Structure of Microbial Nanowires Reveals Stacked Hemes that Transport Electrons over Micrometers

Highlights

- *Geobacter* nanowires are made up of micrometer-long polymerization of cytochrome OmcS

- All hemes are closely stacked (<4–6 Å), providing a continuous path for electron flow

- We show that these are the same filaments that were earlier thought as type IV pili

- This structure explains the molecular basis for electron conduction in protein wires

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In Brief

Stacked heme filaments form the structural basis for long-range electron transport in bacterial nanowires.
Structure of Microbial Nanowires Reveals Stacked Hemes that Transport Electrons over Micrometers


INTRODUCTION

Conductive filamentous appendages of common soil bacteria Geobacter (Reguera et al., 2005), referred to as microbial nanowires, play a critical role in long-range extracellular electron transfer for respiration (Malvankar et al., 2011) and interspecies electron exchange (Summers et al., 2010). These nanowires have been invoked to explain a wide range of globally important redox phenomena that influence carbon and mineral cycling in soils and sediments, bioremediation, corrosion, and anaerobic conversion of organic wastes to methane or electricity (Malvankar and Lovley, 2014; Malvankar et al., 2011). However, these filaments’ composition, structure, and underlying conduction mechanism have remained uncertain because the filaments are difficult to solubilize for studies using traditional biochemical methods and X-ray crystallography. Geobacter sulfurreducens serves as a model organism for the broader phenomenon of extracellular electron transfer because it produces these conductive filaments (Tan et al., 2017) and has a fully sequenced genome and well-developed genetic system (Reguera et al., 2005). In contrast to other electron-transferring bacteria (Marsili et al., 2008), G. sulfurreducens does not use diffusing shuttle molecules but requires direct contact with an electron acceptor via conductive filaments for long-range extracellular electron transfer (Reguera et al., 2009).

G. sulfurreducens outer-surface c-type cytochromes, including the hexaheme cytochrome Omcs, with hemes packed within ~3.5–6 Å of each other. The inter-subunit interfaces show unique structural elements such as inter-subunit parallel-stacked hemes and axial coordination of heme by histidines from neighboring subunits. Wild-type Omcs filaments show 100-fold greater conductivity than other filaments from a Δomcs strain, highlighting the importance of Omcs to conductivity in these nanowires. This structure explains the remarkable capacity of soil bacteria to transport electrons to remote electron acceptors for respiration and energy sharing.

SUMMARY

Long-range (>10 μm) transport of electrons along networks of Geobacter sulfurreducens protein filaments, known as microbial nanowires, has been invoked to explain a wide range of globally important redox phenomena. These nanowires were previously thought to be type IV pili composed of PilA protein. Here, we report a 3.7 Å resolution cryoelectron microscopy structure, which surprisingly reveals that, rather than PilA, G. sulfurreducens nanowires are assembled by micrometer-long polymerization of the hexaheme cytochrome OmcS, with hemes packed within ~3.5–6 Å of each other. The inter-subunit interfaces show unique structural elements such as inter-subunit parallel-stacked hemes and axial coordination of heme by histidines from neighboring subunits. Wild-type Omcs filaments show 100-fold greater conductivity than other filaments from a Δomcs strain, highlighting the importance of Omcs to conductivity in these nanowires. This structure explains the remarkable capacity of soil bacteria to transport electrons to remote electron acceptors for respiration and energy sharing.

Article

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of PilA. Instead, the filament composition was inferred from indirect evidence, including the presence of PilA in biochemical analyses (Tan et al., 2016b), or from low-resolution imaging by atomic force microscopy (AFM) and negative-staining transmission electron microscopy, which suggested filament dimensions similar to type IV pili (Reguera et al., 2005).

On the other hand, alternative functional roles have been suggested for PilA (Reguera et al., 2007). PilA is implicated in the secretion of OmcS to the outer surface (Liu et al., 2018; Richter et al., 2012), and overexpression of PilA is accompanied by overproduction of OmcS and extracellular filaments (Leang et al., 2013; Summers et al., 2010). Here, we reconcile previous observations by directly identifying the composition and structure of extracellular appendages of wild-type G. sulfurreducens using cryo-electron microscopy (cryo-EM). These appendages are polymerized filaments of OmcS with unique structural features that also provide a molecular basis for understanding long-range electronic transport in proteins. Gel electrophoresis, mass spectrometry, AFM, and conductivity measurements show that these nanowires are the same filaments that were previously thought to be type IV pili. Data from our measurements are consistent with previous studies, and our results establish a previously unknown class of protein-based nanowires based on cytochrome polymerization.

