Hassan, Ghada; Forsman, Nina; Wan, Xing; Keurulainen, Leena; Bimbo, Luis M.; Johansson, Leena Sisko; Sipari, Nina; Yli-Kauhaluoma, Jari; Zimmermann, Ralf; Stehl, Susanne; Werner, Carsten; Saris, Per E.J.; Österberg, Monika; Moreira, Vânia M.

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*Published in:*  
ACS Sustainable Chemistry and Engineering

*DOI:*  
10.1021/acssuschemeng.8b05658

Published: 04/03/2019

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

*Please cite the original version:*  
Dehydroabietylamine-Based Cellulose Nanofibril Films: A New Class of Sustainable Biomaterials for Highly Efficient, Broad-Spectrum Antimicrobial Effects

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Supporting Information

ABSTRACT: The design of antimicrobial surfaces as integral parts of advanced biomaterials is nowadays a high research priority, as the accumulation of microorganisms on surfaces incurs substantial costs on the health and industry sectors. At present, there is a growing interest in designing functional materials from polymers abundant in nature, such as cellulose, that combine sustainability with outstanding mechanical properties and economic production. There is also the need to find suitable replacements for antimicrobial silver-based agents due to environmental toxicity and spread of resistance to metal antimicrobials. Herein we report the unprecedented decoration of cellulose nanofibril (CNF) films with dehydroabietylamine 1 (CNF-CMC-1), to give an innovative contact-active surface active against Gram-positive and Gram-negative bacteria including the methicillin-resistant S. aureus MRSA14TK301, with low potential to spread resistance and good biocompatibility, all achieved with low surface coverage. CNF-CMC-1 was particularly effective against S. aureus ATCC12528, causing virtually complete reduction of the total cells from 10⁵ colony forming units (CFU)/mL bacterial suspensions, after 24 h of contact. This gentle chemical modification of the surface of CNF fully retained the beneficial properties of the original film, including moisture buffering and strength, relevant in many potential applications. Our originally designed surface represents a new class of ecofriendly biomaterials that optimizes the performance of CNF by adding antimicrobial properties without the need for environmentally toxic silver.

KEYWORDS: Nanocellulose, Dehydroabietylamine, Antimicrobial, Drug-resistant, Silver, Biomaterials

INTRODUCTION

Preventing the accumulation of microorganisms on surfaces is critical in many different applications including textiles, furniture, food packaging, and industrial and marine equipment.1,2 Therefore, there is a pressing need for antimicrobial surfaces that can efficiently prevent and/or eliminate fouling by bacteria, to reduce health risks, extend the life span of consumer/industrial goods, and allow for general convenience. In the design of new antibacterial surfaces or as a means to enhance the performance of existing ones, a plethora of strategies have been reported that include surface coating, chemical modification, or patterning.3–9 Common drawbacks include failure in achieving efficacy and biocompatibility levels suitable for specific target applications, extreme chemical complexity that reflects on cost-effectiveness, and, last but not least, the use of synthetic polymers and/or environmentally hazardous reaction conditions.
Cellulose, with an estimated production of $1.5 \times 10^{12}$ tons in the biosphere annually, is the most abundant naturally occurring polymer on earth and a virtually inexhaustible source of raw material for the design of environmentally friendly and biocompatible products to replace fossil fuel derived ones. Cellulose nanofibril (CNF) is considered as an advanced, fully renewable material with high potential to reach commercial applications in a nearby future. CNF is prepared by mechanical disintegration of cellulose fibers via high shearing followed by homogenization and can nowadays be obtained from either wood or crops with reasonably low energy consumption. The appealing properties of CNF, including global abundance and renewability as well as high stiffness and lightweight character, are a consequence of both its natural origin and its chemical structure, i.e. of the way it is organized into fibrils with length in the micrometer and width in the nanometer range, forming a network with both amorphous and crystalline regions. In particular, thin films can be made of CNF that are robust, solvent-resistant, and can be surface-modified for acquiring functionality.

