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Abstract. In this study, we investigate the interactions between the cellulose surface and Ag nanoparticles (AgNPs) for the purpose of manufacturing hybrid nanomaterials using bacterial cellulose nanocrystals (BCNs) as a model substrate. We focus on the role of the BCN surface chemistry on the AgNP nucleation obtained by chemical reduction of Ag\textsuperscript{+} ions. Homogeneous hybrid suspensions of BCN/AgNP are produced, regardless of whether the BCNs are quasi-neutral, negatively (TBCNs) or positively charged (ABCNs). The characterization of BCN/AgNP hybrids identifies the \textasciitilde OH surface groups as nucleation points for AgNPs, of about 20 nm revealing that surface charges only improve the accessibility to OH groups. X-ray Absorption technics (XANES and EXAFS) revealed a high metallic Ag\textsubscript{0} content ranging from 88\% to 97\%. Moreover, the grafting of hydrophobic molecules on a BCN surface (HBCNs) does not prevent AgNP nucleation, illustrating the versatility of our method and the possibility to obtain bifunctional NPs. A H\textsubscript{2}O\textsubscript{2} redox post-treatment on the hybrid induces an increase in AgNPs size, up to 90 nm as well as a shape variation (i.e., triangular). In contrast, H\textsubscript{2}O\textsubscript{2} induces no size/shape variation for aggregated hybrids, emphasizing that the accessibility to \textasciitilde OH groups ensures the nucleation of bigger Ag nano-objects.
Keywords. Bacterial cellulose nanocrystal (BCN), hydroxyl groups, surface modification, silver nanoparticles (AgNPs), AgNP size-shape variation, hydrophobic BCN, hybrid nanomaterial.
1. Introduction

Cellulose is an almost inexhaustible biopolymer extracted from wood, cotton, algae, tunicates or bacteria [1–3], leading to variation in dimensions and structural organization [4]. Cellulose displays interesting properties such as high water retention capacity, high wet strength, low density, biocompatibility, non-toxicity and biodegradability, which make it very appropriate for several applications in biomedicine and pharmacology [5–7], cosmetics [8], paper and textiles [9]. Cellulose can also be hydrolyzed to increase its crystallinity, commonly by acid hydrolysis, leading to the so-called cellulose nanocrystals (CNCs) [10]. While the extraction of plant-sourced cellulose often requires the use of hazardous chemicals, bacterial cellulose (BC) is mainly secreted extracellularly by certain bacteria such as *Gluconacetobacter xylinus* [1]. Therefore, bacterial cellulose nanocrystal (BCN) contain no functional groups other than hydroxyl groups [11] and thanks to its high chemical purity, they are particularly interesting for food and medical applications.

To confer antimicrobial properties to CNCs, as to BCNs, it is quite common to graft or nucleate metallic nanoparticles (e.g., Cu, Ag, ZnO, Au) on their surface [12–15]. Among others, silver nanoparticles (AgNPs) have been widely studied. A silver precursor is generally incorporated by addition of a salt such as silver nitrate (AgNO$_3$), and several experimental methods (e.g., UV radiation [16,17], hydrazine reduction [18]) have been proposed to nucleate AgNPs on the cellulose nanocrystal surface, thus obtaining a hybrid material (i.e., CNC/AgNPs). Chemical reduction is one of the most common ways to synthesize metallic NPs, and sodium borohydride (NaBH$_4$) [19–22] is reported to be one of the most efficient reducing agents, inducing the rapid formation of 2-3-nm AgNPs [23]. NH$_4$OH [24] and ascorbic acid [25] can also be used to reduce Ag$^+$ ions into AgNPs. To carry out this chemical AgNP synthesis, capping agents or stabilizers (e.g., trisodium citrate: TSC; polyvinylpirrolidine: PVP; Cetrimonium bromide: CTAB) are often introduced to better control the morphological properties of AgNPs, preventing their aggregation [26–28]. In other cases, nanocellulose is proposed as both a support and a reducer for the generation of metallic nanoparticles [29]. Xiong et al. [30] claim that cellulose promotes the generation of AgNPs and dendritic Ag nanostructures thanks to the extensive presence of surface hydroxyl groups. Furthermore, various surface modifications of CNCs can be performed (e.g., amidation, oxidation, esterification, etherification) [10], possibly affecting the reduction of AgNPs on the CNC surface. Most of the studies concerning CNC/AgNP hybrids [31–33] indicate how the introduction of negative surface charges (i.e., via TEMPO-mediated oxidation [32]) can provide the high-binding capability for the transition of metal species such as Ag$^+$ ions. Moreover, other surface modifications can be performed on the CNC surface to extend the possible application fields. For example, cationized cellulose can be produced by grafting different molecules on the cellulose surface (e.g., HPTMAC: hydroxypropyltrimethylammonium chloride [34], BriBB: a-bromoisobutyryl bromide [35], HDTMA: hexadecyltrimethylammonium [36] or pyridinium [37]) so that it can be used for papermaking, cosmetics and drug-delivery [38]. The work of Shateri Khalil-Abad et al. [39] reveals that surface-
modification of cellulose cotton fibers by 3-chloro-2-hydroxypropyl trimethyl ammonium chloride (CHPTAC) affects the absorption of AgNPs on the fiber surface. Indeed, such an adsorption is greater on cationized cellulose fibers than on untreated ones. To obtain these cationized fibers, Dong et al. [40] propose to graft ammonium ions onto the surface, thus creating two different pathways for the deposition of metal NPs (e.g., Au, Pt, Pd): the first one based on the electrostatic assembly of metal nanoparticles capped with negative citrate ions, and a second one where negative metal complex ions are adsorbed onto the cationic substrate and then reduced. Both approaches lead to a high surface coverage of the CNC surface by metallic nanoparticles.

CNCs can also be made hydrophobic to allow their dispersion in non-aqueous solvents. To do this, one of the most widely used approaches is amidation, where coupling agents are grafted onto TEMPO-oxidized nanocrystals [10]. Araki et al. [41] propose the grafting of poly(ethylene glycol) with a terminal amino group onto the surface of TEMPO-oxidized CNCs. Zhang et al. [42] use a similar experimental approach to covalently functionalize CNCs via peptide coupling chemistry with different alkylamines (e.g., propylamine, n-butyamine, amylamine, hexylamine, heptylamine). Hu et al. [43] propose using tannic acid to covalently attach primary amine with a long alkyl chain (i.e., decylamine) to the CNC surface. In addition, Cunha et al. [44] produced inverse emulsions, tailoring the hydrophobicity of CNCs and nanofibrillated cellulose by chemical modification with lauroyl chloride (C_{12}). However, no studies have proposed bifunctional hybrids, using hydrophobically-modified CNCs as substrates for AgNP nucleation.

As for nanocellulose surface modification, AgNP characteristics (e.g., size, shape, crystalline structure) can strongly influence the final properties of the CNC/AgNP hybrid system. Some works report simple techniques for a shape- and size-selective synthesis of Ag nanostructures (e.g., from spherical AgNPs to triangular-shaped AgNPs or Ag nanoprisms, denoted AgNPrisms [45]) based on the use of hydrogen peroxide (H_{2}O_{2}) redox post-treatment [20,21,46,47], as well as photo-induced [48] or solution-phase [49] methods. In particular, the size-shape transformation of AgNPs achieved with H_{2}O_{2} relies to redox capabilities linked to the autocatalytic decomposition on the Ag surface. These AgNPrisms are anisotropic (i.e., their lateral dimension is greater than their thickness) and are thus characterized by a strong localized surface plasmon resonance (LSPR). Their resulting size- and shape-dependent optical properties make them suitable for sensors [50,51] and biological imaging [52], catalysts [53], nanophotonic devices and circuits [54,55]. Parnklang et al. [46] show the shape transformation from spherical AgNPs to AgNPrisms when H_{2}O_{2} oxidizes the AgNPs to Ag^{+} ions and thus reduces them into Ag atoms in nanometric silver particles. The H_{2}O_{2} redox post-treatment has also been applied to hybrid systems. Jiang et al. [20] propose the fabrication of films made of AgNPrisms and TEMPO-oxidized cellulose nanofibrils, which work as capping and shape-regulating agents. The presence of a predominant (111) peak in the XRD pattern allows them to affirm that AgNPrisms are bounded to cellulose nanofibers.
The result is that hybrids made of AgNPs fixed on the CNC surface have been produced, whereas the interactions and binding mechanisms have not been studied in detail [29,31,56]. In this paper, we propose to perform different surface modifications on BCNs in order to shed light on the effective role of cellulose nanocrystal surface chemistry on the nucleation of AgNPs. We also prepared bifunctional NPs that combine a hydrophobic surface and AgNPs. Moreover, we investigated the impact of the CNC surface modification on the H₂O₂ redox post-treatment, linking the size and shape variation of AgNPs to their physicochemical characteristics (i.e., AgNP structure and oxidation state).

