Amino group in *Leptothrix* sheath skeleton is responsible for direct deposition of Fe(III) minerals onto the sheaths

Tatsuki Kunoh\(^1,2\), Syuji Matsumoto\(^1,2\), Noriyuki Nagaoka\(^3\), Shoko Kanashima\(^4\), Katsuhiko Hino\(^1\), Tetsuya Uchida\(^2\), Katsunori Tamura\(^1,2\), Hitoshi Kunoh\(^1,2\) & Jun Takada\(^1,2\)

*Leptothrix* species produce microtubular organic–inorganic materials that encase the bacterial cells. The skeleton of an immature sheath, consisting of organic exopolymer fibrils of bacterial origin, is formed first, then the sheath becomes encrusted with inorganic material. Functional carboxyl groups of polysaccharides in these fibrils are considered to attract and bind metal cations, including Fe(III) and Fe(III)-mineral phases onto the fibrils, but the detailed mechanism remains elusive. Here we show that NH\(_2\) of the amino-sugar-enriched exopolymer fibrils is involved in interactions with abiotically generated Fe(III) minerals. NH\(_2\)-specific staining of *L. cholodnii* OUMS1 detected a terminal NH\(_2\) on its sheath skeleton. Masking NH\(_2\) with specific reagents abrogated deposition of Fe(III) minerals onto fibrils. Fe(III) minerals were adsorbed on chitosan and NH\(_2\)-coated polystyrene beads but not on cellulose and beads coated with an acetamide group. X-ray photoelectron spectroscopy at the N1s edge revealed that the terminal NH\(_2\) of OUMS1 sheaths, chitosan and NH\(_2\)-coated beads binds to Fe(III)-mineral phases, indicating interaction between the Fe(III) minerals and terminal NH\(_2\). Thus, the terminal NH\(_2\) in the exopolymer fibrils seems critical for Fe encrustation of *Leptothrix* sheaths. These insights should inform artificial synthesis of highly reactive NH\(_2\)-rich polymers for use as absorbents, catalysts and so on.

The biological strategies that living organisms use to produce a wide range of specially designed organic–inorganic materials such as bone, teeth, and shells\(^1\) have been exploited for artificially synthesizing materials for biomedical, industrial, and technological purposes and customizing their biomineral properties\(^1–5\). Of the many types of microbial sorbents (i.e., fungi, bacteria, and yeasts), some bacteria such as Fe/Mn-oxidizing bacteria frequently produce unique, ingenious structures with specific functions for their survival (e.g., to protect themselves, to stock and utilize nutrients, to move and function well) by biomineralization\(^6\). Not only the visual uniqueness of such structures but also their structural and physicochemical properties (e.g., exquisite organic–inorganic materials that are not readily synthesized artificially) have attracted researchers’ attention and provided many insights for developing and/or improving beneficial manufactured goods\(^7–10\).

*Leptothrix* species, common inhabitants of freshwater environments, oxidize Fe(II) in the presence of low oxygen concentrations\(^11–13\) and divide to form catenulate cells. They excrete exopolymer fibrils, which are entangled with interfibrillar cross-linkers such as proteins and disulfide bridges to eventually form a microtubular, immature, organic sheath skeleton\(^7,14,15\). Sugar analysis of sheath fibrils prepared from *L. cholodnii* SP-6 (hereafter referred to as SP-6) has elucidated that the basic structure of sheath fibrils comprise heteropolysaccharides containing galactosamine (GalN) and glucosamine (GlcN), both of which have a terminal NH\(_2\)\(^16\). Another polysaccharide polymer with a terminal NH\(_2\), chitosan, is frequently used to adsorb transition metals from wastewater via the chelating ability of NH\(_2\) and an adjacent hydroxyl group\(^17–20\), suggesting that the terminal NH\(_2\) within the sheath fibrils might play a key role in metal encrustation.

\(^1\)Core Research for Evolutionary Science and Technology (CREST), Japan Science and Technology Agency (JST), Okayama University, Okayama, 700-8530, Japan. \(^2\)Graduate School of Natural Science and Technology, Okayama University, Okayama, 700-8530, Japan. \(^3\)Advanced Research Center for Oral and Craniofacial Sciences, Okayama University Dental School, Okayama, 700-8558, Japan. \(^4\)Hyashibara Co., Ltd., Okayama, 702-8006, Japan. Correspondence and requests for materials should be addressed to J.T. (email: jtakada@cc.okayama-u.ac.jp)
The immature sheath skeleton of another *L. cholodnii* strain, OUMS1 (hereafter OUMS1-WT) becomes encrusted with metal cations and/or metal solid phases to eventually form uniquely structured microtubular sheaths comprising an organic–inorganic complex enriched with Fe, Si, P, and Ca\(^{21–23}\). Since an Fe(II)-oxidizing bacterium oxidizes Fe(II) and uses the generated electron as an energy source in the presence of low concentrations of oxygen\(^{11–13}\), the sheath probably possesses an ecological role in avoiding encrustation in Fe(III) oxyhydroxides. Fe(III) minerals (~50 nm diameter) that are generated abiotically in the culture medium (=complex of ferric oxyhydroxides as major components and inorganic components of the medium components as minors, Fig. S1) were reported to adhere directly to the sheath materials of SP-6\(^{24}\) and OUMS1-WT\(^{25}\). The Fe(III) minerals also directly adhere to cell- and protein-free sheath remnants, indicating that living cells and their proteins conjugated to sheath fibrils are not inevitably required for the Fe(III) mineral encrustation\(^{24,25}\). The metal encrustation of the sheath skeleton is considered to result from the interactions between aquatic phase inorganic cations and the functional groups in the sheath skeleton\(^{22}\). Indeed, a strong correlation exists between the presence of acidic polysaccharides with carboxyl groups (COOH) and the distribution of iron oxyhydroxides in *Leptothrix* sheaths\(^{6,26}\), but little is known about the roles of NH\(_2\) within the sheath fibrils on the metal encrustation.