**RESULTS AND DISCUSSION**

Carbodiimide chemistry was selected in order to couple 1 to CNF after irreversible adsorption of sodium carboxymethyl cellulose (Na-CMC) to the film surface (Figure 1A–C). The
reaction with Na-CMC, carried out in low salt concentration aqueous solution, enriched the surface of the films in carboxyl groups that were then covalently linked to 1 through the amidation reaction. Both methods employ relatively nontoxic and inexpensive reagents and ethanol, an environmentally preferable solvent,31 was used in the second step instead of N,N-dimethylformamide (DMF), the usual solvent of choice for coupling reactions.

Contact angle (CA) measurements at 5 s provided a rough estimate of the success of the modification, as it was expected that the binding of 1 to the surface of the CNF film would render it more hydrophobic due to an increase in carbon content (Figure 2A). Indeed, the CA value was much higher for CNF-CMC-1 than for CNF and CNF-CMC. X-ray photoelectron spectroscopy (XPS) analysis revealed that the modified CNF surface in CNF-CMC-1 contained nitrogen and carbon atoms bound only to other carbon atoms with no bonds to oxygen (C=C), inexistent in cellulose (and therefore in CNF), which could only have originated from our chemical modification (Figure 2B1–B2, Table 1). CNF-CMC-1 also displayed lower content of carbon atoms with one bond to oxygen than the reference film. We estimated that compound 1 covered only 20% of the surface based on the N content and 25% based on C=C content, indicating very low surface coverage.

In line with the low coverage value, atomic force microscopy (AFM) height images (Figure 3A) showed that the surface of cellulose fibrils on CNF-CMC-1 did not significantly differ from that of CNF. Fourier-transform infrared spectroscopy (FTIR) analysis furthermore verified that only the outermost surface was modified, since no dramatic changes were detected between modified and unmodified films (Figure 3B). The ability to permeate gas or vapor and the mechanical properties of CNF-CMC-1 were also not significantly affected when compared to CNF (Table S1) apart from a slightly higher oxygen permeability (OP) value at low (50%) relative humidity (RH), which is consistent with the increase in the hydrophobicity of the films after modification with 1 that results in the ability to attract a higher number of oxygen molecules. It is generally accepted that high surface roughness and low surface energy increase hydrophobicity.32 In CNF-CMC-1, the hydrophobicity is most likely caused by the particular surface chemistry, as the AFM data showed no significant changes in surface roughness. The lack of impact of the modification of CNF with 1 on the barrier performance is yet another reflection of the low surface coverage as an increase in this performance in humid environment can only be attained if the CNF film surface is fully covered.33

Overall, the surface modification of CNF with 1 was gentle, yet effective, producing an increase in hydrophobicity that did not result in deterioration of the breathability, i.e., the film’s water vapor and oxygen gas permeability,34 nor of the moisture buffering properties, i.e., the film’s ability to adsorb and release moisture in the surrounding environment thus dampening changes in relative humidity,35 which are beneficial for many applications.33 This is in sharp contrast with modifications made with hydrophobic compounds prior to film formation that typically disturb the hydrogen bonding network resulting in loss of both barrier and strength performance.36

CNF-CMC-1 showed clear and robust antimicrobial activity against different bacterial strains including the Gram positive Staphylococcus aureus, the Gram negative Escherichia coli, and the methicillin-resistant S. aureus MRSA14TK301, whereas CNF and CNF-CMC were inactive (Figure 4A). CNF-CMC-1 was particularly effective against S. aureus ATCC12528 where reduction of the total cells from 10^5 CFU/mL bacterial suspensions was virtually complete after 24 h of contact. Potent activity was also observed against the methicillin-resistant strain and E. coli, however less pronounced than that observed against the normal S. aureus strains. CNF-CMC-1 did not endure treatment with strong acid but retained antimicrobial activity even after exposure to different solvents, 

Table 1. High Resolution Numerical XPS Data

<table>
<thead>
<tr>
<th>sample</th>
<th>Wide Scan Atomic Concentrations (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C 1s</td>
</tr>
<tr>
<td>CNF</td>
<td>60.7</td>
</tr>
<tr>
<td>CNF-CMC-1</td>
<td>67.0</td>
</tr>
<tr>
<td>CNF-CMC-4</td>
<td>62.2</td>
</tr>
</tbody>
</table>

![Figure 2](image)
including strong base, at room temperature, for 24 h (Table S2).

