Amorphous cellulose nanofiber supercapacitors

Mikio Fukuhara1,2*, Tomoyuki Kuroda1, Fumihiko Hasegawa3, Toshiyuki Hashida2, Mitsuhiro Takeda2, Nobuhisa Fujima4, Masahiro Morita5 & Takeshi Nakatani5

Despite the intense interest in cellulose nanofibers (CNFs) for biomedical and engineering applications, no research findings about the electrical energy storage of CNF have been reported yet. Here, we present the first electroadsorption effects of an amorphous cellulose nanofiber (ACF) supercapacitor, which can store a large amount of electricity (221 mJ m\(^{-2}\), 13.1 W kg\(^{-1}\)). The electric storage can be attributed to the entirely enhanced electroadsorption owing to a quantum-size effect by convexity of 17.9 nm, an offset effect caused by positive polar C=O radicles, and an electrostatic effect by appearance of the localised electrons near the Na ions. The supercapacitor also captures both positive and negative electricity from the atmosphere and in vacuum. The supercapacitor could illuminate a red LED for 1 s after charging it with 2 mA at 10 V. Further gains might be attained by integrating CNF specimens with a nano-electromechanical system (NEMS).

Recently, the examination of material and biomedical properties of biodegradable cellulose nanofibers (CNFs) that are thermally stable, strong durable, and light-weight has gained significant attention\(^1\)\(^-\)\(^4\). The completely individualised CNFs were developed by 2,2,6,6-tetramethylpiperidine-1-oxyl radical (TEMPO)-mediated oxidation under aqueous conditions\(^5\). Although CNFs are assumed to be renewable with innumerable promising applications\(^6\), no research findings about the electrical energy storage of CNF have been reported yet, except for the use of cellulose separators in lithium-ion batteries (LIBs)\(^7\)\(^,\)\(^8\). Recently, we found that amorphous titanium-oxides (golden, a-TiO\(_{2-x}\), ATO)\(^9\)\(^,\)\(^10\), perfluorinated polymer (APP)\(^11\), and aluminium-oxides (blackish, a–Al\(_{2-y}O_3\), AAO) supercapacitors\(^12\)\(^-\)\(^15\), showed one hundred times higher capacitance than that of a conventional aluminium electrolytic capacitor\(^16\), illumination of red LED light\(^17\), and a capturing effect of a positive and negative charges from the atmosphere\(^12\), respectively. Especially, the AAO is suitable for applications of heavy electricity such as stationary storage batteries. These amorphous capacitors are completely different from conventional "wet" cells, such as electric double-layer capacitors (EDLCs) and LIBs, which are controlled through the diffusivity of ions\(^14\)\(^,\)\(^15\). However, they are artificial products which are not environmentally friendly. If we can provide a supercapacitor composed of renewable papers, which are made by dewatering a dilute suspension of cellulose fibres, it could cast new light on paper electronics. Strictly speaking, however, there is no data about electroadsorption of paper, especially CNF. Herein, we report the high-performance electric storage of "dry" amorphous cellulose nanofibers (ACF). Our results show that the insulating ACF specimen with nanometre-sized cavities and high work function, is an ideal candidate for supercapacitors with potential applications of light electricity such as handheld electronic devices, transportation, and renewable energy storage for power grids.

Results and discussion

An electric storage system of ACF. The ACF specimens with a surface area of 2 cm\(^2\) exhibit discharging behaviours under constant currents of 20 nA–12.5 μA, after 2 mA–10 V charging for 50 s (Fig. 1a). All curves show an ohmic IR drop to approximately 3 V through a gradual decrease (see comparison with ATO, APP, and AAO in Supplementary Information hereafter, referred to as (SI) Fig. S4). The IR- drop is due to internal charging of unsaturated cells as well as the EDLC\(^18\). We repeated the test under 2 mA-rapid charging/1μA-discharging up to 30 times (Fig. 1b). The discharging time increases and the IR- drop decreases with every cycle repeated. The characteristic of increase in electric storage with increasing charge/discharge frequency is similar to that of EDLCs and LIBs which take a long time to charge under constant current\(^19\). Contrary to conventional charging, we tried to enhance the charge by voltage application for a 10-s duration. The applied voltage (V) dependency on the stored electric energy (E) (Fig. 1c) can be expressed as \(E = 5 \times 10^{-5} e^{0.014 V}\). The maximum energy at 400 V