In this study, we sought to ascertain the involvement of the terminal NH\(_2\) within the constitutive molecules of the immature sheath skeleton of OUMS1-WT in the adsorption of Fe(III) minerals to the skeleton by differential interference contrast (DIC) and fluorescent microscopy, scanning, transmission, and scanning-transmission electron microscopy (SEM, TEM, and STEM, respectively), energy-dispersive x-ray microanalysis (EDX), x-ray fluorescence and photoelectron spectroscopy (XRF and XPS, respectively) (Fig. 1).

**Results**

**Importance of sheath fibrils for Fe encrustation.** We examined the ultrastructure of OUMS1-WT grown in silicon-glucose-peptone medium (Table S1) (hereafter SGP)\(^{27}\) for 2 days and then in SGP + Fe-plate medium (hereafter SGP + Fe) for 2 more days for comparison with previous TEM images of the cells incubated in SGP + Fe plate for 3 days\(^{28}\). For comparison, a sheathless mutant of OUMS1 (named hereafter OUMS1-SL)\(^{28}\) was incubated and studied in similar conditions.

After 2 days incubation in SGP, chains of OUMS1-WT cells (ca 3–4 μm long) were encased within a thin, immature sheath, while longitudinally extending chains of OUMS1-SL cells (ca 7–8 μm long) were not (Fig. 2a, left). After another 2 days incubation in SGP + Fe plate, a few aggregated fibrils were seen around OUMS1-SL cells, suggesting fibril excretion is enhanced in the presence of the Fe source, but they did not assemble into a sheath (Fig. 2, lower center) as reported previously\(^{29}\). Electron-dense particles were deposited on and near the immature sheath encasing OUMS1-WT cells (Fig. 2a, upper center) and on aggregated fibrils far from OUMS1-SL cells (Fig. 2a, lower center). To confirm the absence of sheaths in OUMS1-SL, the culturing period of both isolates in SGP + Fe plate was extended to 14 days. The OUMS1-WT cells were evidently encased within thick sheaths having lots of hairy fibrils extending outward onto which electron-dense particles deposited (Fig. 2a, upper right). Aggregated fibrils 10–150 μm from the OUMS1-SL cell became electron-dense but never assembled into a sheath (Fig. 2a, lower right).

SEM/EDX analyses of OUMS1-WT cells incubated in SGP + Fe plate as above detected a mass of aggregated sheaths and apparent deposition of Si, P, and Fe on sheaths (Fig. 2b), while chained OUMS1-SL cells aggregated without being encased in sheaths and lacked any distinguishable element deposition, even of Fe (Fig. 2c).

From these microscopic results, we confirmed that (i) OUMS1-WT formed immature thin sheaths within 2 days in SGP, while exopolymer fibrils were not excreted from OUMS1-SL cells, and no sheath formed; (ii) OUMS1-SL cells excreted fibrils but failed to form sheaths even after an additional 2 days incubation in SGP + Fe; and (iii) inorganics including Fe were deposited on OUMS1-WT sheaths within another 2 days in SGP + Fe plate, while Fe was not deposited on chained OUMS1-SL cells, which lacked a sheath.