In contrast, compound 1 is a relatively weak antimicrobial agent against the tested strains, with minimal inhibitory concentrations (MIC) values in the 40–50 μM range (Table S3, Figure 5A). Incubation of either *S. aureus* ATCC12598 or *E. coli* DH5α with 100 μM of 1, for 4 h, resulted in death of some bacteria in the medium, with slight leakage of ATP from *S. aureus* cells, suggesting that 1 is a slow acting compound (Figure 5B).

The factors governing bacterial interaction with surfaces are well-known and include surface charge, hydrophobicity, topography, and chemical environment.2,37 Streaming current measurements (Figure 3C) revealed the typical electrokinetic fingerprint of an oxidized cellulose38 for CNF, CNF-CMC, and CNF-CMC-1. The point of zero streaming current was found below pH 3, and the current reached a plateau in the neutral and alkaline pH range indicating that all surfaces are negatively charged under the conditions of the bacterial assay. Compared to CNF, a slightly lower negative streaming current was measured for CNF-CMC and CNF-CMC-1 as a result of the increased friction exerted by the soft CMC layer39 on the hydrodynamic flow at the interface.40 In view of the electrokinetics and AFM data, surface charge and topography do not seem to be determinants of the antimicrobial activity of CNF-CMC-1.

To probe the impact of small changes in chemical environment at the CNF films surface, we synthesized CNF-CMC-4 using compound 4 (Figure 1C–D), a derivative of 1, in order to determine the effects of exposing a protonable amino group at the surface of the CNF, as the presence of cationic groups in antimicrobials is known to result in increased potency albeit accompanied by potential toxicity.2 Compound 4 was made from 1 as depicted on Figure 1C. As before, the XPS analysis confirmed the desired modification and coverage was calculated as 18% based on N and 14% based on C (Figure 2B1–2B2, Table 1). In this case, the steric orientation of compound 4 at the surface of CNF is unknown and probably differs between molecules, as there is mobility in the carbon chain between the two N atoms, which further hampers calculation. In CNF-CMC-4, the increase in the CA value was less pronounced than in CNF-CMC-1 (Figure 2A) and the electrokinetic fingerprint was similar to that of CNF-CMC and CNF-CMC-1 (Figure 3C), building up the evidence for low decoration of the surface with 1 or 4. Nonetheless, the inclusion of 4 at the surface of CNF films dramatically impacted Gram positive staphylococci cells, which were killed after 24 h of contact with CNF-CMC-4 (Figure 4A). A sharp reduction (98.3% of total cells) from the suspension of methicillin-resistant *S. aureus* MRSA 14TK301 and *E. coli* (99.9%) was also observed. We found that the antimicrobial potency of 4 was only moderately better than that of compound 1, with MIC values still in the high micromolar range (Table S3, Figure 5A). However, bacterial cell death with ATP efflux in *S. aureus* ATCC12598 was much more evident after treatment with 4, indicating that this compound causes bacterial membrane disruption rapidly (Figure 5B).

As depicted on Figure 4B, the cell surface became rough after treatment with compound 1 for 1 h when compared to untreated *S. aureus*, whereas treatment with 4 caused

![Figure 3. AFM images (A1, A3 height; A2, A4 adhesion), FTIR spectra (B), and streaming current vs pressure gradient (C) for representative samples.](image-url)
significant morphological changes on the cells and obvious debris appeared. The bacterial cells contacted with CNF looked unharmed especially when compared to those incubated with CNF-CMC-4, were abundant debris was visible in the background and the few remaining cells look highly deformed. No significant changes where visible on the cells in contact with CNF-CMC-1. Moreover, biocompatibility assessment of CNF-CMC-1 and CNF-CMC-4 (Table 2) revealed that CNF-CMC-1 was generally better tolerated by human erythrocytes than CNF-CMC-4, as the percent hemolysis caused was negligible. Human skin fibroblasts colonized CNF-CMC-1 to an extent comparable to that observed with CNF, which is regarded as a biocompatible biomaterial.\(^{11,14}\) For CNF-CMC-4, the hemolysis rate was 6.5% and in sharp contrast with CNF-CMC-1, fibroblasts did not proliferate on top of its surface.

