilized in fields of wastewater treating, wound healing, packaging, and bacteriostasis (Hu & Wang, 2016; Kong et al., 2014; Lv et al., 2018; Puspasari et al., 2018; Wang et al., 2019; Wen et al., 2019).

Recently, the regenerated cellulose film derived from regenerated method is a promising alternative to polymeric one (Puspasari et al., 2018). Many works have focused on the investigation of regenerated method due to its excellent bio-compatibility, and nontoxicity (Lee & Jeong, 2015; Pang et al., 2015). For instance, Weng et al. prepared the regenerated cellulose film through phase inversion, where N-methylmorpholine-N-oxide (NMMO) was used as a solvent for dissolving cellulose, and the obtained film exhibited high rejection rate (99.7 %) for Congo red (Weng et al., 2017). Nevertheless, the pure regenerated cellulose film is hydrophilic, poor thermal stability, bacteria prevention, and mechanical properties (Mao et al., 2006; Ruan et al., 2004). Chemical modification or blended with other polymers is effective pathway for addressing this issue through introducing different functional groups and modifying the inherent structure.

Sodium lignosulfonate (SLS) is a by-product produced via the production of wood pulp by using sulfite pulping processes. Approximately 50 million tons of SLS can be obtained every year, where 95 % of this resource is consumed by direct combustion for low-efficiency electricity supply (Kim & Um, 2020; Xu et al., 2020), therefore it is meaningful to achieve high efficiency of SLS. Fortunately, abundant presence of
active groups, such as hydroxyl (O-H) and sulfonate (R-SO₃⁻Na⁺), and various linkages, including C-O and C-C, make SLS reactive for oxidation, reduction, and polymerization reactions. In particular, as a promising alternative of polyol, various types of lignin can crosslink with other aldehydes, polyols, acid anhydrides, and amines for preparing bio-based adhesive (Ang et al., 2019; Chen et al., 2020; Vázquez et al., 1995; Zhang et al., 2013). Recently, lignin-containing cellulose mixed films have obtained broad attention due to their wide distribution, low production cost, and environmental friendliness (Rojo et al., 2015). With the incorporation of lignin, the related properties (e.g., mechanical performance, thermal stability, UV transmission, and hydrophobic effect) of mixed films can be improved significantly (Nair & Yan, 2015; Wen et al., 2019). Therefore, combining lignin and cellulose via crosslinking method is a promising and innovative strategy for preparing lignin-containing cellulose mixed films.

Recently, an inorganic salt system (zinc chloride/calcium chloride) was researched for preparing cellulose films (Xu et al., 2016). Therefore, we prepared regenerated films (including pure and mixed films with different content of SLS) through inorganic co-system (ZnCl₂/CaCl₂), and modified these obtained films through crosslinking with assistance of succinic anhydride and triethanolamine. The result showed that the modified mixed film (cellulose/SLS-M) with assistance of succinic anhydride and triethanolamine acting as crosslinking agents exhibited excellent properties (such as thermal stability, hydrophobic effect, mechanical performance, and UV transmission) as compared with those of original one, and the improved performance of bacteriostasis was mainly derived from the addition of SLS.

2. Materials And Methods

2.1. Materials

In this study, α-cellulose with particle size of 250 μm used was purchased from Aladdin (Shanghai, China). Sodium lignosulfonate (SLS, 96 %) was purchased from TCI (Shanghai, China), and other chemical reagents, such as zinc chloride (ZnCl₂, 99 %), calcium chloride (CaCl₂, ≥ 99.9 %), absolute ethanol (≥ 99.5 %), succinic anhydride (98 %), triethanolamine (98 %), anhydrous calcium sulfate (CaSO₄, 97 %), sodium carbonate anhydrous (Na₂CO₃, ≥ 99.5 %) were all purchased from Macklin (Shanghai, China). All chemical reagents were utilized without any pretreatment, and deionized water was produced from our lab.