**NH\(_2\)** functional groups in the amino sugars of the OUMS1-WT sheaths play a role in Fe(III) mineral deposition. We examined whether amino sugars were components of OUMS1-WT sheaths as found for SP-6\(^{15,16}\) by using protein-free sheath remnants that had been chemically prepared from immature sheaths of OUMS1-WT cultured in SGP for 2 days (Fig. 3a). The GC/MS analysis revealed that GaN and GlcN were major saccharic components of the sheath fibrils of OUMS1-WT, as NH\(_2\)-holding materials in addition to amino acids, similar to those in the *L. cholodnii* SP-6 sheath fibrils\(^{15,16}\).
Figure 2. TEM images (a) of OUMS1-WT and OUMS1-SL after incubating in SGP and then with SGP + Fe plate and SEM images with EDX distribution patterns of elements (b) of both strains. (a) Left, SGP incubation for 2 days: cross section of OUMS1-WT cell encased with a thin, immature sheath (arrow) (upper left) and OUMS1-SL cell with no sheath (lower left). Center, additional 2-day incubation in SGP + Fe plate: electron-dense Fe particles on and near the immature sheath encasing OUMS1-WT cells (upper center) and on a few fibril aggregations far from OUMS1-SL cells (lower center, arrowhead). Right, additional 14-day incubation in SGP + Fe plate: deposition of electron-dense Fe particles in a thick sheath encasing OUMS1-WT cells and in abundant fibrils extending outward from the sheath surface (upper right). Electron-dense fibril aggregation seen apart from the OUMS1-SL cell (lower right), but no sheath has formed. (b) Aggregated immature sheaths of OUMS1-WT after incubating in SGP + Fe plate for 14 days with EDX distribution patterns of S, Si, Ca, P, and Fe showing apparent deposition of Si, P, and Fe on sheaths. (c) Aggregated chains of cells of OUMS1-SL after incubating in SGP + Fe plate for 14 days with EDX distribution patterns of S, Si, Ca, P, and Fe, lacking any distinguishable deposition, even of Fe.
Figure 3. GC/MS spectrograph showing sugar composition in protein-free OUMS1-WT sheath remnants and attachment of Fe(III) minerals on OUMS1-WT sheath skeleton treated with NH2-binding fluorescent reagents. (a) GalN and GlcN were detected in the sheath remnants as major saccharic materials. (b) Schematic figure of allophycocyanin-conjugated NH2-reactive reagent (APC-reagent) masking NH2 group. (c) Top, DIC images of immature sheaths encasing OUMS1-WT cell chains and longitudinally extending, sheath-uncovered OUMS1-SL cell chains after treatment with APC-reagent. Bottom, APC-reagent-treated OUMS1-WT sheaths turned blue within 30 min (inset) and fluoresced red; similarly treated OUMS1-SL were visibly unchanged (inset) and did not fluoresce. (d) Fe ratios in OUMS1-WT sheaths and those treated with APC-reagent, sulfo-NHS-acetate, or acetic anhydride after incubation in Fe(III)SGP relative to those in Fe-free SGP (with At% of Fe = 1). Note the suppressed Fe levels in these NH2-masked sheaths. (e) TEM images of OUMS1-WT cells. None (APC reagent-untreated) 1st, a cell encased with thin, immature sheath (arrow) after incubation in SGP; 2nd, electron-dense Fe(III) minerals (arrowheads) attached to surface of immature sheath and fibrils extending from the surface after incubation in Fe(III)SGP (inset, enlarged image showing mineral attachment to the extending fibrils [arrowhead]). APC reagent 3rd, APC-reagent-treated cell encased with a thin, immature sheath (arrow) incubated in SGP; 4th, no Fe minerals on or near the sheath even after incubation in Fe(III)SGP.

Whitish, immature sheaths encasing OUMS1-WT cells that had been incubated in SGP for 2 days were treated with allophycocyanin (APC)-conjugated NH$_2$-reactive reagent (hereafter APC-reagent) (Fig. 3b). Within 3 h, the OUMS1-WT sheaths turned visibly blue and, viewed with fluorescence microscopy, fluoresced an intense red (Fig. 3c, lower left), demonstrating the presence of terminal NH$_2$ in the sheaths. In comparison, cells of OUM1-SL, incubated in SGP and similarly treated with the APC-reagent, remained white and did not fluoresce (Fig. 3c, lower right). The lack of excretion of exopolymer fibrils from OUM1-SL cells within 2 days incubation in SGP (Fig. 2a) accounted for the lack of a positive response (change in color) or any notable deposition of elements such as Fe (Fig. 2c). Subsequently, the APC-reagent-treated OUM1-SL sheaths were incubated with a suspension of abiotic Fe(III) minerals in SGP [hereafter Fe(III)SGP] to examine the influence of APC-masked NH$_2$ in the sheath skeleton on Fe(III) mineral encrustation (Fig. 3b). XRF analysis of washed and freeze-dried untreated control sheaths revealed that the ratio of Fe was nearly 50 times higher in Fe(III)SGP relative to an atomic (At) % of Fe = 1 in Fe-free SGP (Fig. 3d). However, in the APC-reagent-treated sheaths, the relative ratio was enhanced approximately 10-fold more than after the SGP incubation (Fig. 3d). Similarly, the increase in Fe percentage after the Fe(III)SGP incubation was suppressed by two other NH$_2$-blocking reagents, sulfo-NHS-acetate or acetic anhydride (Figs 3d and S2a,b). Electron-dense particles were apparently deposited around the immature control sheaths after incubation in Fe(III)SGP (Fig. 3e, left two), while such particles were not seen on or around APC-reagent-, sulfo-NHS-acetate-, or acetic anhydride-treated OUM1-SL WT cells (Fig. 3e, right two, S2a,b). To confirm whether the deposited electron-dense particles corresponded to Fe minerals, we incubated the SGP-precultured OUM1-SL WT cells with Fe(III)SGP for 2 days. EDX analyses of cross-sectioned cells (Fig. 4a) showed that the electron-dense particles on or around the exopolymer fibrils were composed of Fe, bound to the medium components such as P and S, as judged by the similar distribution of Fe/O and Fe/Ps (Fig. 4c). The XRD analysis provided evidence that the Fe detected by EDX was actually Fe(III) oxihydroxide (Fig. S1a–c).