RESULTS AND DISCUSSION

We grew WT cultures using anodes of microbial fuel cells as the sole electron acceptors (O’Brien and Malvankar, 2017). These growth conditions promote production of conductive biofilms and filaments as well as overexpression of PilA in comparison with growth on soluble electron acceptors such as fumarate. (Malvankar et al., 2011). Consistent with previous studies (Tan et al., 2016b), we confirmed the presence of both PilA and OmcS with expected molecular weights of ~6.5 kDa and ~45 kDa, respectively, in our filament preparations using polyacrylamide gel electrophoresis (SDS-PAGE), peptide mass spectrometry, and western immunoblotting (Figure S1; Method Details).

Cryo-EM images of filaments purified from the WT strain showed a sinusoidal morphology with a period of ~200 Å (Figure 1A). Averaged power spectra from multiple filaments show a meridional layer line at ~1/(47 Å) (Figure S4B), establishing that there are ~4.3 subunits per turn of a ~200 Å pitch 1-start helix (Figure 1B). Using the iterative helical real space reconstruction (IHRSR) approach (Egelman, 2000), we were able to reach a resolution where the handedness of α helices was clearly visible. The 1-start filament helix was left handed, with a rise per subunit of 46.7 Å and a rotation of ~83.1° (Figure 1B), substantially different than type IV pili that typically show a rise of ~10 Å and a right-handed helix (Wang et al., 2017). The tracing of the Cα backbone of the protein subunit at this resolution revealed that the asymmetric unit contained at least 380 residues, which contrasts with the 61 residues present in PilA (Reardon and Mueller, 2013; Reguera et al., 2005). Further, there was no apparent internal symmetry that would arise in the asymmetric unit if it contained multiple copies of identical chains. The NMR
structure of PilA (Reardon and Mueller, 2013) also failed to fit into the observed EM density map. Surprisingly, we found that there were six hemes per asymmetric unit, with the highest densities in the volume at the centers of these hemes, suggesting the presence of metal atoms. Given that the protein contains at least 380 residues and would therefore have a likely molecular weight between 40 and 50 kDa, we used SDS-PAGE and cut out the strongest band in the gel, which was at 45 kDa (Figure S1A), analyzing this band by mass spectrometry. Five proteins were identified, four of which had heme-binding motifs (CXXCH) and had masses between 45 and 49 kDa (Figure S1B). Three of these proteins (OmcS, OmcT, and GSU2501) had a very similar pattern of heme-binding motifs that approximately matched the initial Cα trace (Figure 2B), and their sequences could be easily aligned (Figure 2A). These three proteins had between 45% and 63% sequence identity with each other. Another protein found in mass spectrometry, OmcZ (Inoue et al., 2010), contained 30 additional residues compared to the other three, and thus its sequence could not be aligned to the others, and furthermore, the pattern of eight heme-binding motifs in OmcZ did not match that found in the map. The remaining protein, OmpJ, contains no heme (Afkar et al., 2005). Of the three possible candidates, only the OmcS sequence could be threaded through the map without any conflicts (Figures 2C, 2D, and S2). There are nine regions of OmcT, GSU2501, or both that prevented their sequences from being fit into the map (Figures 2A, highlighted in blue, and S2). In addition, the pattern of bulky amino acids clearly established that only the OmcS sequence was consistent with the map. For example, the elongated density from arginine (R) 256 is clearly seen in the map (Figure 2D). But in the sequences of OmcT and GSU2501, this residue would be a valine (Figure 2A, highlighted in purple) that will not be able to explain the map density. These studies show that the only cytochrome, found in our filament preparations by mass spectrometry, that is consistent with the cryo-EM map is OmcS.

We built an ab initio atomic model of OmcS, which was then used to refine a filament model. We directly estimated its resolution using a map:model comparison (Neumann et al., 2018; Subramaniam et al., 2016), which yielded an estimate of 3.7 Å (Figure S3A). Strikingly, the “gold standard” measure of resolution employed by Relion (Scheres, 2012), which is based upon reproducibility, yielded an estimate of 3.2 Å. Comparing the Relion map (filtered to 3.2 Å) and our map generated in Spider (filtered to 3.6 Å) showed no significant differences (Figures S3B and S3D), suggesting that the Relion estimate of resolution is overly optimistic. Overall, the fit of the model to the map (Figures S3E and S3F), the good density for many side chains, and the refinement statistics (Table 1) all validate our model of the OmcS filament.