2. Materials and Methods

**Chemicals.** Food grade nata de coco was purchased from AROY-D (Thailand). All the other products were purchased from Sigma-Aldrich (France) and used as received without further purification: silver nitrate (AgNO₃ ≥ 99%), sodium borohydride (NaBH₄ ≥ 96%), 2,2,6,6-Tetramethylpiperidine 1-oxyl (TEMPO: 98.8%), sodium bromide (NaBr: 99%), sodium hypochlorite solution (NaClO: 10% RT), cholaminchloride hydrochloride (AETMA: 99%; pKa = 7), octylamine (C₈: 99%; pKa = 10), N-hydroxysuccinimide (NHS), N-(3-dimethylaminopropyl)-N′-ethylicarbodiimide (EDC), hydrogen peroxide 30% (H₂O₂). For all the aqueous suspensions, ultra-pure water was used.

**Synthesis of uncharged BCNs.** For the preparation of BCNs, the experimental procedure proposed by Kalashnikova et al. [57] was followed. Briefly, nata de coco cubes were dialyzed in ultra-pure water for 9 days, after which they were ground in a Waring blender, suspended in 0.5 N NaOH solution, and stirred for 2 h at 70°C. Finally, the sample was rinsed using ultra-pure water. A whitening treatment was performed twice, dispersing the slurry in a NaClO₂ solution (8.5 g/L) in a sodium acetate buffer (pH = 4.5) and stirring it for 2 h at 70°C. The bleached cellulose was rinsed with distilled water and then hydrolyzed with 2.5 M HCl solution at 70°C for 2 h. The suspension was centrifuged three times (10 min., 10000 g) and dispersed in ultra-pure water up to pH > 5. The final BCN suspension was then dialyzed against ultra-pure water for one week and finally homogenized by ultrasound (amplitude = 20; 5 min). The BCN content in the suspension (around 2 g/L) was determined by a drying test.

**Synthesis of anionic BCNs mediated via TEMPO oxidation (TBCNs).** The experimental protocol for the BCN surface carboxylation via TEMPO-mediated oxidation was adapted from the method proposed by Araki et al. [58]. A quantity of 100 mL of BCN suspension (2 g/L) was mixed with 0.25 g of NaBr and 0.05 g of TEMPO, after which 2.5 mL of NaClO were added to start the BCN carboxylation, leading to a pH increase. The resulting suspension was stirred for 24 h at room temperature while maintaining the pH at 10.3 through the controlled addition of 0.1 M NaOH solution using an automatic titrator (Metrohm 901 Titrando, France). The final suspension was centrifuged twice (10 min; 10000 g), dispersed in ultra-pure water while maintaining the BCN concentration at 2 g/L and, finally, homogenized by ultrasound (amplitude = 10; 5 min). The sample was dialyzed
against ultra-pure water for one week to remove any residual contaminants (dialysis bath volume to sample volume = 10:1).

**Synthesis of cationic BCNs (ABCNs).** For the BCN surface modification by AETMA, the coupling method proposed by Zhang et al. [42] was used with several modifications, working in a H₂O medium instead of N,N-dimethylformamide (DMF). A quantity of 100 mL of TBCN suspension (2 g/L) was mixed for 5 min at room temperature with 230 mg of EDC. Then, 180 mg of NHS were added and the suspension was stirred for a further 30 min. Finally, a quantity of 210 mg of AETMA were introduced and the sample was stirred for 24 h at room temperature while maintaining the pH at 7 through the controlled addition of 0.1 M NaOH solution using an automatic titrator (Metrohm 901 Titrando, France). The final suspension was centrifuged to remove unreacted residual molecules and dispersed in ultra-pure water to be homogenized by ultra-sound (amplitude = 10; 5 min). The suspension was dialyzed for one day against a KNO₃-saturated solution to promote ion exchange from chloride to nitrate, and then against ultra-pure water for one week (dialysis bath volume to sample volume = 10:1).

**Synthesis of hydrophobic BCNs (HBCNs).** To obtain HBCNs, the TBCNs were functionalized with alkyl chains (C₈) using a coupling method similar to the one previously proposed for the preparation of the ABCNs in suspension. A quantity of 60 mL of TBCN suspension (2 g/L) was stirred for 5 min at room temperature with 230 mg of EDC and for an additional 30 min after the addition of 180 mg of NHS. In another beaker, a volume of 270 μL of octylamine (C₈) were dissolved in 40 mL of absolute ethanol and added drop-by-drop to the TBCN suspension. The sample was stirred for 24 h at room temperature while maintaining the pH at 10 by the addition of 0.1 M NaOH solution controlled by an automatic titrator (Metrohm 901 Titrando, France). The suspension was then dialyzed against a 60/40 %v/v water/ethanol mixture for 3 days to remove the unreacted residual molecules. The final HBCN concentration was found to be around 2 g/L.

A schematic representation of the various BCN surface modifications and the corresponding scanning transmission electron microscopy (STEM) images are reported in Fig. 1.
Synthesis of BCN/AgNP hybrids. Both AgNO$_3$ and NaBH$_4$ solutions were freshly prepared just before the experimental session, and the NaBH$_4$ aqueous solution was placed in ice to minimize its decomposition. AgNO$_3$ aqueous solution (300 μL, 50 mM) was added to 10 mL of BCN water suspension and mixed for 1 min at room temperature. The NaBH$_4$ aqueous solution (500 μL, 100 mM) was then introduced to reduce Ag$^+$ ions to AgNPs, immediately turning the suspension from translucent to light-yellow. The final hybrid suspension was stirred at room temperature for 24 h, protected by light using aluminum foil to prevent silver oxidation, and then dialyzed against water for 24 h. The procedure produced hybrid suspensions with BCNs, independently of prior surface modifications of the BCN.

H$_2$O$_2$ redox post-treatment. The modification of AgNP characteristics was performed by adding H$_2$O$_2$ to the BCN/AgNP initial hybrid suspension. From now on, the AgNPs obtained after the addition of H$_2$O$_2$ will be generically indicated as AgNPs$_{H_2O_2}$ and, more specifically, as AgNPrisms when they reach a triangular shape. Immediately after the reduction by NaBH$_4$, various amounts of H$_2$O$_2$ (i.e., 0, 40, 80, 120, 160, 250 μL) were added to obtain various H$_2$O$_2$/AgNP mass ratios, $\alpha$ (i.e., 0, 0.08, 0.17, 0.25, 0.33, 0.52). The H$_2$O$_2$ introduction induces an exothermic reaction associated with the gas bubbles formation associated to the H$_2$O$_2$ decomposition [21]. The color of the sample gradually varied from yellow to blue with the introduction of H$_2$O$_2$. The final hybrid was dialyzed against ultra-pure water for 24 h.
Characterization. For FTIR measurements, the pure and surface-modified BCN suspensions were freeze-dried using a LYO GT2 lyophilizer (SRK System Technik, Germany; T = -90°C; 1.5 \(10^{-2}\) mbar). A quantity of 2 mg of freeze-dried sample was homogenized with 120 mg of KBr and the resulting powder was compressed (5 tons; 5 s) to obtain a circular pellet. The FTIR analysis were performed using a Nicolet iS50 FTIR spectrometer (Thermo Fisher Scientific, USA) and the spectra were determined to be in the 400 – 4000 cm\(^{-1}\) range, with a resolution of 4 cm\(^{-1}\). Each spectrum was the average of 200 scans collected in absorbance mode and examined using OMNIC\textsuperscript{TM} software (Thermo Fisher Scientific, USA).

The conductivity of BCN, TBCN, ABCN and HBCN aqueous suspensions was automatically measured by a Metrohm 856 Conductivity Module (France). The suspensions were titrated using 0.1 M aqueous NaOH (addition rate of 0.1 mL/30 s) [58]. The data were recorded by Tiamo\textsuperscript{TM} Titration software and the conductivity values were corrected from dilution effects and thus plotted against the added volume of sodium hydroxide solution. The inflection point was graphically determined from the intersection of the least squares regression lines fit and the data points in the distinct regions of the titration curves, evaluating the NaOH volume required for neutralization and, thus, the surface charge density [59,60], \(\rho\) (mol/g). Using \(\rho\) values, the degree of oxidation [61] (DO) was calculated as:

\[
\text{DO} = \frac{n_{\text{COOH}} \times M_{\text{AGU}}}{m_{\text{cellulose}} - n_{\text{COOH}} \times M_{\text{COOH}}}
\]

where \(n_{\text{COOH}}\) represents COOH moles (mol) deduced from the surface charge density, \(\rho\); \(M_{\text{AGU}}\) was the molecular weight of an anhydroglucose unit (162 g/mol); \(m_{\text{cellulose}}\) corresponded to BCN mass and \(M_{\text{COOH}}\) was the molecular weight of a carboxylic group.

The zeta potential values were measured using a ZetaSizer Nano ZS (Malvern, UK). BCN suspensions were diluted at 0.1 g/L and filtered (pore diam. = 5 µm). Five measurements at T = 20°C were performed for each sample. For the case of HBCNs suspended in a H\(_2\)O/EtOH mixture, the residual ethanol was removed by dialysis of several milliliters of suspension against water for 2 days. All the suspensions were sonicated just before the measurements.