**Binding energy shift on the sheath surface caused by incubation with Fe(III)SGP.** Since XPS can determine the intrinsic binding energy of the atomic orbital, which shifts chemically with the surroundings of the atom, the electronic state of the material surface was analyzed using XPS. From the XPS measurement of SGP-incubated OUM1-SL sheaths, peaks of O1s (532 eV), N1s (399 eV), and C1s (284 eV) were detected, suggesting the organic nature of the sheaths (Fig. 5a). Additional peaks of Fe2p (710 and 723 eV) were detected in the sheaths incubated in Fe(III)SGP (Fig. 5b). The N1s peak yielded by Fe(III)SGP-incubated sheaths shifted toward a binding energy higher than that yielded by the SGP-incubated sheaths (Fig. 5c), suggesting that Fe(III) minerals affected the N-related functional group such as NH$_2$.60,31

**Binding energy shift on the surface of NH$_2$-holding C-polymers and NH$_2$-coated polystyrene beads caused by incubation with Fe(III)SGP.** The described results led us to consider that NH$_2$ in the OUM1-SL WT sheath skeleton plausibly contributes to encrustation of the skeleton with Fe(III) minerals. We verified this possibility in model experiments to compare the affinity of cellulose (β-glucose polymer) and chitosan (GlcN polymer) and that of NH$_2$-coated polystyrene beads and uncoated (plain) beads for Fe(III) minerals. The Fe ratio of chitosan increased drastically after incubation in Fe(III)SGP relative to At% of Fe = 1 in Fe-free SGP, while that of cellulose after the incubation in SGP was comparable to that in Fe(III)SGP (Fig. 6a). SEM and EDX indicated that aggregated Fe(III) minerals were attached to the chitosan fibrils after incubation, while the smooth surfaces of cellulose fibers were unchanged after incubation with SGP and Fe(CIII)SGP (Fig. 6b,c). In XPS, peaks of photoelectron O1s and C1s were detected for cellulose, while an additional N1s peak was detected for chitosan (Fig. 6d–g). The incubation in Fe(III)SGP yielded additional peaks of Fe2p for chitosan, but not for cellulose (Fig. 6f,g), suggesting the possible binding of Fe(III) minerals to NH$_2$ of chitosan. Similar to the results for OUM1-SL WT sheaths, the binding energy of N1s in chitosan shifted to a higher level (Fig. 6h), suggesting that NH$_2$ of chitosan was influenced by Fe(III) minerals.

To further confirm the involvement of NH$_2$ with the Fe encrustation, we examined the affinity of NH$_2$-coated polystyrene beads (hereafter NH$_2$ beads) and uncoated beads (hereafter plain beads) for Fe(III) minerals. When NH$_2$ beads were incubated with Fe(III)SGP, a precipitate formed within 30 min, while the suspension of plain beads remained turbid (Fig. 7a). SEM revealed that the surfaces of the NH$_2$-beads after Fe(III)SGP incubation were heavily coated with granular or rod-shaped Fe particles in contrast to the smooth surface of the plain and NH$_2$ beads after SGP incubation (Figs 7b and S3c). The Fe atomic percentage determined by XRF was ~6.0 for the NH$_2$-coated beads and ~1.3 for the plain after incubation in Fe(III)SGP (Fig. 7b). XPS detected photoelectron Cls and O1s peaks derived from polystyrene in plain beads and an additional N1s peak from the NH$_2$ beads (Fig. 7c,e). The peak patterns from the plain beads in SGP and in Fe(III)SGP were comparable (Fig. 7d). In contrast, additional Fe2p peaks were detected in NH$_2$ beads after incubation in Fe(III)SGP (Fig. 7e,f). Notably, incubation of NH$_2$ beads in Fe(III)SGP shifted the N1s peak toward a higher binding energy (Fig. 7g), again suggesting an interaction between Fe(III) minerals and NH$_2$. When acetylated NH$_2$ beads (NH-Ac beads) were incubated with Fe(III)SGP, lack of Fe attachment to these beads was confirmed by SEM, EDX, and XPS (Fig. S3b–e).