While it has been known for more than half a century that cytochromes can polymerize in ethanolic solutions (Margoliash and Lustgarten, 1962) and structures have been determined for aggregates up to tetramers (Hirotá et al., 2010), natural
polymerization of the type we observe here has not been previously described to our knowledge. Stacking arrangements of aromatic rings generally prefer parallel (offset face-to-face) or perpendicular (T-shaped) conformations (Janiak, 2000). The parallel stacking yields the highest electronic coupling, which maximizes electron transfer (Jiang et al., 2017), whereas the T-shape enhances structural stability (Janiak, 2000). Hemes in the OmcS nanowires form parallel-stacked pairs, with each pair perpendicular to the next, forming a continuous chain over the entire length of the filament (Figure 1D). The minimum edge-to-edge distances is 3.4–4.1 Å between the parallel hemes and 5.4–6.1 Å between the perpendicular stacked pairs (Figure 1D). For all hemes in OmcS nanowires, two histidines axially coordinate iron at the center of each heme, and the vinyl groups of each heme form covalent thioether bonds with cysteines (Figure 3). The bis-histidine axial ligation of the heme iron atoms are consistent with the coordination found in other multi-heme c-type cytochromes (Clarke et al., 2011), and the cysteine linkages are consistent with the c-type hemes reported to occupy six heme-binding motifs of OmcS (Qian et al., 2011).

Our OmcS model filament has a low percentage of α and 3_10 helices (~13%) as well as β strands (~8%), leaving ~81% of the model as turns and coil, which is consistent with previous secondary-structure studies of OmcS (Qian et al., 2011). We compared the OmcS protomer within the filament with a group of three crystallographic structures of other multi-heme c-type cytochromes (PDB: 1OFW, 3UCP, and 3OV0). These structures showed 45%, 49%, and 60% turns and coils, respectively, with uniformly hydrophobic cores surrounding the hemes and heme-binding residues. Cores in our model of OmcS also included buried charges (arginine at locations 333, 344, and 375) that lack proximal compensating charges, as well as buried side-chain hydroxyls (tyrosine at locations 186, 231, and 385). In addition, our model of OmcS has a salt bridge between protein chains, aspartate 407 to arginine 151, which along with cysteines in heme binding motifs, are highly conserved amino acids in OmcS (Ashkenazy et al., 2016). The lack of structural homology seen with these other c-type cytochromes is consistent with the observation that there is no conserved fold for this family of proteins (Bertini et al., 2006).

The model shows that each OmcS subunit contacts only one subunit on either side, so that all connectivity in the filament is along the left-handed 1-start helix (Figure 1B). The interface between adjacent subunits is extensive, with ~2,600 Å² of surface area buried per subunit (Figure 3A). In addition to the buried surface area, interactions between adjacent subunits incorporate additional stabilizing elements unique to the filament structure. Histidine 16 in each subunit coordinates the iron in heme 5 of an adjacent subunit (Figure 3B). Furthermore, heme pairs at the interface are parallel rather than T-shaped, with ~4 Å edge-to-edge distance (Figures 1D, 3B, and 3C). This parallel stacking and inter-subunit coordination of heme may contribute substantially to the stability of the protein-protein interface. In addition, the presence of parallel-stacked hemes at the interface suggests facile transport of electrons between monomers.

There is no precedent for such seamless micrometer-long polymerization of hundreds of cytochromes to our knowledge. Based on previous studies that have shown that cytochromes could form tetramers, the OmcS polymerization could be due to successive domain swapping, where the c-terminal helix can be displaced from its original position in the monomer and histidine-heme coordination can be perturbed significantly (Hirota et al., 2010). We have determined the structure for the OmcS protomer within the filament, but the structure of isolated OmcS monomer is needed to provide insight into this surprising polymerization process.