X-ray Absorption Near-Edge Structure (XANES) measurements were performed to investigate the AgNP oxidization state (i.e., metallic silver, Ag\(_0\); ionic silver, Ag\(^+\)) while Extended X-ray Absorption Fine Structure (EXAFS) data made it possible to shed light on the AgNP bulk atomic structure (e.g., bond length, interatomic distance). Both XANES and EXAFS spectra were simultaneously recorded in transmission mode at the Ag K-edge (25250 to 27750 eV) on the SAMBA beamline at the SOLEIL synchrotron (Saint Aubin, France). The Si (220) monochromator was calibrated to 25515.6 eV at the first inflection point of the Ag foil XANES spectrum. To be analyzed, all the hybrids were freeze-dried and then compressed to obtain a circular pellet of 6 mm where the quantity of AgNP was high enough to reach an absorption edge jump close to 1. These pellets were placed on a sample rod,
quenched in liquid nitrogen and then introduced into the He cryostat (T = 20 K). Silver foil (Agfoil) and AgNO₃ aqueous solution with 1 wt% glycerol (AgH₂O) were used as standards. One continuous scan was recorded for each hybrid sample in the 25250 to 27750 eV energy range with a monochromator velocity of 5 eV/s and an integration time of 0.08 s/point. The obtained scans were then normalized and background-subtracted using the Athena software package [62]. The XANES spectra were analyzed by a linear combination fitting (LCF) procedure in the E₀ - 20 eV, E₀ + 50 eV energy range, with E₀ set to 25514 eV and using Agfoil and AgH₂O standards as components. All component weights were forced to be positive and the relative proportions of the components were forced to add up to 100%. Concerning the EXAFS oscillation, a background subtraction was performed before applying an autobk algorithm (Rbkb = 1, k-weight = 3). The Fourier transform of the k³-weighted EXAFS spectra was then calculated over a k range of 2.5-19.5 Å⁻¹, using a Hanning apodization window (width of the transition region window parameter = 1). The k³ EXAFS fitting was performed in the 2.35-7.7 Å distance range with the Artemis interface [62] to IFEFFIT using least-squares refinements. Paths used for fitting standards and samples were obtained from a metallic silver crystallographic model [63] using the FEFF6 algorithm included in the Artemis interface. Only paths with a rank higher than 7% were considered, and the E₀ value was set to 25520 eV. The amplitude reduction factor S₀² was determined to be equal to 0.978 by fitting the 1ˢᵗ coordination sphere of the Agfoil spectrum over a range of 2.30-2.83 Å. This value was used in all the fitting procedures. Degeneracy of the paths, energy shift ΔE₀, radial distance shift ΔR, and thermal and static disorder σ² were fitted for each of the selected paths for a total of 57 independent points and 19 variables. All R-factors were lower than 0.02.

The pellets prepared for XANES-EXAFS measurements were then analyzed by X-ray powder diffraction (XRD). The XRD diffractograms were recorded in 10 min on a Bruker D8 Discover diffractometer (France). Cu-Kα₁ radiation (Cu Kα₁, 1.5405 Å) produced in a sealed tube at 40 kV and 40 mA was selected and parallelized using a Gobël mirror parallel optics system and collimated to produce a 500-mm beam diameter. The data were collected in the 3°-70° 2θ range. XRD measurements were also performed for initial BCN suspensions with different surface modifications. The crystallinity index (CrI, %) was calculated as a function of the maximum intensity of the diffraction peak from the crystalline region (I₂₀₀) at 2θ = 22.5° and the minimum intensity from the amorphous region (Iₐₐₐ) at 2θ = 18°, according to the Segal equation [64]:

\[
    \text{CrI} = \frac{I_{200} - I_{am}}{I_{200}}
\]

The crystallite size (CS) was determined using Scherrer’s equation [65]:
where $K$ is the shape factor (0.9), $\lambda$ is the X-ray wavelength (1.54 Å), $\beta$ is the full-width at half-maximum (FWHM) and $\theta$ is the angle of the diffraction peak of the crystalline phase (Bragg’s angle). The FWHM was determined considering the characteristic peak at $2\theta = 22.5^\circ$ for initial BCN suspensions and the peak at $2\theta = 38^\circ$ for the AgNPs in hybrids.

The light-visible absorbance of hybrid suspensions was measured in the 300–800 nm range using a Mettler-Toledo UV7 (USA) spectrophotometer equipped with a 10-mm quartz cell. All the samples were water-diluted (1:10) and ultra-pure water was used as a blank reference.

To determine the real content of AgNPs in hybrid suspensions, a volume of 1 mL of sample was digested by 40 mL water/aqua regia mixture (i.e., 30% v aqua regia, HCl/HNO$_3$: 3/1). The resulting suspensions were then analyzed by atomic absorption spectroscopy, AAS (ICE 3300 AAS, Thermo Fisher Scientific, USA). A calibration curve was obtained by the measurement of a digested sample of silver standard solution (1000 $\mu$g/mL, Chem-Lab NV, Belgium) at different concentrations, from 0.5 to 10 ppm. Two independent measurements were repeated for each sample.

For scanning transmission electron microscope (STEM) observations, the as-synthesized suspensions were water-diluted at 0.5 g/L in BCN content and 10 $\mu$L were then deposited onto glow-discharged carbon coated grids (200 meshes, Delta Microscopies, France) for two minutes, removing the excess by touching the edge of the drop with Whatman filter paper. The grids were dried overnight in air and then coated with a platinum layer (thickness = 0.5 nm) by an ion-sputter coater (LEICA EM ACE600, Germany). Brightfield images were recorded using a field emission gun scanning electron microscope (Quattro S, Thermo Fischer Scientific, USA) operating at 10 kV with a STEM detector. For each sample, the acquired images were analyzed by ImageJ software. The AgNP Feret diameter (i.e., the largest distance between two tangents to the contour of the measured particle) was determined considering the largest possible number of AgNPs (from 35 to 100, depending on the sample).

3. Results

**Characteristics of surface-modified BCNs.** To elucidate the role of functional groups on the surface of the BCNs in AgNP nucleation, different surface modifications were performed. Firstly, a suspension of neutral BCNs was obtained from HCl acid hydrolysis. BCNs were then carboxylated via the TEMPO oxidation method (TBCN) obtaining a DO of 0.1 estimated from conductometric data. FTIR spectroscopy assessed the effective surface-modifications. As shown in Fig. 2a, the unmodified BCNs showed well-defined peaks at 1000 cm$^{-1}$ and 3000-3500 cm$^{-1}$, typical of –OH and –C–O–C– groups, respectively [66]. The TBCN spectrum displayed a strong absorption band between 1630 cm$^{-1}$ (carboxylate) and 1730 cm$^{-1}$ (acid), which was characteristic of the carboxyl groups in their acidic form, validating a successful TEMPO oxidation of BCNs [61]. TBCNs were further surface-modified to obtain two types of surface chemistry. Firstly, a choline chloride derivative (AETMA) was used...
to synthesize aminated cationic BCNs (ABCNs). For this sample, the FTIR spectrum revealed the presence of two new bands at around 1580 cm\(^{-1}\) assigned to the C=O stretching of the amide I band and to the N–H vibration of the amide II band, respectively [67]. Secondly, an octylamine C\(_8\) was covalently grafted onto the carboxyl group of TBCNs using NHS/EDC, leading to a hydrophobic BCN (HBCN). The good degree of coupling between TBCNs and C\(_8\) was proved by the very low surface charge density measured by conductometric titration (i.e., 0.010 mmol/g). The detection of these same bands in the spectrum of HBCNs and the presence of asymmetrical and symmetrical CH\(_2\) stretching from the C\(_8\) alkyl chain at 2850 and 2930 cm\(^{-1}\) indicated the successful octylamine grafting [43]. Nevertheless, a residual peak at 1730 cm\(^{-1}\) was still visible probably due to remaining free COOH groups. FTIR results were corroborated by the variation of the zeta potential (\(\zeta\)), implying that a modification of the electrical charge environment of the BCN was induced by the different chemical surface modifications. While pure BCNs were quasi-neutral, TBCNs showed a negative \(\zeta\) value of -25 mV, correlated to the variation of surface charge density (\(\rho\)). On the other hand, a clear increase in \(\zeta\) to +16 mV was observed for the aminated ABCNs due to the substitution of COO\(^-\) groups by the N\(^+\) of the AETMA molecules. The slightly negative value of -9 mV was measured for the alkyl hydrophobized HBCNs because of the grafting of octylamine onto the surface carboxyl groups. This low value indicated that HBCNs were highly substituted and no longer electrostatically but, instead, sterically stabilized. These experimental evidences were consistent with a stable suspension of highly-charged colloids for TBCNs and ABCNs with respect to BCNs.