**Discussion**

The behavior of dissolved metals in natural bodies of water is strongly influenced by particular inorganic and organic materials, suggesting complex interactions of various metal-complexing agents in aquatic systems with microbes and/or their constituent polymers. Generally, extracellular polymeric substances (EPS) of bacteria contain carbohydrates, proteins, lipids, extracellular DNA, and humic substances, which possess various types of terminal functional groups in the molecules. The exopolymer fibrils of the _Leptothrix_ sheaths are made of EPS, suggesting an interaction between Fe(III) minerals and NH$_2$. When acetylated NH$_2$ beads (NH-Ac beads) were incubated with Fe(III)SGP, lack of Fe attachment to these beads was confirmed by SEM, EDX, and XPS (Fig. S3b–e).
in the *Leptothrix* sheaths and *Gallionella* stalks contribute to attracting dissolved cations\(^6,7\). Several lines of evidence presented here indicate that NH\(_2\) in amino sugars and amino acids in the *Leptothrix* sheath is involved at least in direct deposition of Fe(III) minerals onto the sheath skeleton, which is composed of organic exofibrils.

**Figure 4.** STEM image and EDX element distribution pattern on immature sheaths encasing OUMS1-WT cell incubated in Fe(III)SGP. (a) Top, STEM image of electron-dense particles attached to the sheath and distribution patterns of N and O. Bottom, distribution of P, S, and Fe. (b) Semiquantitative distribution maps of O and Fe determined by the Cliff-Lorimer method. The signals for both elements at the same location strongly suggest that Fe could exist as iron oxides. (c) Merged images for distribution of O/Fe, P/Fe, and S/Fe, suggesting that the electron-dense Fe particles plausibly bind the medium components.
The Fe(III) minerals generated in this study are a complex of Fe oxyhydroxides and light elements such as P, S, K, Ca, and Cl from the surrounding medium (Fig. S1). Fe oxyhydroxides tend to be negatively charged below pH 7.0. In addition, binding of Fe oxyhydroxides with the added P was proved to cause a more negative surface charge. Therefore, we infer that the interaction between negatively and positively charged Fe(III) minerals and NH$_2$, respectively, could be one of the driving forces for binding the minerals to *Leptothrix* sheath fibrils at pH 7.0 used in this study, although this assumption should be verified in the near future.

Note that we do not interpret our present results as meaning that NH$_2$ is the only functional group that interacts with Fe(III) minerals, because masking of NH$_2$ in sheaths with specific NH$_2$-reactive reagents did not completely block the Fe(III) minerals deposition onto the immature sheaths (Fig. 3d), although the reagents appeared to block most of the terminal NH$_2$ in the sheath skeleton. This incomplete blocking suggests the possible involvement of other functional groups in the deposition. On the basis of earlier reports, there is no reason to rule out possible synergistic reactions between NH$_2$ and other functional groups such as COOH. At present, we infer that polymer-directed mineralization is a general phenomenon that could occur in any system containing NH$_2$-enriched polysaccharides originating from an organism.

The energy shift that was detected in the OUMS1-WT sheaths, chitosan, and NH$_2$ beads incubated with Fe(III) minerals (Figs 5–7) could be due to the influence of Fe(III) minerals as they approach the N-atomic orbital, which could readily shift chemically in the atom. For example, the electron density around N is reduced when the covalent bond forms between NH$_2$ and Fe(III) minerals or from the intense attractive force of charged particles. On the basis of earlier reports, we infer that in the aqueous phase the positive charge from NH$_2$ and the negative charge from ferric oxyhydroxides may interact and that the chemical shift of N2p may reflect

**Figure 5.** Photoelectron spectra acquired by XPS from OUMS1-WT sheaths incubated in SGP or in Fe(III) SGP. SGP: (a) Peaks of C1s, O1s, and N1s detected. Fe(III)SGP: (b) Additional Fe2p peaks detected. (c) Shift of N1s peak toward higher binding energy.
Figure 6. Behavior of Fe(III) minerals on cellulose and chitosan. (a) Fe ratios on cellulose and chitosan after incubating in Fe(III)SGP relative to At% of Fe = 1 in Fe-free SGP. (b) Top, merged images of SEM and EDX Fe distribution on cellulose and chitosan incubated in SGP. Bottom, Merged images of cellulose and chitosan incubated in Fe(III)SGP, showing Fe deposition on chitosan. (c) Left, SGP-incubated chitosan without any fine particles attached. Right, Fe(III)SGP-incubated chitosan with abundant fine particles (arrowheads), especially on aggregated fibrils (arrows). (d,f) XPS spectra of cellulose mixed with either SGP or Fe(III)SGP; note O1s and C1s peaks but none for N and Fe. (e,g) XPS spectra for chitosan after incubation in either SGP or Fe(III)SGP; note O1s, C1s, and N1s peaks in SGP incubation, additional Fe2p peaks in Fe(III)SGP. (h) N1s peak shifted toward higher binding energy after Fe(III)SGP incubation.
this interaction, although further detailed analyses are necessary to verify these and/or other possibilities such as a covalent bond, Van der Waals force, and condensation reaction of OH⁻.