2.2. Preparation of mixed cellulose films

These cellulose films (CFs, including pure and mixed cellulose films which contained different amounts of SLS) were prepared via a regeneration method reported in a previously reported literature (Hou et al., 2019). Typically, 0.45 g α-cellulose was added into 1.05 g deionized water to obtain a cellulose suspension liquid. Afterwards, continuous stir was performed in a beaker (10 mL) for 30 min to achieve uniform dispersion of cellulose, which was recorded as solution 1. Then, 9.87 g ZnCl₂ and 0.30 g CaCl₂ were mixed into 3.63 g deionized water. After 30 min stir, different amounts of SLS (10, 20, 30, and 40 wt%, based on the input of α-cellulose) were added into this homodisperse metal precursor solution, and
were stirred at 75 °C for 30 min together, which was named as solution 2. After that, solution 2 was dropwise added into solution 1 with intense stir and was further stirred at room temperature for 12 h to make these two solutions mix well. Finally, the obtained solution was placed under vacuum atmosphere to remove air bubbles and was casted on a glass plate to prepare the film with thickness of 450 µm, and then it was immersed in 450 mL absolute ethanol for 1 h to separate the film from glass plate, which was immediately fixed with two splints and dried at room temperature overnight. The prepared film was tailored with size of 2 cm × 2 cm for the next characterization and was denoted as SLS \(_x\)-CF, where \(x\) means the mixed addition of SLS (wt%). For instance, SLS\(_0\)-CF represents the pure CF without extra addition of SLS, while SLS\(_{10}\)-CF means the mixed CF with the addition of 10 wt% SLS.

### 2.3. Modification process

Due to the presence of hydroxyl groups (O-H) in structures of cellulose and lignin, they have the potential to be utilized as polyols for the achievement of crosslinking (Feng et al., 2021; Lamaming et al., 2020; Yin et al., 2020). Typically, it is difficult to realize mutual crosslinking between two macromolecules through chemical pathways due to their complex structures and unclear reaction mechanism. Therefore, the introduction of “bridge monomers” (succinic anhydride and triethanolamine) is beneficial for improving the interconnection between these two macromolecules. Generally, the polymerization reaction occurs between carboxylic acids and alcohols at a certain temperature to form polymeric esters. In theory, it is feasible to combine cellulose and lignin through this mechanism mentioned above. In this section, succinic anhydride and triethanolamine were utilized as “bridge monomers” (i.e., crosslinking agents) to improve the interconnection between \(\alpha\)-cellulose and SLS to form the crosslinking structure, and the specific modification steps are as follow.

First, succinic anhydride (0.18 g), triethanolamine (0.45 g), and different volumes of absolute ethanol (100, 200, and 300 mL) were blended uniformly to prepare organic precursor solutions with different concentrations. Afterwards, the pure CF (i.e., SLS\(_0\)-CF) was immersed into different organic precursor solutions for different time (1, 2, and 3 days). Finally, the obtained film after immersion was cured at different temperatures (50, 60, and 70 °C) for 6 h to make the modification complete. Finally, the film after modification was washed by ethanol for three times to remove unreacted molecules, and dried at curing temperatures (50, 60, and 70 °C) for 12 h, which was named as CF-M. The performance of obtained films was preliminarily evaluated via their mechanical properties to optimize the modification condition, and results were presented in the next discussion. Other blending films, which contained different amounts of SLS (10, 20, 30, and 40 wt%), were all modified by the optimized condition (300 mL ethanol, 70 °C curing temperature, and 1 day immersing time), which were assigned as SLS\(_x\)-CF-M. During the curing at mild conditions (below 100 °C), succinic anhydride was hydrolyzed as succinic acid, which can be polymerized with polyols (such as triethanolamine, and hydroxyl groups in structures of cellulose and lignin) to achieve the interconnection of several molecules, and the illustration of modification process is presented in Scheme 1. From the structure displayed in Scheme 1, two internal bonding (interaction
hydrogen bonding and crosslinking derived from “bridge monomers”) can be obtained to fabricate the crosslinking structure of modified blending films.