The surface modifications affected the nanocrystal dispersions, as shown from STEM images of dry samples in Fig. 1. Even though the global morphology of the nanocrystals remained unchanged after modification, it could be observed that TBCNs and ABCNs were better dispersed in comparison to quasi-neutral unmodified BCNs and HBCNs. Furthermore, TBCN and ABCN suspensions were less opaque than those of pure BCNs and HBCNs (Fig. 1). Analysis of STEM images indicated an average length of between 850 and 1750 nm and a width of between 20 and 40 nm for all surface-modification treatments, which was in agreement with several studies in the literature [57,68,69]. All the outcomes are summarized in Table 1, indicating that the BCN surface was successfully modified for all of the cases considered.
Fig. 2. Characterization of unmodified BCNs and BCNs with TEMPO oxidation, aminated or hydrophobic surface modifications: (a) FTIR spectra; (b) XRD patterns.

Table 1. Crystalline and surface charge characteristics of native and modified BCNs

| Sample | Crystallinity, CrI (%)<sup>1</sup> | Crystallite size, CS (nm)<sup>1</sup> | Surface charge density, ρ (mmol/g)<sup>2</sup> | Zeta potential, ζ (mV) |
|--------|---------------------------------|---------------------------------|---------------------------------|----------------|
| BCN    | 59                              | 6.1                             | 0.006 ± 0.001                   | -11.2 ± 1.1   |
| TBCN   | 57                              | 5.8                             | 0.630 ± 0.003                   | -25.4 ± 0.8   |
| ABCN   | 62                              | 5.7                             | 0.540 ± 0.002                   | +15.9 ± 0.7   |
| HBCN   | 58                              | 6.1                             | 0.010 ± 0.002                   | -9.3 ± 1.9    |

- <sup>1</sup> by XRD; <sup>2</sup> by conductometric titration, two measurements were performed for each sample.

The XRD patterns of unmodified BCNs and surface-modified BCNs (Fig. 2b) showed 2θ diffraction peaks at 14.5°, 16.4°, 22.5° and 34.8°, related to (101), (010), (200) and (040) crystalline planes, respectively, according to the triclinic indexation of Nishiyama et al.[70]. As reported in Table 1, the crystallinity was unchanged around 60% after correction of the amorphous for all the samples, and the chemical treatment did not affect the size of crystallites, which was 5.9 ± 0.2 nm for all the samples, in agreement with the value found by Vasconcelos et al. [71]. These results are consistent with the fact that TEMPO oxidation and other post treatments do not affect the crystalline part of the samples [72,73].

**AgNP nucleation on BCNs with various surface modifications.** The nucleation of AgNP was investigated on the four BCN types of interest (i.e., native BCNs, TBCNs, ABCNs and HBCNs). Firstly, AgNO₃ aqueous solution was added to the BCN suspension and NaBH₄ aqueous solution was then introduced to reduce Ag⁺ into AgNPs. As soon as the silver precursor was reduced, the initial
translucent aqueous suspension turned light or dark yellow (inset, Fig. 3a), giving different shades of color, probably because of the initial dispersion state of the type of BCN used. However, such a color variation was not reflected in the UV-Vis spectra (Fig. 3a) since the effective AgNP content was very similar in all the hybrids (i.e., 7.9 wt% AgNP in BCN/AgNP; 6.8 wt% AgNP in TBCN/AgNP; 8.5 wt% AgNP in ABCN/AgNP; 8.6 wt% AgNP in HBCN/AgNP) with a same dominant in-plane absorption peak at $\lambda_{\text{max}} \approx 400$ nm. It confirmed the synthesis of well-dispersed AgNPs derived from the coalescence of monomeric Ag particles obtained by a reduction to a zero-valence Ag atom [20].

Even if the surface modification did not seem to affect the intensity of the spectra (i.e., similar AgNP content for all the hybrids), the full width of the main peak and the background intensity increased for HBCN/AgNP. This type of behavior is attributed to the synthesis of HBCN/AgNP in a H$_2$O/EtOH mixture since the AgNO$_3$ is not well-soluble in ethanol and could affect AgNP nucleation, leading to a larger distribution of AgNP sizes or heterogeneities. However, the use of EtOH was essential since it allowed the C$_8$ dissolution and helped hydrophobic nanocrystal dispersion after the synthesis, as was the presence of H$_2$O, which ensured the correct dissolution of silver precursor and, consequently, AgNP formation.

STEM images of dry BCN/AgNP, TBCN/AgNP, ABCN/AgNP and HBCN/AgNP hybrids are shown in Fig. 3b. The analysis of these images confirmed that all the AgNPs had the same particle size, which was thus not affected by the different surface modifications (i.e., BCN/AgNP = 17.1 ± 12 nm; TBCN/AgNP = 17.5 ± 12 nm; ABCN/AgNP = 16.1 ± 13 nm; HBCN/AgNP = 18.0 ± 12 nm). Complete size distributions are reported in Fig. S1. The nucleation growing method resulted in well-grafted AgNPs on the cellulose surface for the neutral and modified CNCs. In contrast, AgNPs synthesized in the same conditions without BCN rapidly aggregated. This proved that BCNs form a perfect substrate for the nucleation point to obtain well-dispersed quite-monodispersed AgNPs without the need for any other capping agents or stabilizers [74].

Nevertheless, these results suggested that AgNP nucleation on the cellulose nanocrystal surface non-specifically occurred on hydroxyl groups and/or on the additional negative surface charges. The aim was to determine if these surface charges represent an additional nucleation point that interacts with Ag$^+$ ions or if they just acted as promoters of nanocrystal dispersion, facilitating the accessibility to the hydroxyl surface groups. The TEMPO carboxylated TBCNs were then compared to aminated ABCNs. In this case, the functionalization by the positive surface charges preserved the electrostatic repulsion and, consequently, the good dispersion. At the same time, these positive charges could not interact with Ag$^+$ ions and thus did not represent a possible nucleation point for AgNPs. The STEM image of the ABCN/AgNP hybrid (Fig. 3b) showed that AgNPs were well-nucleated on both surfaces, indicating that the –OH surface groups represented the effective AgNP nucleation point. The OH groups and Ag$^+$ ions are complexed throught ion-dipole interactions [22] and the extensive number of hydroxyl groups on the BCN surface can promote the complexation of Ag$^+$ ions also acting as passivation contacts for AgNP stabilization [75].
To our knowledge, such a result represents the first experimental proof that hydroxyl surface groups on cellulosic surfaces are the real nucleation points for metallic nanoparticles and that additional negative surface charges just improve the dispersion state, thereby increasing the accessibility to the nucleation sites. This result was corroborated by the fact that for the HBCN/AgNP hybrid, most of the AgNPs were well-grafted onto the HBCN surface as well. In this case, the only real possible nucleation point was represented by –OH surface groups since most of the carboxylate groups were removed by grafting of C8 molecules. The presence of several AgNPs not anchored at the HBCN surface was visible in STEM images and could be linked to the suboptimal AgNO3 dissolution due to the presence of EtOH in the suspending medium.

Concerning the structural characterization, all the XRD patterns of BCN/AgNPs (Fig. 3c) showed characteristic peaks of crystalline silver, especially at 37.9° which is usually attributed to the (111) lattice plane of face-centered cubic (fcc) silver (JCPDS Card No. 89-3722). Other characteristic silver diffraction peaks at 46.2° and 64.4° related to other lattice planes (i.e., (200) and (220) crystalline planes) were detected with a weaker intensity. The presence of fcc crystal facets confirmed the isotropic nature of the crystals [20]. For all the samples, the same average AgNP diameter of 3 nm was estimated from XRD patterns. The nucleation mechanism of AgNPs could affect their growth on the nanocrystal surface and, thus, their final characteristics. Therefore, the oxidation state of AgNPs grafted onto BCN with different surface modifications (i.e., Ag0 and Ag+ contents) was determined by a linear combination fitting (LCF) of XANES spectra of the hybrids (Fig. S2a and Fig. S2b). For each sample, the R-factor and the Chi-square values of the fits are reported in Table S1. A quantity of 92 ± 9% of Ag0 was found for all the hybrids, showing the efficiency of the AgNP reduction and nucleation, independently of the surface treatment of the initial BCNs. All Fourier transform spectra of BCN/AgNP (Fig. S2c and Fig. S2d) were fitted with the crystallographic structure of metallic silver with an R-factor systematically lower than 0.015. The shifts in R values (i.e., interatomic distance) obtained from the fits were systematically negligible (< 0.06 Å⁻¹; all of the values are presented in Table S2), showing that the interatomic distances in BCN/AgNP case did not significantly change in comparison to the metallic silver distances and that the space group of BCN/AgNP sample still corresponded to the fcc silver structure, as suggested by XRD. It follows that the initial and final crystal structural organization of AgNPs were not affected by the various surface-modification treatments. These results showed the NaBH4 reduction with –OH surface groups as nucleation points represented an efficient nucleation mechanism where the speciation and the crystalline structure of AgNPs were not affected by the surface modifications performed on cellulose nanocrystals.