Overall, the combination of 1 and 4 with CNF films resulted in hybrid surfaces displaying broad and potent antimicrobial
DH5 ATP from DH5 concentration was used as control. and 4 for the functionalization, more than charge, topography or clearly show that small changes in the diterpene used CMC-4 μ (100 ATCC12598; II
compared with the percent of viable a fibroblasts grown on a sterile tissue culture-treated 24-well plate. Figure 5. A
incubated with compound α incubated with compound α incubated with compound 1 h; III S. aureus ATCC12598 incubated with compound 4 (100 μM) for 1 h; IV nontreated. E. coli DHSer incubated with compound 1 (100 μM) for 1 h; VI E. coli DHSer incubated with compound 4 (100 μM) for 1 h. B. Efflux of ATP from S. aureus ATCC12598 after incubation with compounds I and 4 (100 μM). Bacteriostatic vancomycin with 100 μM final concentration was used as control.

activity despite the low coverage. The differences observed in the activity and biocompatibility of CNF-CMC-1 and CNF-CMC-4 clearly show that small changes in the diterpene used for the functionalization, more than charge, topography or hydrophobicity, will significantly impact the chemical environ-

**CONCLUSION**

Herein we set the grounds for a facile, ecofriendly, and mild strategy for immobilization of antimicrobial diterpenes onto CNF, leading to innovative and sustainable silver-free antibacterial films, without compromising the original physicochemical properties of the nanocellulose. The proposed mode of action, along with the fact that the films are based on compounds from a new chemical class, should account for a low potential to spread resistance. Furthermore, the potency and biocompatibility of CNF-CMC-1 warrant additional investigation for applications in the biomedical field where surfaces able to minimize the risk of infection at relevant bacterial bioburden are needed. Efforts are currently underway to investigate this possibility as well as the applicability of our design strategy to other biopolymers.

**EXPERIMENTAL SECTION**

Chemistry. General. (+)-Dehydroabietylamine was purchased from TCI, Europe. N,N-Diisopropylethylamine (DIEA), 1-hydroxybenzotriazole hydrate (HOBt), di-i-butyl dicarbonate, cesium carbonate, sodium carboxymethyl cellulose (DS 0.70–0.85, MW ~ 250 000), N,N-dimethylformamide (DMF), and dichloromethane were acquired from Sigma-Aldrich Co. 3-Bromopropylamine hydrobromide was purchased from Ega-Chemie. N-(3-Dimethylaminopropyl)-N′-ethylcarboxydimide hydrochloride (EDC) and trifluoroacetic acid (TFA) were purchased from Fluorochem Ltd. Calcium chloride was purchased from Merck, sodium hydrogen carbonate, from VWR International Oy, methanol, from VWR Chemicals, and ethanol (Etch A, 94% w/w), from Altia Oy. All reagents were used without purification apart from dehydroabietylamine, which was purified by flash column chromatography (FCC) using an ethyl acetate: 10% w/w NH₄OH in methanol gradient 10 → 60%, when used as the starting material for the synthesis of compound 4. For thin layer chromatography (TLC) silica gel 60 F254 was used. FCC was made with a Biotage high-performance flash chromatography Sp4-system (Uppsala, Sweden) using a 0.1 mm path length flow cell UV detector/recorder module (fixed wavelength: 254 nm), and 25, 50, or 100 g SNAP cartridges (25–100 mL/min flow rate).

![Figure S. A. Fluorescence microscopy image of bacteria counts with/ without treatment with compounds 1 and 4. I nontreated S. aureus ATCC12598; II S. aureus ATCC12598 incubated with compound 1 (100 μM) for 1 h; III S. aureus ATCC12598 incubated with compound 4 (100 μM) for 1 h; IV nontreated E. coli DHSer; V E. coli DHSer incubated with compound 1 (100 μM) for 1 h; VI E. coli DHSer incubated with compound 4 (100 μM) for 1 h. B. Efflux of ATP from S. aureus ATCC12598 after incubation with compounds I and 4 (100 μM). Bacteriostatic vancomycin with 100 μM final concentration was used as control.](image)