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1 New Industry Creation Hatchery Center, Tohoku University, Sendai 980-8579, Japan. 2 Fracture and Reliability Research Institute, Graduate School of Engineering, Tohoku University, Sendai 980-8579, Japan. 3 National Institute of Technology, Sendai College, Natori 981-1239, Japan. 4 Faculty of Engineering, Shizuoka University, Hamamatsu 432-8561, Japan. 5 Cellulose Nanofiber Research Laboratory, Nippon Paper Industries, Fuji 417-8520, Japan. **email: mikio.fukuhara.b2@tohoku.ac.jp**
corresponds to 221 mJ m⁻² (13.1 W kg⁻¹, and 1.6 Wh kg⁻¹). We illuminated a red LED to prove the electric energy storage of ACF. The device with a surface area of 2400 mm² lit the LED for 1 s (Fig. 1d) after it was charged with 2 mA at 10 V. Further gains might be attained by integrating CNF specimens with a nano-electromechanical system (NEMS) (see Fig. S9 in SI).

Complex evaluation of electric storage and I–V characteristics. To analyse non-destructively the electrostatic contribution of the specimen, we measured the AC impedance from 1 to 1 MHz. A Nyquist plot of the impedance data is shown in Fig. 2a. The impedance variation with frequency data of ACF follows the combined pattern of a quadrant, straight horizontal line (inset of Fig. 2a), line with a slope of π/4 rad, and near-vertical line, attributing to a series-RC circuit. The π/4 rad region (Warburg region) is a consequence of the distributed resistance/capacitance in the porous electrode. The imaginary and real impedances rapidly increase up to 4 and 2 MΩ in the lower-frequency region of the Bode diagram, respectively (Fig. 2b). Moreover, the decrement in phase angle to -90° with decreasing frequency is another evidence of DC charging (Fig. 2c). This means that each capacitor on the ACF specimen is connected to a series circuit, \( C = \sum_{k=1}^{n} C_k = nC \). Therefore, the ACF offers an approximately ideal electric distributed constant (EDC) structure for enhancing electrical power storage. The experimental curve of series capacitance \( C_s \) can be expressed as \( C_s = 1.85f^{-0.494} \) \( (r^2 = 0.9984) \), where modulating the frequency \( f \) allows a considerable increase in DC capacitance. Figure 2d represents double I–V and R–V characteristics between -200 and +200 V in air. The curves are asymmetric relative to zero bias (inset of Fig. 2d), which is similar to the Coulomb blockade behaviour. The current \( I \) reached zero at -6.5 and +6.5 V upon increasing and decreasing the applied voltage \( V \), respectively. The zero current at -6.5 and +6.5 V correspond to the emission of electrons from the negative to the positive electrode and from the concave to the convex portions, respectively. This proves the electricity switching effect in rechargeable dry solid AAO supercapacitors.

Structural morphologies and surface characteristics by TEM and AFM. We investigated the structural morphologies and surface characteristics of the ACF specimen. The wide-field X-ray diffraction pattern (Fig. 3a) shows that the specimen consists of an amorphous cellulose phase, characterised by two broad peaks at
approximately 16 and 22°. Few nanocrystals are recognised from the continuous Debye rings in the selected-area electron diffraction pattern (inset of Fig. 3a) (corresponding to the red circle zone of Fig. S2 in SI). However, the contribution of nanocrystals to electric energy storage is unclear (SI Fig. S5). Figure 3b shows the changes in atomic pair distribution functions under a strong irradiation of 100–200 keV (SI Fig. S3). The intensity peaks of the C1–O5, C1–C3, and C3–C6 bonds are at approximately 0.14, 0.26, and 0.39 nm, respectively\textsuperscript{20}, and the corresponding change rates of distance as functions of the applied voltage are presented in Fig. 3c. Electroadsorption\textsuperscript{21} causes volume shrinkage of C3–C6 bonds to 2% due to a -1.98 GPa Maxwell compressive stress (electric field stress) (SI S6). This could be caused by the offset effect of positive polar C6=O6 radicles of carbonyl groups\textsuperscript{11,22}, with a rotation of C6–sodium carboxylate side chain about the C5–C6 bond\textsuperscript{23}. Amorphous structure prefers to rotate. Figure 3d shows an atomic force microscopy (AFM) image of the surface structure of specimen, with a convex diameter of 17.9 nm and a concave diameter of 13.1 nm. The fibrous appearance of the outer-surface resembles the uneven surface of ATO\textsuperscript{9,10}, APP\textsuperscript{11}, and AAO\textsuperscript{12,13,15} (SI Fig. S6). When the applied voltage changed from −20 V to +40 V in 600 s in air (SI Fig. S7), the electrostatic potential distribution histogram shifted to negative values (Fig. 3e). The negative shift (inset of Fig. 3e) can be attributed to the electron emission from the negative cantilever and an electrostatic induction from the positive cantilever in noncontact AFM.