**AgNP redox post-treatment by H$_2$O$_2$.** H$_2$O$_2$ was defined as an efficient etching agent producing shape and structural modifications of AgNP into larger and more robust AgNPs (e.g., AgNPrisms) isolated or even attached to surface-carboxylated cellulose nanofibrils [20]. In this case, we proposed a more detailed study on the effect of the H$_2$O$_2$ redox post-treatment on properties of AgNPs (i.e., AgNP-H$_2$O$_2$) in the different BCN/AgNP hybrids. The outcomes reported in this section will support the understanding of the impact of the H$_2$O$_2$ action for BCN/AgNP where BCN was surface-modified. Even if all the prepared samples were able to undergo the same H$_2$O$_2$-induced structural change, the TBCN/AgNP hybrid system was chosen as a case study since the TBCNs led to good dispersion. Such a TBCN/AgNP hybrid suspension at constant initial AgNP content (i.e., 7 wt%) was mixed with different amounts of H$_2$O$_2$ (40 – 250 μL), thus obtaining a variation of the H$_2$O$_2$/AgNP mass ratio, α, from 0 to 0.52, low enough to balance the H$_2$O$_2$ oxidative activity and its reduction capability. With the addition of H$_2$O$_2$, the color solution drastically changed, ranging from orange to blue (inset, Fig. 3).

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**Fig. 3.** Characterizations of BCN-, TBCN-, ABCN-, HBCN-/AgNP hybrid suspensions: (a) UV-Vis spectra; inset: sample pictures; (b) STEM images; (c) XRD patterns.
The color change was associated with a modification of the UV-Vis spectra of the H₂O₂-containing suspensions with respect to the reference one (i.e., 0 μL H₂O₂), as shown in Fig. 4a. The addition of H₂O₂ induced a redshift of the in-plane dipole surface plasmon resonance peak at 400 nm. This was already visible at α equal to 0.17, since the absorption peak became broader and shifted to 495 nm, showing the presence of two shoulders at 345 nm and 385 nm related to an early modification of primary AgNPs. At α = 0.25, the in-plane plasmon peak was further shifted to 638 nm and finally moved out of the measurement window for higher α values. In this case, a lower intensity peak was detected at λ_max = 335 nm that is usually associated with an out-of-plane quadrupole resonance peak, thus representing a good indicator for general prismatic architectures since it strongly depends on aspect ratio [48]. Such a peak became sharper for α values of 0.33 and 0.52 (i.e., 160 μL and 250 μL of H₂O₂, respectively). The presence of a low-intensity peak at 335 nm, at the same time as the shift of the in-plane resonance peak out of the measurement window, and the absence of the primary AgNP peak is proof that an AgNP size-shape variation is induced, as already observed in literature [20]. To better describe the impact of the H₂O₂ redox post-treatment on the AgNP morphology STEM acquisition were performed (Fig. 4b). The analysis of STEM images indicated that AgNPs initially had an average diameter of 17.5 ± 12 nm that regularly increased to reach 94.1 ± 69 nm for α equal to 0.52, where several AgNPs_H₂O₂ with quite irregular edges and a sometimes vaguely triangular shape could be observed. These results agreed with the experimental evidence reported in a recent study of our group [76], in which we investigated the effect of the H₂O₂ redox treatment in hybrid suspensions where primary 10-nm AgNPs are nucleated on wood cellulose NCs, proving the efficient size-shape transition from 10-nm spherical AgNPs into 300 nm AgNPrisms. In this work, we proposed a H₂O₂ redox mechanism where H₂O₂ induced the oxidative dissolution of primary AgNPs, generating Ag⁺ ions. When α equal to or greater than 0.20, the H₂O₂ oxidation interest most of the AgNPs, except the contact location where AgNPs are effectively grafted onto the NC surface which could actually work as nucleation sites for the formation of newly formed AgNPs with a triangular shape. Finally, all the images clearly showed that AgNPs_H₂O₂, even the bigger ones, were still attached to the TBCN surface, exactly as for the reference sample before the H₂O₂ post-treatment, showing the importance of cellulose as a support for AgNPs_H₂O₂ nucleation, growth and stabilization. The formation of non-perfectly prismatic silver objects and the increase in the average diameter associated with an increase in the polydispersity in size (Fig. S3a and Fig. S3b, Table S3) could be linked to the absence of any additional stabilizer, which implied that the structural change was heavily affected by the aggregation of newly-formed AgNPs_H₂O₂ [46].

To establish how the H₂O₂ post-treatment affected the oxidation state of AgNPs, XANES spectra were recorded and then fitted using the LCF procedure (Fig. S4a and Table S4). Data showed that the amount of Ag⁰ in AgNPs_H₂O₂ slightly decreased with respect to the reference case, from 88% before H₂O₂ post-treatment down to 69% at α = 0.08, as long as the H₂O₂ addition did not induce a size-shape
variation (corresponding to $\alpha$ from 0.08 to 0.25), while remaining equal to 91% and 87% for $\alpha$ values of 0.33 and 0.52, respectively (Table S4). The formation of AgNPs$_{H_2O_2}$ mostly composed of Ag$_0$ silver was also corroborated by the presence in the XRD patterns of characteristic (111), (200) and (220) Ag$_0$ peaks, confirming the presence of a lattice plane of a face-centered cubic structure (Fig. S4b). It could be observed that the H$_2$O$_2$ post-treatment did not induce a structural modification of AgNPs$_{H_2O_2}$ with respect to the primary AgNPs before the H$_2$O$_2$ treatment and that a H$_2$O$_2$/AgNP mass ratio equal to 0.33 and 0.52 determined an increase in the intensity of the characteristic fcc Ag (111) peak, which corresponded to the highest recorded values of Ag$_0$. Such a result suggested that a $\alpha$ parameter at least equal to 0.33 represented the optimum value necessary to reach an effective size-shape variation that maintains the conversion of the initial Ag$^+$ to Ag$_0$ ratio. Finally, the EXAFS Fourier transform spectra (Fig. S4c and Fig. S4d) of the AgNP$_{H_2O_2}$ in hybrids were fitted with the crystallographic structure of metallic silver, (R-factor systematically lower than 0.020), presenting systematically negligible shifts in R space (< 0.06 Å$^{-1}$; all values are in Table S5). This result confirmed that the interatomic distances in AgNP$_{H_2O_2}$ did not significantly change in comparison to the metallic silver distances and that the space group of the samples still corresponded to the fcc silver structure, as observed by XRD. The final crystal structural organization was not affected by the H$_2$O$_2$ redox post-reaction nor by the particle size variation, as observed by Ma et al. [77] Conversely, the AgNP diameter measured by XRD was about 10 nm, which was consistently smaller than the 40-80 nm measured by STEM. Such a difference could be explained by the fact that XRD makes it possible to evaluate the size of the coherent diffraction domains in AgNPs, i.e., perfect repetition in single-crystal particles, twinned or imperfect particles that have more than a single diffraction domain [77]. The characteristics of all the investigated samples are reported in Table 2.

All these results, coupled with the experimental proof that -OH groups on the CNC surface are the AgNP nucleation points, allowed us to conclude that our approach to tune the morphology and the oxidation state of the AgNP grafted onto cellulosic support may be extended to all the hybrid systems where metallic NPs are anchored on a polysaccharide substrate (e.g., cellulose fibers, films or chitin NCs).
Fig. 4. (a) UV-Vis spectra and (b) STEM images of TBCN/AgNP hybrid suspension mixed with different volumes of H$_2$O$_2$ and, thus, at various $\alpha$ values. (c) Ag$_0$ content of AgNP-H$_2$O$_2$ by XANES fitting, and average AgNP-H$_2$O$_2$ diameter determined by STEM as a function of the $\alpha$ parameter in TBCN/AgNP-H$_2$O$_2$ hybrids. For Ag$_0$ value, standard error as considered as the 10% of the measured value.

Table 2. Summary of characteristics of BCN/AgNP hybrids with and without H$_2$O$_2$ redox post-treatment.