Mineral precipitation could be an important determinant of microbial activity levels in the environment, and thus, spatially resolved analyses of organic compound combined with high-resolution mineralogical information should enhance our understanding of biomineralization mechanisms and promote the development of templated models to use in fabricating new materials¹⁵. Detailed information on the synthetic mechanism for biological organic–inorganic complexes will help expand the use of these ingenious complexes to benefit life. A variety of industrial applications such as lithium-ion battery anode material, catalyst enhancer, and pigment have been developed for metal-encrusted *Leptothrix* sheaths⁷–¹⁰. Current information on the roles of NH₂ in Fe encrustation of the *Leptothrix* sheaths will greatly help in harnessing biomineralization mechanisms to create novel functional

Figure 7. Behavior of Fe(III) minerals on plain and NH₂-coated polystyrene beads. (a) Precipitation only on NH₂-coated beads (NH₂ beads) within 30 min after incubating in the suspension of Fe(III) minerals. (b) Top, merged images of SEM and EDX Fe distribution on plain and NH₂ beads incubated with SGP. Bottom, merged image of the respective beads incubated with Fe(III)SGP, showing Fe deposition on NH₂ beads. Fe ratios in Fe(III)SGP detected by XRF relative to At% of Fe = 1 on SGP-incubated beads: ~1.3 for plain beads, ~6.0 for NH₂ beads. (c,d) XPS spectra of plain beads incubated with SGP and Fe(III)SGP, respectively, detecting O₁s and C₁s but no N and Fe peaks. (e,f) XPS spectra of NH₂ beads incubated in SGP and Fe(III)SGP, respectively, detecting O₁s, C₁s, and N₁s with SGP incubation and additional Fe₂p peaks with Fe(III)SGP. (g) N₁s peak shifted toward higher binding energy after incubation in Fe(III)SGP.
materials. Toward creating novel materials, we should examine whether NH₂-blocking affects the potential of other functional groups to bind inorganics including Fe, how NH₂-blocking influences previously reported the functioning of sheath fibrils, and how we can artificially synthesize NH₂-rich polymers with high reactivity to inorganics for use as adsorbents and/or catalysts.

Organic polymers play important roles in ecosystems by adsorbing biologically important elements. Fe cycling, mediated by microbiological oxidation-reduction of the element, is a vital environmental process, both on the micro and global scales, and iron-oxidizing bacteria such as *Leptothrix* are significant facilitators of ferric iron reduction in ecosystems. A step-by-step approach such as the present study should improve our understanding of the complex systems of Fe circulation.

**Methods**

**Strains, medium, and culturing.** *Leptothrix cholodnii* strain OUMS1 (NITE BP-860) (hereafter OUMS1-WT) and its sheathless mutant (OUMS1-SL) were used. Cells of OUMS1-WT and OUMS1-SL, obtained from frozen stock cultures, were independently streaked onto SGP agar plates and incubated at 20 °C for 7 days. Single colonies were transferred separately to 25 ml of SGP in 200 ml aluminum foil-capped Erlenmeyer flasks and cultured on a rotary shaker at 20 °C and 70 rpm. After 2 days, 1–5 ml of the cell suspension (adjusted to 10 cfu ml⁻¹ by densitometry) was used to inoculate 25 ml of (i) SGP and (ii) SGP containing 500 μM FeSO₄ (SGP + FeSO₄), or (iii) 100 ml SGP containing three small pieces of Fe plate (SGP + Fe plate). Samples were examined after 2 days of incubation. Viability of OUMS1-WT in these culture conditions was confirmed by live/dead staining after a 4-day culture (Fig. S1g). Specimens incubated in SGP + Fe plate were used for preliminary microscopic observations, and those in SGP or SGP + FeSO₄ were mainly used for the following affinity tests and electron microscopy, unless otherwise stated.

**Abiotic preparation of Fe(III) minerals in SGP.** Our previous work provided evidence that in a shaken solution of SGP + FeSO₄, Fe(II) was oxidized to Fe(III) almost completely within 6 h. In the present study, uninoculated SGP + FeSO₄ was shaken similarly for 8-12 h to ensure generation of abiotic Fe(III) minerals. The suspension of Fe(III) minerals in SGP (hereafter Fe(III)SGP) was used for the affinity tests, as follows.

**Determination of amino sugar composition in protein-free OUMS1-WT sheath remnants.** Protein-free sheath remnants were prepared according to a previous protocol. OUMS1-WT cells, pelletized from 4–6 l of a 2-day-old culture in SGP, were washed once with ultrapure water (H₂O) and resuspended in 22.5 ml of the lysis solution containing 2.5 mM EDTA and 150–500 μg lysosome (Sigma-Aldrich), and incubated at 37 °C for 0.5–2 h. Then 2.5 ml of 10 % w v⁻¹ SDS was added, and the suspension was shaken for 0.5–2 h at room temperature. Thus-prepared sheath remnants were washed twice with H₂O, then treated with proteinase K (50–100 μg ml⁻¹, Nacalai Tesque, Kyoto, Japan) at 37 °C for 12 h. The final protein-free sheath remnants were washed with H₂O six times and analyzed for sugar composition as follows.

