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Published in:
ACS Sustainable Chemistry and Engineering

DOI:
10.1021/acssuschemeng.9b04005

Published: 16/08/2019

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

Please cite the original version:
Sustainable High Yield Route to Cellulose Nanocrystals from Bacterial Cellulose

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Supporting Information

ABSTRACT: HCl gas hydrolysis of a bacterial cellulose (BC) aerogel followed by 2,2,6,6-tetramethylpiperidine-1-oxyl radical-mediated oxidation was used to produce hydrolyzed BC with carboxylate groups, which subsequently disintegrated into a stable dispersion of cellulose nanocrystals (CNCs). The degree of polymerization was successfully reduced from 2160 to 220 with a CNC yield of >80%.

KEYWORDS: Colloidal dispersion, Gaseous acid, Hydrolysis, Nanocellulose, TEMPO-oxidation

INTRODUCTION

Cellulose nanocrystals (CNCs), isolated from plant-based fibers or a microbial cultivation product, are a premium example of a modern renewable nanomaterial. Aside from its fully biobased origin, it is biodegradable, nontoxic, and possesses chirality as well as high mechanical strength. Proposed applications for CNCs include composites, chiral templates, biomedical devices, and responsive materials, to name a few. However, the sustainable aspect of CNCs is compromised by its production, generally performed with ca. 65% aqueous sulfuric acid, leading to difficult purification procedures, high water consumption, and poor yields. To tackle these problems, our group recently introduced a method that utilizes HCl vapor or gas for cellulose hydrolysis and subsequent CNC manufacturing. The approach has several advantages: it leaves solid fibers as a product and they are far easier to purify than an aqueous reaction mixture, resulting in higher yields and minimal water consumption. Moreover, the gaseous acid is easier to recover and recycle than a liquid/solid reaction product. On the other hand, the major problem with the HCl gas process lies in the difficult dispersion of uncharged CNCs from the already hydrolyzed fiber matrix. A proof-of-concept with dispersion in formic acid exhibited sizable CNC yields of >97% but formic acid with heavy (tens of hours) sonication is scarcely a viable dispersion medium even under laboratory conditions. Attempts with alternative dispersion methods have failed to reach the remarkable yields. Here, we want to communicate a straightforward way to utilize HCl gas for manufacturing CNCs in high yields from a commercially available bacterial cellulose (BC) substrate. Although more than often quoted to be highly crystalline, BC has been shown to possess a so-called leveling-off degree of polymerization (LODP) and it also has been utilized in CNC preparation. The actual isolation of CNCs was facilitated by a second step with oxidation catalyzed by N-oxoammonium cation of 2,2,6,6-tetramethylpiperidine-1-oxyl (TEMPO) that imparts charge on the CNC surface, enabling aqueous dispersion of CNCs.

RESULTS AND DISCUSSION

The full procedure for obtaining CNCs from BC is depicted in Figure 1. Unlike hierarchically arranged cellulose microfibrils in plant cell walls, BC is readily excreted by certain microbes into an isotropic network of microfibrils, forming a hydrogel of pure cellulose with high accessibility. First, the BC cubes intended for dessert foodstuff were washed with mild alkali to remove sugars and other taste ingredients to obtain a pure BC hydrogel...
The following drying step from hydrogel into aerogel was crucial: freeze-drying had to be performed to pristine, wet BC, which was then frozen by the emerging vacuum during the first seconds of the process (Figure 2d, Figure S2). Usually in freeze-drying, the sample is frozen before placing it in the vacuum. However, when a wet, unfrozen sample is positioned under a vacuum of a freeze-drier, it freezes almost immediately due to three different factors: (i) depression of the melting point, (ii) evaporative cooling of the sample as the water immediately starts to vaporize, and (iii) a compressive decrease of temperature as pressure is decreased. This resulted in Brunauer–Emmett–Teller (BET) surface area of 29 m²/g (Figure S5). By contrast, when the sample was frozen before placing it in a freeze-dryer, as is the case in conventional freeze-drying, the surface area was far smaller (BET area 16 m²/g, Figures S2, S6). This can also be visually observed from Figure 2e. Furthermore, merging several cubes together by blending and subsequent drying led to severe loss in accessibility, as attested by a weak hydrolysis effect (see Figure 3). The reason behind the distinctions in the freezing methods (Figure 2d vs 2e) may lie in different routes to crystallization of water but a systematic study is outside the scope of this letter.