The atomic structure was solved with filaments from conductive biofilms of electrode-grown cells that require long-distance electron transport. However, fumarate-grown cells can also produce conductive filaments (Ing et al., 2017; Leang et al., 2010, 2013; Malvankar et al., 2011; Reguera et al., 2005). Filaments purified with fumarate-grown cells showed similar structure to filaments purified from electrode-grown cells (Figures S4A and S4B). Moreover, previously published images of intact G. sulfurreducens filaments attached to cells (Leang et al., 2013) showed structural features similar to purified OmcS filaments such as an identical helical rise of 47 Å (Figure S4C). These results showed that purified OmcS filaments are similar in dimensions and structure to cell-attached filaments that were previously thought to be type IV pili (Leang et al., 2013). Importantly, these studies also indicate that the formation of OmcS filaments is a natural process and not due to artificial preparation or pH conditions that can cause cytochrome c to form filamentous structures under extremely denaturing conditions (Haldar et al., 2015).

The cryo-EM images also showed another filamentous structure that was thinner than the OmcS filament (Figure S5). The averaged power spectrum of these filaments showed similar layer lines to the OmcS filament but with a slightly different axial rise of ~57 Å and rotation of ~160°, suggesting that this thinner filament is also not a type IV pilus and could potentially be another cytochrome filament. Due to lower abundance of this filament in our cryo-EM images, it was not possible to build an
atomic model or to determine its composition. No filaments with power spectra consistent with type IV pili were observed in purified filaments preparations or in previously published images of intact, cell-attached filaments (Leang et al., 2013).

To evaluate the contribution of OmcS to the conductivity of these filaments, the direct current (DC) conductivity of individual OmcS filaments of WT strain was compared with sparse filaments produced by a \( \Delta \text{omcS} \) strain (Figure 4B). \( \text{G. sulfurreducens} \) forms a variety of filaments in response to genetic mutations (Klimes et al., 2010). Therefore, it is not surprising that the \( \Delta \text{omcS} \) strain also forms filaments that were previously thought to be conductive type IV pili (Leang et al., 2010). AFM (Figures 4A–4E) revealed distinct structural features for WT \( \text{G. sulfurreducens} \) filaments versus the type IV pili of other species (Wang et al., 2017) and filaments from the \( \Delta \text{omcS} \) strain (Leang et al., 2010). In contrast to the linear and smooth-surfaced structure of \( \Delta \text{omcS} \) filaments, WT OmcS filaments exhibited an axial periodicity with a 20 nm pitch (Figure 4E), consistent with the helical pitch determined by cryo-EM (Figure 1B). Filaments of the \( \Delta \text{omcS} \) strain showed no apparent axial periodicity under AFM. Moreover, \( \Delta \text{omcS} \) filament thickness (\( \sim 1.7 \) nm height measured by AFM) was half that of WT OmcS filaments (\( \sim 4 \) nm) (Figures 4D and 4E). This substantial thickness difference and distinct axial periodicity observed for OmcS filaments was used to confirm that OmcS filaments studied for electrical measurements are the same OmcS nanowires characterized by cryo-EM.

AFM was further used to locate an individual filament bridging two gold electrodes (Figure 4F, inset). Our DC-conductivity measurements of individual OmcS filaments fully hydrated in buffer yielded values (Figures 4F and 4G) comparable to previous measurements of WT filaments (Adhikari et al., 2016), further suggesting that OmcS filaments are similar to the WT nanowires discussed in previous studies as type IV pili. DC-conductivity measurements of individual \( \Delta \text{omcS} \) filaments showed a very low conductivity that was more than 100-fold lower than OmcS filaments (Figures 4F and 4G). Our conductivity measurements thus show that OmcS is required for the high conductivity of these \( \sim 4 \) nm-thick filaments.

Our finding that WT \( \text{G. sulfurreducens} \) nanowires are OmcS filaments is consistent with previous physiological studies. These studies highlight the importance of OmcS in extracellular electron transfer (Holmes et al., 2006; Leang et al., 2010, 2013; Mehta et al., 2005; Summers et al., 2010). OmcS is one of the most abundant cytochromes found in the proteome for electricity-producing \( \text{G. sulfurreducens} \) and is required only during extracellular electron transfer to insoluble electron acceptors such as Fe (III) oxide (Holmes et al., 2006; Mehta et al., 2005). It is also critical for direct interspecies electron transfer between syntrophic \( \text{Geobacter} \) co-cultures as evolved co-culture overexpressed OmcS and deletion of the \( \text{omcS} \) gene inhibited bacterial ability to exchange electrons (Summers et al., 2010).