2.4. Film characterization

The film thickness was measured by an ID-C112XBS micrometer (Mitutoyo Corp., Tokyo, Japan) and presented as average value of five points. FTIR (Fourier transform infrared spectrometer) was performed on a VERTEX 70 spectrometer. The range of measured wavenumbers was between 500–4000 cm⁻¹, and 32 scans per spectrum were collected with a resolution of 4 wavenumbers. The surface morphology was recorded by S-4800 cold field emission SEM (scanning electron microscope). The thermal stability of prepared films was carried out on a TA Instruments TGA (thermo gravimetric analyzer) Q500 (TA Instruments, USA) with a fixed heating rate of 10 °C/min from room temperature to 600 °C with nitrogen flow of 40 mL/min. Tensile test was conducted on the film with a strain rate of 30 mm/min at 25 °C via an auto tensile tester (SANS CMT4000). The film specimens contained width and length of 10 and 35 mm, respectively. The water contact angle was determined by Kruss DSA100. Light transmission was carried out through the film (2 cm × 2 cm) on a Lambda 950 UV-vis spectrometer in the wavelength range of 200–600 cm⁻¹.

The water absorption was measured via Eq. (1):

\[
\text{Water absorption} = \frac{W_t - W_0}{W_0} \times 100\%
\]

where \( W_t \) was the mass of film measured at different immersion time at room temperature after excess surface water was removed.

Water vapor permeance (WVP) was tested by the standard ASTM method E96. Typical bottles with diameter of 2.7 cm and volume of 20 mL were selected to determine the WVP of films. First, the film was cut into a circle with a diameter of 2.8 cm (slight larger than bottle used). After that, a determined amount of anhydrous CaSO₄ (RH = 0 %) was added into the bottle until two-thirds volume (approximately 3 g), which was covered by different films. Each bottle was placed into a desiccator (a 100 mL beaker) containing saturated Na₂CO₃ solution at the bottom. Excess amount of Na₂CO₃ was added to ensure the saturability of Na₂CO₃ solution, providing a constant RH of 92 % at 25 °C. The bottle was weighted every 12 h to record the change of mass, which was recorded as a function of time, and the slope of line (mass change vs. time) was obtained by linear regression. The WVP was determined as the slope divided by the transfer area (g/day m²).

Antimicrobial activity of films was evaluated by measuring the diameter (cm) of inhibition zone. The bacteria Escherichia coli (E. coli) was selected as researching object, which was grown in nutrient broth
liquid medium at 37 °C for 12 h and further diluted to create a bacteria suspension with the concentration of $1 \times 10^6$ CFU (colony forming unit m/L). Diluted bacteria suspension (0.15 mL) of *E. coli* was combined with nutrient agar. Control sample (CF) and modified mixed samples (SLS$_x$-CF-M) were employed to evaluate effects of the presence of SLS on film antimicrobial activity. Films were cut into circle discs at 6 mm in diameter, and further placed on central location of solid agar plates inoculated with testing bacteria *E. coli*, where each solid agar plate was incubated at 37 °C for 24 h before experiments for the purpose of cleaning zones around the testing film. After experiments, the observed clear zones were places where bacteria growth was inhibited.

All measuring data (e.g., water absorption, WVP, and the diameter of inhibition zone) was recorded in triplicates for each group.

### 3. Results And Discussion

#### 3.1. FTIR

FTIR spectroscopy is a common characterization for recording the structural change of testing materials, which is presented in Fig. 1. As shown in Fig. 1a, the pure CF exhibits a strong and broad absorbance at around 3300–3350 cm$^{-1}$, which is assigned to O-H stretching derived from the structure of cellulose. The peak located at 2902 cm$^{-1}$ is attributed to C-H stretching. Other bands for CF mainly appear at 1310, 1102, 102, and 895 cm$^{-1}$, which are related to the stretching of glucopyranose ring in cellulose (Hu & Wang, 2016). After modification, the peak at 1456 cm$^{-1}$ is due to the presence of C-N originating from triethanolamine, while peaks at 1725 and 1570 cm$^{-1}$ are characteristic of C=O stretching vibration and carboxylate groups derived from succinic anhydride (Cao et al., 2011; Liu et al., 2009; Sehaqui et al., 2017). Additionally, the peak of CF-M at around 1047 cm$^{-1}$ becomes weaker as compared to that of pure CF, indicating the re-formation of hydrogen bonds caused by the addition of crosslinking agents (Cai et al., 2018). Furthermore, the O-H stretching absorbance of modified film located around 3300–3350 cm$^{-1}$ becomes stronger and shifts to higher wavenumber region, demonstrating the strong intermolecular hydrogen linkage between “bridge monomers” and cellulose, which is a reason for the change of tensile strength (Hu et al., 2016). After being blended with SLS, several changes of functional groups (such as the absorbance of O-H, C-H, and C-O groups) are observed in Fig. 1b. Moreover, the characteristic peak of SLS (i.e., sulfonate group) is located at 615 cm$^{-1}$, which increases with the increase of SLS content (Liu et al., 2018).