| Sample     | H$_2$O$_2$/AgNP mass ratio ($\alpha$) | Avg diam. (nm)$^1$ | Ag$_0$ (%)$^2$ |
|------------|--------------------------------------|--------------------|---------------|
| BCN/AgNP   | 0                                    | 17.1 ± 12          | 90 ± 9        |
Impact of BCN surface modification on H\textsubscript{2}O\textsubscript{2} redox post-treatment. Hybrids prepared with various BCN types were submitted to the H\textsubscript{2}O\textsubscript{2} redox post-treatment, keeping the H\textsubscript{2}O\textsubscript{2}/AgNP ratio, $\alpha$, equal to 0.33, the value at which an effective size-shape modification of the initial AgNPs was found in the case of TBCN/AgNP (Table 2). In the case of aminated surfaces, the color variation of the ABCN/AgNP\textsubscript{H\textsubscript{2}O\textsubscript{2}} from yellow to blue was observed after the addition of H\textsubscript{2}O\textsubscript{2}, as observed for the TBCN/AgNP\textsubscript{H\textsubscript{2}O\textsubscript{2}} (Fig. 5a). This suspension showed an in-plane absorption peak shifted to higher $\lambda_{\text{max}}$ (i.e., out of the measurement window), and a sharp out-of-plane quadrupole resonance peak was visible at 333 nm (Fig. 5b). This result agreed with the transformation of the primary AgNPs of 16.1 ± 13 nm into bigger AgNPs\textsubscript{H\textsubscript{2}O\textsubscript{2}} of 95.4 ± 87 nm, as confirmed by STEM (Fig. 5c). On the other hand, the addition of H\textsubscript{2}O\textsubscript{2} to the BCN/AgNP turned the suspension a darker gray color, and the particle size slightly changed with respect to the reference case (i.e., 32.5 ± 19 nm). Moreover, the $\lambda_{\text{max}}$ in the UV-Vis spectra remained at 402 nm and a poorly-defined low-intensity broad peak appeared at 340 nm. These results indicated an uncomplete transformation of the AgNPs, suggesting that the BCN surface modification affected the H\textsubscript{2}O\textsubscript{2} transformation of primary AgNPs since the BCN surface modification influenced the BCN dispersion rather than its chemical structure. When the surface charges provided a good repulsion between nanocrystals (the cases of TBCN and ABCN), the addition of H\textsubscript{2}O\textsubscript{2} made it possible to obtain the formation of AgNPs\textsubscript{H\textsubscript{2}O\textsubscript{2}} of about 100 nm, whereas the size of AgNPs\textsubscript{H\textsubscript{2}O\textsubscript{2}} for the almost neutral surface charge in the BCN case remained constant (i.e., about 20 nm). It appeared that well-dispersed nanocrystals facilitate the accessibility to the surface groups, thereby promoting the H\textsubscript{2}O\textsubscript{2} redox action.

Concerning the hydrophobically-modified surface (i.e., HBCN/AgNP case), the introduction of H\textsubscript{2}O\textsubscript{2} did not induce a size-shape modification of primary AgNPs, as shown in the STEM image (Fig. 5c).
Furthermore, the UV-Vis spectrum of the HBCN/AgNP\textsubscript{H\textsubscript{2}O\textsubscript{2}} suspension did not change with respect to the reference HBCN/AgNP. In this case, not only the absence of repulsion prevented good nanocrystal dispersion, but the suspending medium containing EtOH could promote aggregation and affect the efficiency of the H\textsubscript{2}O\textsubscript{2} redox reaction as well. To check the influence of EtOH on the H\textsubscript{2}O\textsubscript{2} treatment, we compared aqueous TBCN/AgNPs suspension (highly charged nanocrystals to ensure dispersion) to the same TBCN/AgNPs diluted in a H\textsubscript{2}O/EtOH (60/40 v/v) mixture, like in the case of the HBCN/AgNPs. In pure water, the addition of H\textsubscript{2}O\textsubscript{2} induced a color variation from yellow to blue, whereas the hybrid prepared in the H\textsubscript{2}O/EtOH mixture just turned a dark yellow (Fig. S5). This result confirmed that the presence of ethanol limited the formation of AgNPs\textsubscript{H\textsubscript{2}O\textsubscript{2}} that could be linked to a weak oxidation of AgNPs to Ag\textsuperscript{+} and the subsequent reduction in a non-aqueous medium. Size distributions of AgNPs\textsubscript{H\textsubscript{2}O\textsubscript{2}} in BCN hybrids are reported in Fig. S6 and the average diameters are summarized in Table S6.

Interestingly, all the reformed AgNPs\textsubscript{H\textsubscript{2}O\textsubscript{2}} appeared to be nucleated on nanocrystal surfaces, irrespective of the applied surface modification. This means that the BCN still serves as a substrate for AgNPs\textsubscript{H\textsubscript{2}O\textsubscript{2}} formation, ensuring adequate AgNP dispersion. We hypothesized that the H\textsubscript{2}O\textsubscript{2} did not completely oxidize the AgNPs, leaving hooks available as a large amount of re-nucleation points for AgNP\textsubscript{H\textsubscript{2}O\textsubscript{2}} nanoparticles in addition to the \textendash OH surface groups, as reported in another work of our group [76].
Fig. 5. (a) Color variation of hybrid suspensions; (b) UV-Vis spectra; (c) STEM images; (d) XRD patterns of BCN-, TBCN-, ABCN-, HBCN-/AgNP-H$_2$O$_2$ hybrid suspensions with the addition of 160 $\mu$L of H$_2$O$_2$ ($\alpha = 0.33$).

Moreover, the analysis of XANES spectra (Fig. S7a, Table S7) showed that the addition of H$_2$O$_2$ did not affect the Ag$_{0}$ content (i.e., 95 ± 10% compared to 92 ± 9% of the reference cases), indicating that the oxidation state did not change. Finally, the XRD patterns of hybrid suspensions without ($\alpha = 0$) and with the addition of H$_2$O$_2$ ($\alpha = 0.33$), Fig. 5d., indicating the presence of the Ag (111), Ag (200) and Ag (220) peaks, characteristic of the fcc silver model. This confirmed the formation of metallic AgNPs-H$_2$O$_2$ well-grafted onto the BCN substrate, irrespective of the surface modification. Moreover, the fitting of EXAFS Fourier transform spectra (R-factor < 0.016) did not reveal any significant modification of the crystallographic structure in comparison to fcc metallic silver (Fig. S8a and Fig.
S8b, data in Table S8). These studies confirmed that all native and modified BCN/AgNP samples were composed quasi-exclusively of Ag$_0$ with similar crystallographic organization.

4. Conclusion

In this study, we investigated the interactions involving well-dispersed AgNPs grafted on a bio-based substrate, focusing on the role of polysaccharide surface chemistry. Working with bacterial cellulose nanocrystals (BCNs) as model substrate, we experimentally showed that the hydroxyl groups serve as nucleation points for AgNPs through ion-dipole interaction and that the surface charges only promoted nanocrystal dispersion, improving accessibility to the OH groups. To our knowledge, this is the first experimental identification of the effective nucleation point for metallic nanoparticles on a solid cellulose surface. This was investigated modifying native quasi-neutral BCNs dispersed in aqueous suspension into negatively or positively charged, and hydrophobic BCNs then used to prepare hybrid nanomaterials (i.e., BCN/AgNP, TBCN/AgNP, ABCN/AgNP, HBCN/AgNP) where highly controlled AgNPs are anchored on the BCN surface. This work proved that no surface modification of cellulose (e.g., TEMPO oxidation [32]) is necessary to efficiently nucleate and anchor AgNPs on their surface as long as the reduction step is carried out. Furthermore, we prove that the grafting of hydrophobic molecules on BCNs does not prevent the nucleation and the growth of AgNPs, thus opening the way to the formulation of bifunctional nanoparticles in non-aqueous media.

The investigated hybrid suspensions were subjected to an H$_2$O$_2$ redox post-treatment. An optimal H$_2$O$_2$/AgNP mass ratio ($\alpha = 0.33$) was defined at which a size-shape transition of AgNPs, from 10-nm spherical NPs to 100-nm NPs with a triangular shape, while maintaining the initial fcc crystal structure and increasing the Ag$_0$ content in AgNPs. Here, we showed that the dispersion of the BCNs improved the efficiency of the H$_2$O$_2$ post-treatment, with an increase in AgNP size detected only in presence of repulsive surface charges (i.e., TBCNs and ABCNs).

As a result, this highly controlled synthesis of AgNPs well-grafted onto the cellulose NC surface can promote the development and design of new bio-based hybrid materials, such as sensors and biological imaging, catalysts, conductive material. Furthermore, a controlled release of Ag$^+$ ions from well-characterized AgNPs would allow producing bio-based nanomaterials for biocidal properties that can be used for several applications, such as food packaging, paints or surface treatment.

CRediT authorship contribution statement

D. Musino: Conceptualization, methodology, investigation, formal analysis, visualization, data curation, writing – original draft. C. Rivard: Formal analysis, investigation, writing – review and editing. G. Landrot: Formal analysis, investigation, writing – review and editing. B. Novales: Investigation, writing – review and editing. T. Rabilloud: Supervision, investigation, writing – review and editing. I. Capron: Conceptualization, investigation, supervision, project administration, writing - review & editing.
Declaration of competing interest

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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Appendix A. Supplementary data

Supplementary data to this article can be found online at: 10.1016/x0xx00000x

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Table 1. Crystalline and surface charge characteristics of native and modified BCNs

Table 2. Summary of characteristics of BCN/AgNP hybrids with and without H₂O₂ redox post-treatment.
Electronic Supplementary Information

Hydroxyl Groups on Cellulose Nanocrystal Surfaces form Nucleation Points for Silver Nanoparticles of Varying Shapes and Sizes

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Fig. S1. Size distribution histograms of AgNPs nucleated on BCNs, TBCNs, ABCNs and HBCNs. A total of 100 AgNPs were counted for each sample.
Fig. S2. (a) Example of XANES spectrum fitted by linear combination (LCF) analysis using Agfoil and AgNO$_3$ aqueous solutions as standards; (b) XANES data; (c) EXAFS spectra Fourier transform (solid gray lines) and fit (dotted lines); (d) magnitude and imaginary part (solid gray lines) and fit (dotted lines) of the Fourier transform of BCN-, TBCN-, ABCN- and HBCN/AgNP hybrids.
### Table S1. R-factor and Chi-square values for linear combination fitting procedure applied to the XANES region of BCN-, TBCN-, ABCN- and HBCN-/AgNP hybrids.