Purified and freeze-dried bacterial cellulose (BC) cubes were hydrolyzed (dry matter content 94%) with HCl gas (1 bar pressure). The degree of polymerization (DP) was decreased from 2590 to 220 over 21 h (Figure 3), indicating the cleavage of disordered cellulose between the crystallites in semicrystalline cellulose microfibrils, i.e., LODP, necessary for CNC isolation. This DP decrease correlates well with the LODP value reported earlier for BC. The yield of 21 h hydrolysis after rinsing with water was 86.8%. Akin to the plant-based cellulose in fiber form, the bulk of the hydrolysis occurred within minutes of exposing the correctly dried BC to 1 bar HCl (g). Moreover, the BET surface area of BC increased from 29 to 52 m²/g upon hydrolysis (Figures S5, S7). After TEMPO-mediated oxidation, the final product with a carboxylate content of 1.0 mmol COOH/g BC (Figure S1) turned to gel after purification. The zeta potential of the CNCs amounted to ca. 40 mV (Table S2). The yield of TEMPO-mediated oxidation was 93.1%, resulting in a total yield (hydrolysis and oxidation) of 80.8% after centrifugation.

Hydrolyzed and TEMPO-oxidized BC was finally disintegrated into CNCs by sonication, as visualized with atomic force microscopy (AFM) imaging and transmission electron microscopy (TEM) (Figure 4a,b). To prevent aggregation artifacts, cryo-TEM images (Figure S3) were used for image analysis (Figure S4) which revealed rather narrow distributions of width and length (Figure 4c,d). Although a small number of
longer fibrils could be discerned from the images, the vast majority of CNCs were within 100−300 nm length with the mean at 170 nm, correlating fairly well with the DP value (Figure 3). The measured length for CNCs is somewhat shorter than in a number of previous accounts on BC-based CNCs, but these works have generally not presented statistical histograms on the CNC dimensions. Furthermore, the high yield and high charge due to TEMPO-oxidation may also affect the size distribution of the CNCs in this study.

It is curious that in HCl gas hydrolysis followed by TEMPO-oxidation, the CNC yield here from BC (>80%) is markedly higher than previously from plant-based (cotton) fibers (~50%). Because the DP reduction and carboxylic contents are roughly similar in both cases, the reason for the higher CNC yield must lie in the physical structure: the correctly chosen drying method. When the hydrolyzed BC was leached in water and the filtrate was analyzed with high performance anion exchange chromatography (HPAEC), the data showed that 13 wt % glucose could be released from the substrate after hydrolysis (Table S3). This is 1 order of magnitude different from cotton-based fibers where few wt % of glucose could be leached from the fibers hydrolyzed under similar conditions.

Overall, it appears that HCl gas hydrolysis on BC is faster and more efficient than on plant-based fibers.

A highly porous structure of the freeze-dried BC (Figure 2d) is also more accessible to cavitation during sonication. The hierarchical structure of the plant fiber is largely resistant to sonication, as shown earlier with TEMPO-oxidized microcrystalline cellulose (MCC) with maximal carboxylate contents where the yields stalled at a maximum of 20%. By contrast, a more recent work with TEMPO-oxidation amounted to 70% yields from MCC, but only after intensive high-energy sonication for >30 min, which may prove to be a bottleneck yielding from MCC, but only after intensive high-energy sonication for ≥30 min, which may prove to be a bottleneck. Here, utilizing a BC intended for foodstuffs is certainly not a viable route beyond demonstration but recent research has shown how BC can be produced from, e.g., agricultural waste resources as well as minimizes chemical and water consumption; (iii) dry gaseous acid can be easily recycled and it further minimizes the water consumption when purification procedures, such as dialysis in liquid/solid systems, can be
replaced by controlled rinsing with moderate amounts of water as nearly all acid is removed by desorption after the hydrolysis. Indeed, the full recovery of aqueous acid has prevented sustainable, large scale use of acid in hydrolysis of cellulose in the past.\textsuperscript{35} Furthermore, the thermal stability of carboxylic groups should be higher than those of sulfate groups which are the norm for charged groups on CNCs.\textsuperscript{33}

**REFERENCES**

Experimental details, conductometric titrations curves, SEM images, cryo TEM images and background for image analysis, BET raw data, WAXS data analysis, zeta potential analysis, HPAEC analysis (PDF)

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**Funding**

P.P. acknowledges Academy of Finland (Grant no. 315768) and T.P. Acknowledges Academy of Finland (Grant no. 300364).

**Notes**

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

We acknowledge Ms. Suvi Muukka and Ms. Xuetong Shi for auxiliary laboratory work with HCl gas hydrolysis and TEMPO-mediated oxidation. We acknowledge Ms. Tuyen Nguyen for auxiliary laboratory work with zeta potential measurement. We acknowledge Ms. Yingfeng Wang for auxiliary laboratory work with HPAEC analysis. T.P. acknowledges Academy of Finland (Grant no. 300364) for financial support. HYBER Centre of Excellence (2014–2019) and the Aalto University Nanomicroscopy Center (Aalto-NMC) are acknowledged for the use of its facilities. Moreover, E.K. is grateful for the support by the FinnCERES Materials Bioeconomy Ecosystem.

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**ASSOCIATED CONTENT**

Supporting Information The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acssuschemeng.9b04005.


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