### Table 2. Biocompatibility Assessment

<table>
<thead>
<tr>
<th>assay</th>
<th>CNF</th>
<th>CNF-CMC-1</th>
<th>CNF-CMC-4</th>
<th>Triton X-100 %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemolysis*</td>
<td>0</td>
<td>0.9 ± 0.6</td>
<td>6.5 ± 1.9**</td>
<td>100 ± 3.0***</td>
</tr>
<tr>
<td>Fibroblast proliferation 72 h**</td>
<td>59.8 ± 7.3</td>
<td>47.2 ± 17.7</td>
<td>2.7 ± 0.4****</td>
<td>12 ± 0.4***</td>
</tr>
</tbody>
</table>

*Measured in terms of percent of hemolyzed erythrocytes. **Measured in terms of percent of viable fibroblasts placed on top of each material compared with the percent of viable fibroblasts grown on a sterile tissue culture-treated 24-well plate. *Statistical analysis was made by ANOVA, followed by a Dunnett’s multiple comparison test. All data sets were compared with CNF for both hemolysis and fibroblast proliferation. The level of significance was set at probabilities of * p < 0.05, ** p < 0.01, and *** p < 0.001.
Infrared (IR) spectra were recorded on a Vertex 70 (Bruker Optics Inc., MA, USA) FTIR instrument with a horizontal attenuated total reflectance (ATR) accessory (MIRacle, Pike Technology, Inc., WI, USA). The transmission spectra were recorded at a 4 cm\(^{-1}\) resolution, between 4000 and 600 cm\(^{-1}\), using the OPUS S.5 software (Bruker Optics Inc., MA, USA). The graphs in Figure 3B are normalized by the 1159 cm\(^{-1}\) band. Nuclear magnetic resonance (NMR) spectra were recorded on a Bruker Ascend 400 spectrometer, with tetramethylsilane (TMS) as the internal standard. The coupling constants (J) are quoted in Hertz (Hz). Exact mass analyses were performed with a Waters Acquity UPLC attached to Synapt G2 HDMS in positive ionization (ESI) mode (Waters, MA, USA).

**Preparation of CNF films.** Never-dried birch kraft pulp was used to prepare a CNF dispersion. To control the counterion type and ionic strength, the pulp was washed into sodium form prior to preparing 100 mL of 0.85% CNF through a Sefar Nitex uidizer. Free-standing CNF films were prepared by filtering 100 mL of 0.85% CNF through a Sefar Nitex polyamine monofilm (24 μm pore size, at 2.5 bar pressure, for 30 min. The film was then hot-pressed for 2 h at 100 °C and 1800 kg/cm\(^2\) pressure in a Carver Laboratory press (Fred S. Carver Inc.). The prepared films were stored at standard conditions (23 °C and 50% RH). More detailed information about freestanding CNF films and their production can be found elsewhere.4 CNF films were cut into circles with a 5 cm diameter for the experiments reported below.

**Activation of the CNF Film Surface by Carboxymethylation (CNF-CMC).** To a round-bottomed reaction flask containing a 0.05 M/0.01 M solution of CaCl\(_2\)/NaHCO\(_3\) in water (75 mL), Na-CMC (270 mg) was added. The mixture was stirred for 5 min, at 80 °C. One CNF film circle was then added to the mixture and the left to agitate with magnetic stirring at 80 °C, for 4 h. The reaction mixture was poured out from the flask, and the CNF film was washed with the following solutions under magnetic stirring at room temperature: deionized water (150 mL, 10 min), a 0.1 M solution of CH\(_3\)COOH in water (75 mL, 10 min), deionized water (150 mL, 10 min), a 0.4% solution of NaHCO\(_3\) in water (75 mL, 1 h), and deionized water (150 mL, 10 min). The CNF film circle was recovered from the aqueous solution, placed between 2 blotting sheets, and left in an oven to dry overnight at 103 °C. Amidation of Activated CNF Films (CNF-CMC-1 and CNF-CMC-4). One CNF film circle activated as described above was added to a round-bottomed flask containing EDC hydrochloride (0.02 M) and DIPEA (0.04 M) were added, and the mixture was left to agitate at room temperature for additional 24 h. The reaction mixture was then poured out from the reaction flask and the CNF film circle was first rinsed with 94% w/w ethanol (20 mL) and then washed under magnetic stirring with deionized water (4 × 150 mL, 10 min each washing). The modified CNF film circle was recovered from the aqueous solution, placed between 2 blotting papers, and left in an oven to dry overnight at 103 °C.

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