The sheath remnants (20 mg) were vacuum-dehydrated at 50 °C for 2 h, followed by gas phase hydrazinolysis at 110 °C for 1 h with Hydrachl S-204 (J-OI Mills, Tokyo, Japan) to release oligosaccharides from the remnants. The resultant product was dissolved in 1 ml of 0.2 N sodium bicarbonate (NaHCO₃, pH 6.7), followed by filtration (0.8 μm pore, Advantest, Tokyo, Japan) and subsequent overnight dialysis against 100 mM 0.2 N NaHCO₃ using a dialysis membrane (Molecular Weight Cut-off 12,000–14,000, Wako Pure Chemical, Osaka, Japan). For N-acetylation and demineralization, the dialyzed, freeze-dried sample was combined with 250 μl of 0.2 M ammonium acetate (pH 6.7) and 25 μl of acetic anhydride at room temperature for 30 min. This step was repeated for another 30 min. With the gradual addition of 100 % ethanol, the sample was freeze-dried to eliminate both reagents, then dissolved in 200 μl of H₂O, demineralized using an acilyzer G0 (Asahi-kasei, Tokyo, Japan), and freeze-drying. The dried specimens were treated in 1 ml of 2 M trifluoroacetic acid at 100 °C for 2 h for acid hydrolysis. Subsequently, the hydrolyzed specimens were dissolved in 200 μl of H₂O. Ten microliters of the solution was freeze-dried, followed by complete dissolution in 100 μl of pyridine and then combined with 20 μl of N,O-bis(trimethylsilyl) trifluoroacetamide (BSTFA) and 2 μl of trimethylchlorosilane (TMCS) at 60 °C for 30 min for trimethylsilylation. The resultant specimens were subjected to GC/MS analysis (Clarus SQ8T, Perkin Elmer, Waltham, MA, USA). The mass spectra were compared with those of a standard sugar mixture (Glc, Gal, and GaIA: Wako Pure Chem.; GlcA: Sigma-Aldrich; GlcN: Yaizu Suisankagaku, Shizuoka, Japan).

**Fluorescent labeling to mask NH₂ in sheath skeletons.** The fluorescent protein, allophycocyanin (APC) conjugated reagent (Dojindo, Kumamoto, Japan) (APC-reagent) was used to confirm the presence of NH₂ in sheath skeletons of OUMS1-WT (Fig. 3b). In parallel, OUMS1-SL, which does not excrete exofibrils, was used as a negative control for exofibril absence. OUMS1-WT and OUMS1-SL, which had been precultured in SGP for 2 days, were separately mixed with this reagent at dilutions from 10⁻² to 10⁻³ and incubated at 37 °C for 1–12 h. After four washes in SGP, the specimens were observed microscopically. For monitoring their affinity for Fe(III) minerals, they were incubated with Fe(III)SGP for 2 days before observation. For XRF analysis, the Fe(III)SGP-incubated specimens were washed six times in H₂O and freeze-dried.

Two other reagents that can mask NH₂ were similarly tested: sulfo-NHS-acetate (1–10 mM, Tokyo Kasei, Tokyo, Japan) and acetic anhydride (200 mM, Nacalai Tesque), were used for masking NH₂ in the OUMS1-WT sheath skeleton. OUMS1-WT, precultured in SGP for 2 days, was incubated with either reagent at 37 °C for 1–12 h. After washing with SGP four times, the respective specimens were incubated in Fe(III)SGP for 2 days, then washed six times in H₂O, freeze-dried, and analyzed by XRF. For TEM/EDX imaging, the incubated specimens were chemically fixed as described below.
Reference experiments using polysaccharides and polystyrene beads. Chitosan (Wako Pure Chem.) was incubated in SGP or Fe(III) SGP at a final concentration of 4 mg ml\(^{-1}\) for 12–16 h to examine its affinity for Fe(III) minerals. For comparison, the affinity of cellulose (Nacalai Tesque) for the minerals was tested. After incubation in SGP or Fe(III) SGP, these polymers were washed at least six times in H\(_2\)O, freeze-dried, and subjected to XRF analysis, SEM/EDX imaging, and XPS measurement.

Additionally, three types of polystyrene beads (plain, NH\(_2\)-coated, and acetylated-NH\(_2\)-coated: 10 \(\mu\)m diameter each) (Micro mod, Rostock, Germany) were incubated in SGP or Fe(III) SGP at a final concentration of 2.5 mg ml\(^{-1}\). After incubation for 12–16 h, the beads were washed twice in H\(_2\)O to remove SGP or Fe(III) SGP, then washed four times in 100% ethanol and allowed to air-dry. Either ethanol-suspended or air-dried beads were subjected to the following XRF analysis, SEM/EDX imaging, and XPS measurement. The NH\(_2\) on acetylated-NH\(_2\)-coated beads is supposed to be blocked by the acetylation.