OmcS also plays a critical role in electron transport to electrodes in current-producing biofilms. Both microarray and quantitative reverse transcription polymerase chain reaction (qRT-PCR) analysis have demonstrated that cells show greatest increase in transcript levels for OmcS during the early stages of growth on electrodes (Holmes et al., 2006). Furthermore, immunogold localization has shown that OmcS is distributed throughout conductive \( \text{G. sulfurreducens} \) biofilms (Leang et al., 2013) and that deletion of the \( \text{omcS} \) gene inhibits their
production of electricity under some conditions (Holmes et al., 2006). However, the role of Omcs in conductivity of nanowires was overlooked because ∆omcs biofilms were conductive and produced high current densities (Malvankar et al., 2011). These biofilm results might be due to a reciprocal relationship (Park and Kim, 2011) between the expression of Omcs and that of Omcz, a cytochrome essential for current production (Nevin et al., 2009). The ∆omcs biofilms may compensate for the loss of Omcs by increasing the production of Omcz.

Using immunogold localization, previous studies found that Omcs is associated with filaments (Leang et al., 2010, 2013; Summers et al., 2010). As cytochromes were not known to form filaments before our work, AFM images of filaments (Malvankar et al., 2012) as well as these antibody-labeling results (Leang et al., 2010, 2013; Summers et al., 2010) were interpreted as showing isolated Omcs monomers binding to the surfaces of the PilA filaments rather than showing antibodies directly binding to Omcs filaments. In light of the result presented here, a reinterpretation of these previous studies suggests that the antibodies may have been directly binding to the subunits of the Omcs filaments.

While no evidence of PilA was found in the structure of any of the purified filaments, non-filamentous PilA was present in our samples in lower abundance compared to Omcs (Figure S1A), consistent with prior studies (Tan et al., 2016b). Multiple studies have shown that PilA is required for secretion of Omcs to the extracellular environment, as pilA deletion eliminated the presence of Omcs in outer-surface preparations (Liu et al., 2018; Richter et al., 2012). Overexpression of PilA is also accompanied by overproduction of Omcs and filaments (Leang et al., 2013; Summers et al., 2010), further suggesting that PilA is involved in secretion of Omcs filaments that may explain previous correlations found between PilA and biofilm conductivity (Malvankar et al., 2011). The requirement of PilA for the synthesis of Omcs filaments thus also explains the inability of ∆pilA cells to grow on insoluble electron acceptors such as Fe (III) oxides (Reguera et al., 2005) and electrodes (Reguera et al., 2006). A number of other bacteria have also been shown to require non-filamentous type IV pilins for the secretion of extracellular proteins (Hager et al., 2006). One possibility is that PilA is a pseudopilin as a part of a type 2 secretion system (T2SS) (Nivaskumar and Francetic, 2014) and that Omcs is exported by this T2SS. Previous studies have shown that T2SS pseudopilin can be secreted outside the cell (Nivaskumar and Francetic, 2014; Vignon et al., 2003). A similar mechanism could explain the presence of non-filamentous PilA in our filament preparations.

This potential ability of PilA to regulate the secretion and assembly of the Omcs nanowires, as well as that of other multiheme cytochromes (Liu et al., 2018; Richter et al., 2012), could explain how point mutations in pilA caused cells to produce filaments with different conductivities than WT filaments. Filaments produced by cells with point mutations in pilA showed very different morphology than WT filaments (Tan et al., 2016a). For example, substitution of two residues in PilA with tryptophan yielded mutant cells with filaments that surprisingly had half the diameter of WT filaments (Tan et al., 2016a). Based on our finding that conductive filaments are composed of cytochromes, this very large structural change in the filaments of mutant strains suggests that the observed change in conductivity could be due to filaments of different cytochromes or of different conformations of Omcs. Further structural studies on the filaments produced by these pilA mutants are needed to fully reconcile these studies with our finding of Omcs filaments functioning as nanowires.
In summary, our findings show that conductive G. sulfurreducens filaments are polymerized chains of OmcS. The filament structure has hemes closely stacked along the micrometer length of the filament, establishing the molecular basis for electronic conductivity in these nanowires. Functional characterization of conductivity in individual filaments shows that OmcS is required for the filament conductivity. The structure presented here provides insights into supramolecular protein nanowires, explaining the remarkable capacity of soil bacteria to transport electrons to extracellular electron acceptors for respiration (Malvankar et al., 2011) and for sharing of energy and nutrients with syntrophic partners (Summers et al., 2010) that are hundreds of micrometers away. The advances in understanding of the structural basis for conductivity in microbial nanowires presented here can provide design principles for development of future bioelectronic interfaces between living cells and devices.