**Molecular structure and quantum-size effect for electric storage.** The ACF structure, consisting of sodium (1→4)-β-D-poly-glucuronate containing C=O radicles\textsuperscript{24} with permanent dipoles (Fig. 4a)\textsuperscript{22,25}, is similar to the APP structure\textsuperscript{11}. Therefore, we inferred that the superior electric storage on the uneven ACF specimen is derived from the same quantum-size effect\textsuperscript{26,27} described in Fig. 4b, which induces a relative increase in the combined electrons. This leads to fewer free outer electrons due to the screening effect. Figure 4c presents the convex diameter dependencies of the calculated electrostatic potential and the induced outer electron pressure of carbon atoms surrounding the ACF structure, by help of the Thomas–Fermi screening effect (discussed in the SI S9). The decrease in diameter make increase the negative potential and the positive pressure. The calculated potential is −22.5 eV at 17.9 nm convex diameter. Therefore, the work function of ACF is negatively higher than those (−5.5, −10.1, and −20.7 eV) of ATO\textsuperscript{19}, APP\textsuperscript{11}, and AAO\textsuperscript{15}, respectively (SI Fig. S9). Figure 4d shows a morphological schematic proposing a possible mechanism for large electrical charges. The uneven surface serves as the EDC circuit (Fig. 4e) with an insulating layer containing tiny capacitors throughout the bulk. Similar to how
plant-produced cellulose absorbs atmospheric CO₂ via photosynthesis, ACF is also expected to capture both positive and negative electricity from the atmosphere, preventing the greenhouse effect. Thus, the ACF supercapacitor would be suitable for applications of light electricity such as handheld electronic devices, transportation, and renewable energy storage for power grids. However, this paper is the first report that presents the high-performance electric storage of “dry” ACF supercapacitor. We must next investigate electrical characteristics such as specific capacitance, energy efficiency of the charge–discharge process, cyclic stability for practical use.

**Optimised structure of ACF and its electronic role.** Finally, we investigate the reason why the CNF possesses superior electric adsorption. We optimized the local structures around CH₂OH and COONa radicals in TEMPO-oxidised native cellulose (C₁₂H₁₀O₁₀) and NaOH-extracts of TEMPO-oxidised native cellulose (C₁₂H₁₂O₁₁Na), respectively. The local structures of these celluloses are depicted in Fig. 5a,b, respectively. We then simulated the density of states (DOS) for C₁₂H₁₀O₁₀ and C₁₂H₁₂O₁₁Na, respectively. In sharp contrast to DOS of C₁₂H₁₀O₁₀ with OH radical, an isolated electronic state appears at − 4.5 eV in the band gap for C₁₂H₁₂O₁₁Na with COONa radical (Fig. 5d). This localized state corresponds to the empty 3 s orbital of Na cation shown in Fig. 5b. Here, it should be noted that the localized electrons present near the two-atomic vacancies in the AlO₆ cluster of AAO induce positive charges (electrostatic effect) on the inside of the insulating oxide surface, resulting in the adsorption of many electrons under electron-beam irradiation. Hence, we infer by analogy that the occurrence of the localized electron in the vicinity of COONa radical induces positive charges (electrostatic effect) on the inside of the insulating ACF surface, leading to high adsorption of many electrons from the atmosphere and in vacuum. Thus, the COONa radical in flexible ACF plays an important role for superior electric adsorption.

**Conclusions**

We demonstrated the high-performance electric energy storage (221 mJm⁻², 13.1 Wkg⁻¹) of “dry” amorphous nanocellulose fiber supercapacitor with nanometre-sized cavities and high work function, based on the quantum-size and enhanced electroabsorption effects, respectively. From appearance of localized electrons, C₆–sodium
carboxylate (COONa) radicals in ACF play crucial role (electrostatic effect) for high-performance electric storage. The integration of the film with NEMS is likely to provide potential applications in light electricity such as handheld electronic devices, transportation, and renewable energy storage for power grids.