| Hybrid sample | R-factor  | Chi-square | Ag\(_0\) (%)\(^1\) |
|---------------|-----------|------------|---------------------|
| BCN/AgNP      | 0.0003215 | 0.01114    | 90 ± 9              |
| TBCN/AgNP     | 0.0004912 | 0.01317    | 88 ± 9              |
| ABCN/AgNP     | 0.0003987 | 0.01086    | 94 ± 9              |
| HBCN/AgNP     | 0.0002244 | 0.00613    | 97 ± 10             |

\(^1\) by XANES, the standard error as 10% of the measured value.
Table S2. EXAFS fit results for BCN-, TBCN-, ABCN- and HBCN-/AgNP hybrids.

| Degeneracy of the paths | Interatomic distance R (Å) | Variation in interatomic distance ΔR (Å) | Debye-Waller factor σ² |
|--------------------------|-----------------------------|----------------------------------------|------------------------|
| BCN/AgNP                 | TBCN/AgNP                   | ABCN/AgNP                              | HBCN/AgNP              |
| Ag1 ss                   | 9.5 ± 0.5                   | 10.4 ± 0.5                             | 10.9 ± 0.6             | 0.003                   | -0.029 | -0.030 | -0.028 | 2.876 |
| Ag2 ss                   | 2.5 ± 1.1                   | 3.9 ± 1.5                              | 4.4 ± 1.7              | 0.003                   | -0.033 | -0.037 | -0.038 | 4.076 |
| Ag3 ss                   | 25.7 ± 3.7                  | 25.9 ± 3.6                             | 26.4 ± 3.7             | 0.006                   | -0.035 | -0.036 | -0.035 | 4.997 |
| Ag4 ss                   | 6.1 ± 2.3                   | 2.7 ± 4.8                              | 5.5 ± 2.5              | 0.003                   | -0.010 | -0.021 | -0.012 | 5.801 |
| Ag5 ss                   | 10.4 ± 6.3                  | 8.4 ± 6.2                              | 11.6 ± 6.4             | 0.004                   | -0.056 | -0.057 | -0.057 | 6.439 |
| Ag7 ss                   | 38.8 ± 8.7                  | 36.9 ± 7.8                             | 43.2 ± 10.6            | 0.003                   | -0.053 | -0.057 | -0.057 | 7.628 |
| Ag1 Ag7 ot               | 96*                         | 96*                                    | 96*                    | 0.003                   | -0.028 | -0.027 | -0.027 | 7.784 |
| Ag3 Ag7 ot               | 96*                         | 96*                                    | 96*                    | 0.004                   | -0.035 | -0.034 | -0.035 | 7.777 |

BCN/AgNP  TBCN/AgNP  ABCN/AgNP  HBCN/AgNP
| R-factor   | 0.015 | 0.014 | 0.012 | 0.015 |
|-----------|-------|-------|-------|-------|
| ΔE₀        | 0.89 ± 0.58 | 0.69 ± 0.54 | 0.49 ± 0.26 | 0.47 ± 0.60 |

ss: single scattering; at: acute triangle; ot: obtuse triangle; fs: forward scattering; dfs: double forward scattering; fta: forward through absorber. Fixed parameters are indicated by a "***". The amplitude reduction factor S₀² was fixed at 0.978 Å. Errors obtained for σ² were systematically lower than 0.0027; errors obtained for ΔR and R were systematically lower than 0.0150.
**Fig. S3.** Size distribution histograms of TBCN/AgNP-H$_2$O$_2$ hybrids with various volumes of H$_2$O$_2$ added and, consequently, various H$_2$O$_2$/AgNP mass ratios. $\alpha$ from 0 to 0.17 (a), and from 0.25 to 0.52 (b).

**Table S3.** Average diameter of AgNPs-H$_2$O$_2$ in TBCN/AgNP-H$_2$O$_2$ hybrid at 7 wt% AgNP treated with different volumes of H$_2$O$_2$ (i.e., various H$_2$O$_2$/AgNP mass ratios, $\alpha$).

| H$_2$O$_2$/AgNP mass ratio ($\alpha$) | Avg AgNP-H$_2$O$_2$ diam. (nm) | AgNP count |
|-------------------------------------|---------------------------------|-------------|
| $\alpha 0$                          | 17.5 ± 12                       | 100         |
| $\alpha 0.08$                       | 21.6 ± 12                       | 100         |
| $\alpha 0.17$                       | 32.5 ± 19                       | 100         |
| $\alpha 0.25$                       | 42.7 ± 28                       | 100         |
| $\alpha 0.33$                       | 97.5 ± 65                       | 80          |
| $\alpha 0.52$                       | 94.1 ± 69                       | 35          |
Fig. S4. (a) XANES spectra; (b) XRD diffractograms; (c) EXAFS spectra Fourier transform (solid gray lines) and fit (dotted lines); (d) magnitude and imaginary part (solid gray lines) and fit (dotted lines) of the Fourier transform of TBCN/AgNP-H$_2$O$_2$ hybrids at 7 wt% with various volumes of H$_2$O$_2$ added (i.e., various H$_2$O$_2$/AgNP mass ratios, $\alpha$).
Table S4. R-factor and Chi-square values for TBCN/AgNP$_2$O$_2$ hybrids at various H$_2$O$_2$/AgNP mass ratios.

| H$_2$O$_2$/AgNP mass ratio ($\alpha$) | R-factor | Chi - square | Ag$_0$ (%) |
|--------------------------------------|----------|--------------|------------|
| $\alpha$0                            | 0.0004912| 0.01317      | 88 ± 9     |
| $\alpha$0.08                         | 0.0006215| 0.01679      | 79 ± 8     |
| $\alpha$0.17                         | 0.0005848| 0.01588      | 79 ± 8     |
| $\alpha$0.25                         | 0.0005803| 0.01580      | 69 ± 7     |
| $\alpha$0.33                         | 0.0003492| 0.00947      | 91 ± 9     |
| $\alpha$0.52                         | 0.0005803| 0.01236      | 87 ± 9     |

1 by XANES, the standard error as 10% of the measured value.
Table S5. EXAFS fit results for TBCN/AgNP$_2$H$_2$O$_2$ hybrid suspensions at various H$_2$O$_2$/AgNP ratios, $\alpha$.

| Path    | $\delta_0$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ | $\alpha$ |
|---------|------------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|
| Ag1 ss  | 9.9 ± 0.5  | 8.4 ± 0.5| 8.5 ± 0.4| 7.3 ± 0.6| 10.4     | 10.0     | 0.004    | 0.003    | 0.003    | 0.003    | 0.003    | 0.003    | -        | -        | -        | -        | -        | -        | -        | -        | -        | -        | -        | 2.876    | 2.877    | 2.876    | 2.876    |
| Ag2 ss  | 2.3 ± 1.0  | 2.3 ± 1.1| 2.4 ± 1.2| 3.4 ± 1.7| 3.4 ± 1.7| 0.003    | 0.003    | 0.003    | 0.003    | 0.003    | 0.003    | -        | -        | -        | -        | -        | -        | -        | -        | -        | -        | -        | 4.072    | 4.077    | 4.077    | 4.078    |
| Ag1 Ag1 at | 48*       | 48*      | 48*      | 48*      | 48*      | 0.005    | 0.005    | 0.005    | 0.005    | 0.005    | 0.005    | -        | -        | -        | -        | -        | -        | -        | -        | -        | -        | -        | 4.314    | 4.316    | 4.315    | 4.314    | 4.316    |
| Ag3 ss  | 25.9      | 23.7     | 23.3     | 21.9     | 26.3     | 26.1     | 0.006    | 0.005    | 0.005    | 0.005    | 0.005    | 0.005    | -        | -        | -        | -        | -        | -        | -        | -        | -        | -        | 4.998    | 4.998    | 4.997    | 4.998    | 4.999    |
| Ag1 Ag3 ot | 96*       | 96*      | 96*      | 96*      | 96*      | 0.005    | 0.004    | 0.004    | 0.004    | 0.004    | 0.004    | -        | -        | -        | -        | -        | -        | -        | -        | -        | -        | 5.398    | 5.399    | 5.398    | 5.398    | 5.400    | 5.399    |
| Ag4 ss  | 4.3 ± 3.6  | 7.0 ± 3.6| 12.8     | 15.2     | 7.1 ± 2.2| 7.1 ± 2.2| 0.004    | 0.003    | 0.002    | 0.003    | 0.002    | 0.002    | -        | -        | -        | -        | -        | -        | -        | -        | -        | 5.795    | 5.804    | 5.830    | 5.840    | 5.807    | 5.806    |
| Ag1 Ag4 fs | 24*       | 24*      | 24*      | 24*      | 24*      | 0.008    | 0.006    | 0.005    | 0.006    | 0.005    | 0.005    | -        | -        | -        | -        | -        | -        | -        | -        | -        | -        | 5.766    | 5.776    | 5.801    | 5.810    | 5.780    | 5.777    |
| Ag1 Ag1 fta | 12*       | 12*      | 12*      | 12*      | 12*      | 0.014    | 0.012    | 0.012    | 0.013    | 0.012    | 0.012    | -        | -        | -        | -        | -        | -        | -        | -        | -        | -        | 5.752    | 5.755    | 5.753    | 5.752    | 5.756    | 5.755    |
| Ag1 Ag4 Ag1 | 12*       | 12*      | 12*      | 12*      | 12*      | 0.008    | 0.006    | 0.005    | 0.006    | 0.005    | 0.005    | -        | -        | -        | -        | -        | -        | -        | -        | -        | -        | 5.766    | 5.776    | 5.801    | 5.810    | 5.780    | 5.777    |
ss: single scattering; at: acute triangle; ot: obtuse triangle; fs: forward scattering; dfs: double forward scattering; fta: forward through absorber. Fixed parameters are indicated by a “*”. The amplitude reduction factor $S_0^2$ was fixed at 0.978 Å. Errors obtained for $\sigma^2$ were systematically lower than 0.0018; errors obtained for $\Delta R$ and $R$ were systematically lower than 0.0096.
Fig. S5. Comparison between TBCN/AgNP prepared with the addition of 160 μL of H₂O₂ (α = 0.33) in a pure water medium or in a H₂O/EtOH mixture. The presence of ethanol seems to affect the H₂O₂ post-treatment since the color suspension did not turn blue like the pure water suspension.