#### 3.2. SEM

The surface morphology of pure CF, CF-M, and SLS$_x$-CF-M was observed by SEM and shown in Fig. 2. Clearly, the surface of pure CF is homogeneous, while the obvious variation on surface morphology appears after modification to make the film heterogeneous, which is due to the co-existence of different crystalline and amorphous regions derived from different components. With the addition of SLS, the film becomes more heterogeneous and phase separation occurs as the content of SLS increases from 10 to
40 wt%, which suggests that CF and SLS become more incompatible. This observation indicates that excess addition of SLS is not beneficial for the effect derived from “bridge monomers” to make the interconnection between cellulose and lignin close, which may be the cause of variation in mechanical property.

### 3.3. TGA

Figure 3 shows thermograms of pure CF and other modified blending films. As presented in Fig. 3, the thermal degradation process is mainly divided into three stages, including moisture evaporation (stage 1, below 150 °C), rapid decomposition (stage 2, from 150–500 °C), and carbonization (stage 3, above 500 °C) stages (Lu et al., 2021). The main decomposition process occurs at stage 2, where the mass loss is mainly attributed to the thermal decomposition of “bridge linking” bonds (e.g., C-O, C = O), cellulose backbone, and lignin structure to form H₂O, CO₂, CH₄, CO, and various aromatic volatiles (de Oliveira Junior et al., 2012; Lu et al., 2020; Mazarin de Moraes et al., 2015). At the stage of carbonization, char formation (around 35 wt%) is inevitable because of the dominant proportion of carbon (C) in lignocellulosic biomass (Laouge & Merdun, 2020; Yeo et al., 2019). In addition, almost no difference is observed during the first stage, while the maximum degradation temperature increases from 304 °C for pure CF to 305 °C for SLS₁₀-CF-M, 308 °C for SLS₂₀-CF-M, 307 °C for SLS₃₀-CF-M, and 310 °C for SLS₄₀-CF-M, respectively. The increasing variation can be attributed by the crosslinking effect derived from “bridge monomers”, the interaction hydrogen bonding between cellulose and lignin, or the deterioration of heat and mass transfer in samples after the addition of lignin (Ma et al., 2015). Moreover, several previous studies had reported that the incorporation of lignin (a three-dimensional structure macromolecule) as a natural carbon resource could improve the thermal stability of raw substrate, especially for the field of flame retardant (Dai et al., 2020; Ferdosian et al., 2016; Podkoscielna et al., 2020). Furthermore, not only the improvement of thermal stability during main decomposition process can be achieved, but also the content of condensed phase obtained after pyrolysis also increases, which is caused by structural collapse of cellulose backbone and lignin aromatic units to form the thermostable char with higher crosslinking density. TGA results illustrate that the thermal stability of pure CF is improved obviously, and predictable reasons are as follows: 1) abundant polar bonds (hydroxyls) were introduced with the addition of lignin, which could interact with cellulose via interaction hydrogen bonds to form crosslinking network; 2) the introduction of “bridge monomers” was beneficial to form the interconnection between lignin and cellulose via “bridge linking” bonds (the polymerization of carboxylic acid and alcoholic hydroxyl), thus strengthening the stability of mixed system; 3) three-dimensional structure of lignin was more stable as compared with cellulose, which was harder to be decomposed at high temperatures, as more stable condensed phase (char) with higher thermal stability was generated due to the repolymerization caused by the instability of active functional groups and the regeneration of carbon-carbon bonds.