Differential interference contrast (DIC) and fluorescence microscopic imaging. OUMS1-WT and OUMS1-SL cells that had been treated with APC-reagent or for live/dead staining were observed with a BX51 system microscope (Olympus, Tokyo, Japan) equipped with DIC optics and a fluorescence attachment with a mercury lamp: U-MWIG3 U-MWU2, and U-MBIW3 (530–550, 330–360, and 460–490 nm excitation and 580, 430, and 520 nm emission, respectively) and dichroic mirror units.

Scanning and transmission electron microscopy. For SEM/EDX, the above-H\(_2\)O-washed specimens were fixed with a mixture of 2.5% v v\(^{-1}\) glutaraldehyde and 1% w v\(^{-1}\) OsO\(_4\) in 0.1 M caddycate buffer (pH 7.0), washed with H\(_2\)O, dehydrated with an increasing series of ethanol solution (30%, 50%, 70%, 95%, and 100%), t-butanol, and critical-point drying, then placed on carbon tape (Nisshin EM, Tokyo, Japan) on a stub. Freeze-dried sugar chains were similarly attached to carbon tape, as were ethanol-suspended polystyrene beads, then air-dried. Specimens were coated with platinum (ca 15 nm thick) using an ion-sputter (E-1030, Hitachi, Tokyo, Japan) and then observed with an SEM (S-4300, Hitachi) equipped with a dispersive X-ray spectrometer (EDX) (Genesis 2000, Ametek-Edax, Berwyn, PA, USA) at 15 kV.

For TEM, the above-treated specimens collected by centrifugation were fixed with a mixture of 2% v v\(^{-1}\) glutaraldehyde and 2% v v\(^{-1}\) paraformaldehyde in 0.1 M phosphate buffer (pH 7.4) at 4°C overnight. After a 30 min buffer wash, the specimens were embedded in 3% agar in the buffer. Small pieces of the agar block were post-fixed with 2% w v\(^{-1}\) OsO\(_4\) in 0.1 M phosphate buffer (pH 7.4) for 1.5 h, then washed with the buffer. Then, the specimens were dehydrated in an increasing series of ethanol solutions (30%, 50%, 70%, 95%, and 100%), treated with propylene oxide, then embedded in Spurr’s resin. Sections (70–80 nm thick) were cut using a ultramicrotome (Leica, Wetzlar, Germany) equipped with a diamond knife (Ultra 45° 3.0 mm, Diatome, Hartfield, PA, USA) and then observed with a TEM (JEM-2100F, JEOL, Tokyo, Japan) equipped with EDX at 200 kV. The semiqualitative distribution maps were calculated by the Clift-Lorimer method\(^{43,44}\) to confirm their location.

X-ray fluorescence (XRF) analysis. To determine the atomic percentage of Fe in the test specimens, freeze-dried OUMS1-WT sheaths and sugar polymers and air-dried polystyrene beads were packed into small aluminum pans for elemental analysis with an Orbis micro x-ray fluorescence (XRF) analyzer (Ametek, Berwyn, PA, USA). Atomic percentage of any detected element with a standard error was expressed as the mean (±SE) of 10 spots.

X-ray photoelectron spectroscopy (XPS). For obtaining photoelectron spectra from OUMS1-WT sheaths, sugar chain polymers and polystyrene beads, XPS measurements were carried out using monochromatic Al K\(_\alpha\) radiation (\(hv = 1486.6\) eV) as described previously\(^{45,46}\). Briefly, specimens were spread on carbon tape (Nisshin EM) and vacuum-dried. In the spectroscopy apparatus, surface charges on specimens were neutralized by using a low energy flood gun (~5 eV). Before evaluating the shifts in binding energy of the N1s in the respective specimens (Figs 5c, 6h and 7g), the detected N1s values in Fe(III)SGP-treated specimens (Figs 5b, 6g and 7f) were compensated on the basis of the value of Cs of the respective SGP-treated specimens (284.3 eV) (Figs 5a, 6e and 7e), because the binding energy levels of all elements varied slightly between Fe(III)SGP- and SGP-treated specimens. The photoelectron O1s spectrum acquired from all specimens was negligible, because it was apparently derived from contaminated carbonate within the x-ray photoelectron spectroscopy vacuum chamber of the XPS device.

X-ray diffraction (XRD) analysis. To examine the crystallinity of the Fe(III) minerals obtained from SGP + Fe plate or SGP + FeSO\(_4\), XRD patterns of ethanol-washed and dried Fe(III) minerals were analyzed using an RINT-2500HF X-ray generator (Rigaku, Tokyo, Japan) with Cu K\(_\alpha\) radiation (voltage: 40 kV; current: 200 mA) as described previously\(^{47}\). The freeze-dried specimens were fixed on a zero background sample holder and scanned continuously from 10° to 90° (2\(\theta\) value) at a rate of 3° min\(^{-1}\). The XRD pattern of the zero background sample holder was also measured.

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Author Contributions
T.K. designed the study, conducted most of the experiments, and wrote the manuscript; S.M. performed XPS measurement; N.N. performed TEM/EDX; S.K. and K.H. determined sugar composition; T.U., K.T., H.K., and J.T. developed the original concept of the project and/or provided technical advice.

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