**STAR METHODS**

Detailed methods are provided in the online version of this paper and include the following:

- **KEY RESOURCES TABLE**
- **CONTACT FOR REAGENT AND RESOURCE SHARING**
- **EXPERIMENTAL MODEL AND SUBJECT DETAILS**
  - **Bacterial strains and growth conditions**
- **METHODS DETAILS**
  - **G. sulfurreducens filament preparation and biochemical characterization**
  - Filament preparation from fumarate-grown cells (Figure S4A)
  - Cryo-EM sample preparation conditions
  - Cryo-EM data collection conditions
  - Cryo-EM image analysis
  - Model building of OmcS filaments
  - Atomic force microscopy
  - Direct current conductivity measurements
  - Conductivity calculations
- **QUANTIFICATION AND STATISTICAL ANALYSIS**
- **DATA AND SOFTWARE AVAILABILITY**

**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found with this article online at https://doi.org/10.1016/j.cell.2019.03.029.

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**AUTHOR CONTRIBUTIONS**

F.W., under the supervision of E.H.E., performed the image analysis, reconstructed the OmcS filament structure, generated, refined the filament model, and imaged and biochemically analyzed filaments from fumarate-grown cells that yielded similar helical parameters as filaments from electrode-grown cells used to build the atomic model. Y.G. prepared and optimized cryo-EM grids, collected data used to build the atomic model, and carried out AFM imaging on wild-type filaments. J.P.O., V.S., and C.S. grew cells for studies on cell-attached filaments and purified filaments used for building the atomic model and for functional studies. Y.G. and V.S. identified cell-attached and purified filaments by negative stain electron microscopy. S.M.Y. carried out biochemical analysis and, with D.V., performed AFM and conductivity measurements of filaments. S.E.Y. performed AFM imaging of cell-attached filaments of the ΔomcS strain, compared their height profiles with filaments of WT strain, and interpreted morphological differences. D.V., Y.G., and C.S. performed biochemically analyzed filaments from fumarate-grown cells. N.S.M. supervised the sample preparation, characterization, and data collection used to build the atomic model; conceived and designed functional studies; and wrote the manuscript with input from all authors.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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**REFERENCES**


## STAR METHODS

### KEY RESOURCES TABLE

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Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Nikhil S. Malvankar (Nikhil.Malvankar@yale.edu).

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Bacterial strains and growth conditions**

The *Geo bacter sulfurreducens* wild-type (WT) strain PCA (designated DL-1) (ATCC 51573, DSMZ 12127) (Coppi et al., 2001) and the omcS deletion mutant strain (Mehta et al., 2005) (designated ΔomcS) were obtained from our laboratory culture collection. The cultures were maintained at 30°C or at 25°C under strictly anaerobic conditions in growth medium supplemented with acetate (10 mM) as the electron donor and fumarate (40 mM) as the electron acceptor in sterilized and degassed NBAF medium (O’Brien and Malvankar, 2017). 1L NBAF medium contained the following: 0.04 g/L calcium chloride dihydrate, 0.1 g/L magnesium sulfate heptahydrate, 1.8 g/L sodium bicarbonate, 0.42 g/L potassium phosphate monobasic, 0.22 g/L potassium phosphate dibasic, 0.2 g/L ammonium chloride, 0.38 g/L potassium chloride, 0.36 g/L sodium chloride, vitamins and minerals as listed in (O’Brien and Malvankar, 2017). Resazurin was omitted and 1 mM cysteine was added as an electron scavenger. All chemicals obtained from Fisher Scientific unless otherwise noted. For filament samples used to build the atomic model, the wild-type strain was grown on electrodes under electron acceptor-limiting conditions that induce filament expression (O’Brien and Malvankar, 2017) (Figure 1A).