### 3.4. Mechanical properties
Table 1
Tensile strength of CF-M prepared via different conditions.

| No. | Ethanol/mL | Temperature/°C | Time/day | Tensile strength/MPa |
|-----|------------|----------------|----------|----------------------|
| 1   | a          | a              | a        | 56.6                 |
| 2   | a          | b              | b        | 46.6                 |
| 3   | a          | c              | c        | 47.9                 |
| 4   | b          | b              | a        | 56.7                 |
| 5   | b          | c              | b        | 60.8                 |
| 6   | b          | a              | c        | 56.4                 |
| 7   | c          | c              | a        | 64.6                 |
| 8   | c          | a              | b        | 50.7                 |
| 9   | c          | b              | c        | 55.0                 |

*a Ethanol: 100 mL; Immersion time: 1 day; Curing temperature: 50 °C.*

*b Ethanol: 200 mL; Immersion time: 2 days; Curing temperature: 60 °C.*

*c Ethanol: 300 mL; Immersion time: 3 days; Curing temperature: 70 °C.*

For obtaining the optimized condition of modification in Sect. 2.3, orthogonal experiments for testing tensile strength were performed with three variables (i.e., ethanol volume/mL, immersing time/day, and curing temperature/°C), and each tensile strength presented in Table 1 was the average of three records. From Table 1, the tensile strength of 64.6 MPa was obtained in 300 mL ethanol at 70 °C for 1 day. More ethanol and less immersion time are beneficial for the film regeneration and maintaining the integrity of obtained films, respectively. Excessing immersing time may slightly destroy the interaction bonding, thus declining the degree of crosslinking (Hu & Wang, 2016). Similarly, higher temperatures will accelerate the crosslinking to strengthen the internal interconnection.

In addition, all films exhibit high tensile strength and relevant strain-to-failure values (see Fig. 4). The pure CF shows a tensile strength of 55.1 MPa and a strain-to-failure value of 18.5 %. After modification, these two values all increase significantly to 66.7 MPa and 20.8 %, respectively, which demonstrates that strong crosslinking bonds are formed after “bridge monomers” addition to extremely improve the mechanical properties. With the incorporation of 10 wt% SLS, the value of tensile strength rises obviously to 67.8 MPa, while that of strain-to-failure declines to 10.6 %. Further increasing SLS addition, both tensile strength and strain-to-failure value all dropped rapidly, which indicated that the strong interaction hydrogen bonding is created between SLS and cellulose film, as proven in FTIR, but phase separation is subsequently formed (see SEM images in Fig. 2), breaking the compatibility of blending system, thus leading to the degradation of mechanical properties. As for SLS10-CF-M, the tensile strength (67.8 MPa) is
higher than that of other reported mixed cellulose films, such as glutaraldehyde-polyvinyl alcohol/cellulose film (5 wt% P-G/RC, 66.3 MPa) (Hou et al., 2019), 5 wt% nanofibrillated cellulose (NFC)/poly(vinyl alcohol) (PVA) film (44.3 MPa) (Xiao et al., 2016), 32 wt% α-cellulose nanofibers (CHFs)/PVA film (39.0 MPa) (Cai et al., 2016).

3.5. Contact angle and water absorption capacity

Figure 5 shows the surface contact angle of different samples, which demonstrates hydrophilic and hydrophobic properties of testing materials, therefore exhibiting water adsorption capacity (see Fig. 6, the average amounts of water absorption of pure CF, SLS$_{10-40}$-CF-M are 66.18, 46.73, 57.44, 56.58, and 64.01 %, respectively). All contact angles are less than 90°, indicating that all samples are hydrophilic (Wang et al., 2019). However, with the incorporation of SLS, the contact angle of mixed films increases by 10–30°, where the highest value of contact angle is obtained at 10 wt% SLS addition. Generally, there are abundant hydrophilic groups in the internal structure of SLS, such as hydroxyl and sulfonate groups. Nevertheless, the worse hydrophilic capacity of mixed films is obtained after SLS addition, and the main reason is that hydroxyl groups in both SLS and cellulose can be interacted with each other to form crosslinking structure. In addition, the formation of “bridge linking” is positive for the decrease of hydroxyl groups, thus lowering hydrophilic ability (Llorens et al., 2015). The excess addition of SLS is negative for the blending film to obtain a compatible system as separation phase (SLS region) is formed on the surface of film, therefore increasing their hydrophilic capacity.