Methods

TEMPO-oxidized CNFs (COONa content: 1.48 mmol g⁻¹, TC-01A) with 3-nm diameters were prepared by Nippon Paper Industries. The 10-μm-thick ACF specimen was fabricated on the Al substrate via slip casting. Sample structure was examined by X-ray diffraction (XRD) in reflection mode with monochromatic Cu Ka radiation. To avoid destroying of nanofibrils under strong and long-term electron-beam irradiation, we conducted selected-area electron diffraction (SAED) analyses at 100–200 keV with the electron density of 3 nA/m². Electron-refraction and -irradiation were performed using transmission electron microscopy (JEM-2100, JEOL). The noncontact-scanning Kelvin probe-atomic probe microscopy (NC-AFM, JSPM-5200, JEOL), based on the measurement of the electrostatic force gradient was applied to measure the absolute electrical potential between the Pt-coated cantilever tip from −100 to +100 V and the ACF surface as the work-function difference. The AC impedance and DC charging/discharging behaviours of each RC combination and illumination test of a red LED light (2 V–100 μA) were analysed using the galvanostatic charge/discharge measured using a potentiostat/galvanostat (SP-150, BioLogic Science Instruments) with DC’s of 10 V, 10 pA ~ 100 mA for ~900 s and charging currents of 10 mA at 293 K. A 2 × 10⁻⁴ W red LED lump was used to verify the electric energy storage. The optimised local atomic configurations of the C₁₂H₂₀O₁₀ and C₁₂H₁₇O₁₁Na were determined through a plane-wave-based first-principles density functional calculation (VASP 5.3)²⁸.

Figure 4. (a) Structure of sodium (1→4)-β-D-poly-glucuronate cell with green carbon, pink oxygen, light-blue hydrogen, and red sodium atoms. (b) Schematic diagram for calculations based on the Thomas–Fermi statistical method. (c) Convex dependences of the electrostatic potential and electron pressure with diameter. (d) Schematic representation of the microscopic electric energy storage used in this study. (e) The electric distributed constant circuit of the amorphous ACF surface.
References

1. Kafy, A. et al. Cellulose long fibers fabricated from cellulose nanofibers and its strong and tough characteristics. Sci. Rep. 7, 17683. https://doi.org/10.1038/s41598-017-17713-3 (2017).

2. Abibol, T. et al. Nanocellulose, a tiny fiber with huge applications. COBIOT 39, 76–88. https://doi.org/10.1016/j.copbio.2016.01.002 (2016).

3. Abdel-karim, A. M., Salama, A. H. & Hassan, M. L. Electrical conductivity and dielectric properties of nanofibrillated cellulose thin films from bagasse. J. Phys. Org. Chem. 31, e3851. https://doi.org/10.1002/poc.3851 (2018).

4. Martin-Martinez, F. J. Designing nanocellulose materials from the molecular scale. PNAS 115, 7174–7175. https://doi.org/10.1073/pnas.1809308115 (2018).

5. Saito, T. & Isogai, A. TEMPO-mediated oxidation of native cellulose. The effect of oxidation conditions on chemical and crystal structures of the water-insoluble fractions. Biomacromol 5, 1983–1989. https://doi.org/10.1021/bm0497769 (2004).

6. Tobjörk, D. & Österbacka, R. Paper electronics. Adv. Mater. 23, 1935–1961. https://doi.org/10.1002/adma.201004692 (2011).

7. Kuribayashi, I. Characterization of composite cellulosic separators for rechargeable lithium-ion batteries. J. Pow. Sour. 63, 87–91 (1996).

8. Chun, S.-J. et al. Eco-friendly cellulose nanofiber paper-derived separator membranes featuring tunable nanoporous network channels for lithium-ion batteries. J. Mater. Chem. 22, 16618–16626. https://doi.org/10.1039/c2jm32416f (2012).

9. Fukuhara, M. et al. Superior electric storage in de-alloyed and anodic oxidized Ti-Ni-Si glassy alloy ribbons. Phys. Status Solidi. RRL 7, 477–480. https://doi.org/10.1002/pssr.201307195 (2013).

10. Fukuhara, M. et al. Amorphous titanium-oxide supercapacitors with high capacitance. EPL 128, 58001. https://doi.org/10.1209/0295-5075/128/58001 (2019).