Table S6. Average diameter of AgNPs H₂O₂ in BCN-. TBCN-. ABCN. HBCN/AgNP H₂O₂ hybrids at H₂O₂/AgNP ratios. α, equal to 0.33.

| Hybrid sample + 160 μL H₂O₂ (α= 0.33) | Avg AgNP H₂O₂ diam. (nm) | AgNP count |
|---------------------------------------|---------------------------|------------|
| BCN/AgNP H₂O₂                         | 32.5 ± 19                 | 100        |
| TBCN/AgNP H₂O₂                        | 97.5 ± 65                 | 80         |
| ABCN/AgNP H₂O₂                        | 95.4 ± 87                 | 100        |
| HBCN/AgNP H₂O₂                        | 29.4 ± 22                 | 100        |
**Fig. S7.** XANES spectra of BCN−, TBCN−, ABCN−, and HBCN/AgNP-H2O2 hybrids mixed with 160 μL of H2O2 (α = 0.33).

**Table S7.** R-factor and Chi-square values for BCN−, TBCN−, ABCN− and HBCN/AgNP-H2O2 hybrid suspensions mixed with 160 μL of H2O2 (α = 0.33).

| Hybrid sample + 160 μL H2O2 (α = 0.33) | R-factor   | Chi-square | Ag0 (%) |
|----------------------------------------|------------|------------|----------|
| BCN/AgNP-H2O2                          | 0.0002663  | 0.00783    | 93 ± 9   |
| TBCN/AgNP-H2O2                         | 0.0003492  | 0.00947    | 91 ± 9   |
| ABCN/AgNP-H2O2                         | 0.0001964  | 0.00538    | 97 ± 10  |
| HBCN/AgNP-H2O2                         | 0.000187   | 0.00509    | 97 ± 10  |

1 by XANES, the standard error as 10% of the measured value.
Fig. S8. (a) EXAFS spectra Fourier transform (solid gray lines) and fit (dotted lines); (d) magnitude and imaginary part (solid gray lines) and fit (dotted lines) of the Fourier transform of BCN-, TBCN-, ABCN-, and HBCN/AgNP\textsubscript{H\textsubscript{2}O\textsubscript{2}} hybrids with 160μL of H\textsubscript{2}O\textsubscript{2} (α = 0.33).
Table S8. EXAFS fit results for BCN-, TBCN-, ABCN- and HBCN- AgNP \( \text{H}_2\text{O}_2 \) hybrid suspensions with 160 \( \mu\text{L} \) of \( \text{H}_2\text{O}_2 \) (\( \alpha = 0.33 \)).

| Degeneracy of the paths | BCN | TBCN | ABCN | HBCN | BCN | TBCN | ABCN | HBCN | BCN | TBCN | ABCN | HBCN | BCN | TBCN | ABCN | HBCN |
|-------------------------|-----|------|------|------|-----|------|------|------|-----|------|------|------|-----|------|------|------|
| Ag1 ss                   | 10.2 ± 0.6 | 10.4 ± 0.5 | 11.5 ± 0.6 | 11.0 ± 0.6 | 0.0030 | 0.0030 | 0.0031 | 0.0030 | -0.027 | -0.027 | -0.028 | -0.030 | 2.879 | 2.878 | 2.878 | 2.876 |
| Ag2 ss                   | 3.3 ±1.5 | 3.4 ± 1.4 | 4.7 ±1.8 | 4.7 ± 1.8 | 0.0032 | 0.0032 | 0.0038 | 0.0038 | -0.032 | -0.034 | -0.035 | -0.038 | 4.077 | 4.075 | 4.074 | 4.071 |
| Ag1 Ag1 at               | 48* | 48* | 48* | 48* | 0.0045 | 0.0045 | 0.0047 | 0.0044 | -0.040 | -0.041 | -0.041 | -0.045 | 4.318 | 4.317 | 4.317 | 4.313 |
| Ag3 ss                   | 27.1 ± 3.9 | 26.3 ± 3.6 | 27.9 ± 4.0 | 26.5 ± 3.7 | 0.0049 | 0.0048 | 0.0049 | 0.0046 | -0.034 | -0.034 | -0.034 | -0.038 | 4.999 | 4.999 | 4.998 | 4.995 |
| Ag1 Ag3 ot               | 96* | 96* | 96* | 96* | 0.0040 | 0.0039 | 0.0040 | 0.0038 | -0.022 | -0.022 | -0.022 | -0.024 | 5.400 | 5.400 | 5.399 | 5.397 |
| Ag4 ss                   | 5.6 ± 2.5 | 7.1 ± 1.8 | 2.8 ± 4.6 | 12.9 ± 1.4 | 0.0023 | 0.0019 | 0.0032 | 0.0015 | -0.009 | -0.004 | -0.017 | 0.016 | 5.802 | 5.807 | 5.794 | 5.827 |
| Ag1 Ag4 fs               | 24* | 24* | 24* | 24* | 0.0053 | 0.0049 | 0.0063 | 0.0044 | -0.036 | -0.031 | -0.044 | -0.014 | 5.775 | 5.780 | 5.767 | 5.797 |
| Ag1 Ag1 fta              | 12* | 12* | 12* | 12* | 0.0120 | 0.0120 | 0.0125 | 0.0118 | -0.054 | -0.055 | -0.055 | -0.060 | 5.757 | 5.756 | 5.756 | 5.751 |
| Ag1 Ag4 Ag1 dfs          | 12* | 12* | 12* | 12* | 0.0053 | 0.0049 | 0.0063 | 0.0044 | -0.036 | -0.031 | -0.044 | -0.014 | 5.775 | 5.780 | 5.767 | 5.797 |
| Ag5 ss                   | 13.0 ± 6.9 | 12.8 ± 6.6 | 12.3 ± 7.6 | 10.7 ± 5.4 | 0.0038 | 0.0038 | 0.0041 | 0.0029 | -0.052 | -0.055 | -0.057 | -0.060 | 6.445 | 6.442 | 6.440 | 6.437 |
| Ag7 ss                   | 38.0 ± 8.7 | 38.9 ± 8.9 | 34.9 ± 7.5 | 44.7 ± 10.7 | 0.0025 | 0.0027 | 0.0022 | 0.0029 | -0.048 | -0.048 | -0.051 | -0.052 | 7.639 | 7.639 | 7.636 | 7.635 |
| Ag1 Ag7 ot               | 96* | 96* | 96* | 96* | 0.0028 | 0.0028 | 0.0027 | 0.0029 | -0.025 | -0.026 | -0.026 | -0.028 | 7.787 | 7.787 | 7.787 | 7.784 |
| Ag3 Ag7 ot               | 96* | 96* | 96* | 96* | 0.0037 | 0.0037 | 0.0036 | 0.0037 | -0.033 | -0.033 | -0.034 | -0.036 | 7.780 | 7.780 | 7.779 | 7.776 |

| R-factor | 0.016 | 0.015 | 0.015 | 0.015 |
| ΔE_0     | 0.99 ± 0.62 | 0.52 ± 0.60 | 0.49 ± 0.60 | 0.21 ± 0.60 |
ss: single scattering; at: acute triangle; ot: obtuse triangle; fs: forward scattering. dfs: double forward scattering; fta: forward through absorber. Fixed parameters are indicated by a "*". The amplitude reduction factor $S_0^2$ was fixed at 0.978 Å. Errors obtained for $\sigma^2$ were systematically lower than 0.0018; errors obtained for $\Delta R$ and $R$ were systematically lower than 0.0097.