3.6. Light transmittance

The UV transmission of pure CF and modified mixed films was recorded in the range of 200–600 nm, and their transmission curves are shown in Fig. 7. The transmission of pure CF reaches 69.6 % at 280 nm, while that of SLS$_{10}$-CF-M is 1.2 %, which almost appears opaque. Further increasing the SLS addition from 10 to 40 wt% significantly decreases the transmission of blending films, almost close to 0 %, showing excellent UV blocking performance. These observations suggest that the extra addition of SLS makes the film surface bumpy (see SEM images), thereby inducing optical scattering and refraction. Therefore, with assistance of “bridge linking” and interaction hydrogen bonding, the presence of SLS plays an important role in blocking UV transmission, which make the blending films potential in UV-shielding application.

3.7. Water vapor permeance (WVP)
The water vapor permeance (WVP) of different samples is shown in Table 2. The pure film exhibits the highest WVP (104.12 g/day m$^2$). With the addition of SLS (10 wt%), the WVP of blending film decreases significantly from 104.12 to 56.49 g/day m$^2$, which may be due to the crosslinking effect from “bridge linking” and intense interaction hydrogen bonding between lignin and cellulose to decrease the inter-chain space, thus declining diffusivity and solubility of water molecules, which is corresponding to the result obtained from water absorption and contact angle. However, the hydrophilic property of all films (the contact angles are all less than 90°) give them appropriate values of WVP, which are much higher than that of other cellulosic composite films (El Miri et al., 2015; Karkhanis et al., 2018; Liu et al., 2018). With the increase of SLS addition (from 10 to 40 wt%), the WVP of composite films increases from 56.49 to 83.79 g/day m$^2$. Excess amount of SLS results in phase separation in the film surface, thus making the gap between molecules larger to accelerate the diffusion of water molecules.

### 3.8. Antibacterial property

The antibacterial ability of different samples was evaluated via measuring the diameter (mm) of inhibition zones, a classic method, which adopted film samples on the central location of agar medium coated with bacteria, and observed the inhibition of growth and propagation of bacteria (*E. coli*) to form a blank circle region around the samples, which was generally named as inhibition zone. According to the result in Fig. 8, the original CF presented as a control sample almost exhibits no inhibition zone against *E. coli*, while an apparent inhibition zone is observed after 10 wt% SLS addition in Fig. 8b, indicating the important role of SLS on antibacterial performance. Increasing the addition of SLS from 10 to 40 wt%, the obtained inhibition zone becomes larger and larger (from 8.0 mm for SLS$_{10}$-CF-M to 11.0 mm for SLS$_{40}$-CF-M), which demonstrates that the presence of SLS is beneficial for inhibiting the growth of testing bacteria. Generally, lignin (from hardwood or softwood) contains abundant phenolic hydroxyl groups, which are effective for inhibiting oxidation and bacteria growth (such as *P. aeruginosa, S. enteritidis, E. coli, B. cereus, S. aureus*) (Lourencon et al., 2021; Wang et al., 2021). In addition, the special presence of
sulfonate group in SLS as compared with other lignin is also an effectively antibacterial factor (Asadullah et al., 2020).

4. Conclusions

In this study, SLS$_x$-CF-M composite films were fabricated, and FTIR result demonstrated the completion of blending film preparation. The crosslinking structure formed by interaction hydrogen bonds and “bridge linkages”, and separation phase induced by the increasing addition of SLS were two main reasons for the change of mechanical and hydrophobic properties, while the UV transmission and antibacterial performance were all influenced by the addition of SLS. Treating SLS as a renewable alternative resource of other synthetic polyols or bacteriostatic agent for film modification meets the requirement of sustainable development

Declarations

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Conflicts of interest/Competing interests

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

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