11. Fukuhara, M., Kuroda, T., Hasegawa, F. & Sueyoshi, T. Superior electric storage on an amorphous fluorinated polymer surface. Sci. Rep. 6, 22012. https://doi.org/10.1038/srep22012 (2016).

12. Fukuhara, M. et al. Amorphous aluminum-oxide superconductors. EPL 123, 58004. https://doi.org/10.1209/0295-5075/123/58004 (2018).

13. Fukuhara, M. et al. Effects of temperatures and carbon dioxide nanobubbles on superior electric storage for anodically oxidized films of AlY amorphous alloy. Adv. Adv. 9, 095202. https://doi.org/10.1063/1.5102082 (2019).

14. Fukuhara, M. et al. Surface analyses of amorphous aluminum oxides with AlO₆ clusters. MRS Com. https://doi.org/10.1557/mrc.2020.82 (2020).

15. Fukuhara, M. et al. AlO₆ clusters’ electric storage effect in amorphous alumina supercapacitors. Sci. Rep. 11, 1699. https://doi.org/10.1038/s41598-021-81483-2 (2021).

16. Fukuhara, M., Kuroda, K. & Hasegawa, F. Amorphous titanium-oxide supercapacitors. Sci. Rep. 6, 35870. https://doi.org/10.1038/srep35870 (2016).

17. Fukuhara, M. et al. Anodic oxidization of Al-Y amorphous alloy ribbons and their capacitive properties. J. Alloy Comp. 776, 757–762. https://doi.org/10.1016/j.jallcom.2018.10.346 (2019).
18. Okamura, M. Electric Double Layer Capacitor and Its Storage System (Nikkkan Kogyo, 2011).
19. Kötz, R. & Carlen, M. Principles and applications of electrochemical capacitors. Electrochim. Acta 45, 2483–2498. https://doi.org/10.1016/S0013-4686(00)00354-6 (2000).
20. Fink, H.-P., Philipp, B., Paul, D., Serima, R. & Paakkari, T. The structure of amorphous cellulose as revealed by wide-angle X-ray scattering. Polymer 28, 1265–1270. https://doi.org/10.1016/0032-3861(87)90435-6 (1987).
21. Dupeyrat, M. & Nakache, E. Electrocapillarity and electroadsorption. J. Colloid Interface Sci. 73, 332–344. https://doi.org/10.1016/0021-9797(80)90080-6 (1980).
22. Hegde, V. Dielectric study of biodegradable and/or bio-based polymeric materials, PhD thesis (HAL, Univ. Grenoble Alpes, 2017).
https://tel.archives-ouvertes.fr/tel-01689886
23. Gardner, K. H. & Blackwell, J. The structure of native cellulose. Biopolymers 13, 1973–2001. https://doi.org/10.1002/bip.1974.360131005 (1974).
24. Iosiga, A. & Kato, Y. Preparation of polyuronic acid from cellulose by TEMPO-mediated oxidation. Cellulose 5, 153–164. https://doi.org/10.1023/A:100928603673 (1998).
25. Kumagai, T., Suga, T. & Kubono, A. High ordered structure of polyuria thin films by vapor deposition polymerization. Inst. Electr. Eng. Jpn. Trans. FM 30, 186–192 (2010).
26. Paulus, P. M. et al. Surface and quantum-size effects in Pt and Au nanoparticles probed by 197Au Mössbauer spectroscopy. Phys. Rev. B 64, 205418. https://doi.org/10.1103/PhysRevB.64.205418 (2001).
27. Tringides, M. C., Jalochowski, M. & Bauer, E. Quantum size effects in metallic nanostructures. Phys. Today 60, 50–54. https://doi.org/10.1063/1.2731973 (2007).
28. Gutiérrez, G. & Johansson, B. Molecular dynamics study of structural properties of amorphous Al2O3. Phys. Rev. B 65, 104202. https://doi.org/10.1103/PhysRevB.65.104202 (2002).

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Author contributions
M.F. carried out the electric storage analysis and wrote the paper. F.H. edited the paper and T.K. carried out the electric measurements. T.H. assisted with the device fabrication. M.T. conducted the TEM and electron irradiation analyses. N.F. performed the MD simulations. M.M. prepared amorphous CNF sheet. T.N. carried out the molecular analysis. All authors discussed the results and commented on the manuscript. M.F. supervised all the work.

Competing interests
The authors declare no competing interests.

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Correspondence and requests for materials should be addressed to M.F.

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