**METHODS DETAILS**

**G. sulfurreducens filament preparation and biochemical characterization**

Filaments were separated from bacteria and extracted via centrifugation (Tan et al., 2016a) and maintained in 150 mM ethanolamine buffer at pH 10.5 in a manner similar to structural studies on bacterial filaments (Wang et al., 2017). Cells were gently scraped from the surface using a plastic spatula and isotonic wash buffer (20.02 × 10⁻³ M morpholinepropanesulfonic acid, 4.35 × 10⁻³ M NaH₂PO₄, 1.34 × 10⁻³ M KCl, 85.56 × 10⁻³ M NaCl, 1.22 × 10⁻³ M MgSO₄·7H₂O, and 0.07 × 10⁻³ M CaCl₂·2H₂O), then collected by centrifugation and re-suspended in 150 × 10⁻³ M ethanolamine (pH 10.5). Filaments were mechanically sheared from the cell surface using a Waring Commercial Blender (Cat. No. 7011S) at low speed for 1 min, and then cells were removed by centrifugation at 13,000 g before collecting filaments with an overnight 10% ammonium sulfate precipitation and subsequent centrifugation at 13,000 g. Collected filaments were re-suspended in ethanolamine buffer then cleaned by centrifugation at 23,000 g to remove debris and a second 10% ammonium sulfate precipitation with centrifugation at 13,000 g (Tan et al., 2016a). The final filament preparation was re-suspended in 200 μL ethanolamine buffer. Filament preparations were further passed through 0.2 μm filters to remove any residual cells and stored at 4°C. Cell-free filament preparations were imaged first with transmission electron microscopy to ensure sample quality (Figure 1A). Dilute 5 μL solutions containing filaments were placed on gold electrodes to achieve individual filaments across two gold electrodes (Figure 4F, Inset). Prior to all measurements, filaments were imaged with AFM and height measurements were performed to confirm the presence of individual filaments. For all conductivity measurements, samples were maintained under hydrated buffer environments (150 mM ethanolamine) and the pH of the buffer was equilibrated to pH 7 using HCl (Malvankar et al., 2011). Transmission electron microscopy imaging was used to confirm that filaments maintain their structure and remain morphologically similar at all pH conditions.

For polyacrylamide gel electrophoretic separation (SDS-PAGE) (Figure S1A), all filament samples were prepared in ethanolamine buffer and dried in a speedvac. Samples were then resuspended in 500 μL ultrapure deionized water and centrifuged at 18,000 x g at 4°C for 1 hour. The pellets were dried again in the speedvac, resuspended in water and dried repetitively 2 times. The final dried samples were resuspended in 5-8 μL 2.5% sodium-dodecyl sulfate (SDS) to disassemble filaments into their constituent monomers. The samples were incubated at room temperature for at least 4 hours, then diluted 3-fold with deionized water. The denatured samples were boiled in 1X SDS sample buffer that included b-mercaptoethanol for 12 min. The samples were run on a 16% Tricine protein gel (ThermoFisher Scientific, Carlsbad, CA) initially at constant voltage of 30 V for 18 min before changing to 190 V for 12 min. Precision Plus Protein Prestained molecular weight standards (BioRad, Hercules, CA) and Low Range Protein Ladder (Thermo Scientific) were used to compare the molecular weight of PilA and cytochromes in the filament preparations. Gels were immediately washed at least 3 times with ultra-pure deionized water over a 1-hour period, stained with Coomassie R-250 stain (Thermo Scientific, Rockford, IL), and destained overnight.

Custom monospecific anti-PilA antibody was synthesized by Pacific Immunology (Ramona, CA) by immunizing two rabbits with synthetic peptide sequence containing targeted epitope on the native protein, PilA, and then affinity purifying the serum against that peptide sequence. Specificity of antibody in the serum was confirmed by ELISA after 1st stage of immune response and then verified again with purified antibody (1:125,000 titer) at the final stage. Synthetic peptide sequence contained following 21 amino acids from C terminus of PilA, Cys-RNLKTALESAFADDQTYPPES, to which a cysteine was added to N terminus and then was conjugated with Keyhole Limpet Hemocyanin prior to immunization. The molecular weight of the synthetic peptide was verified by HPLC and ESI-MS.
For LC-MS/MS analysis of filaments (Figure S1B), the PilA band (~6.5 kDa) and OmcS bands (~50 kDa and 30 kDa) were extracted from the protein gel and treated with trypsin to digest the protein (Figure S1). Proteomic analysis of the cleaved peptides from filaments of electrode-grown cells was performed by the Proteomics Mass Spectrometry Facilities at Yale University. Results gave unique amino acid sequence matches with the OmcS and C-terminal domain of PilA (Figure S1) whereas the hydrophobicity of the N-terminal domain may have interfered with the sequence detection procedure (Ing et al., 2017).

Filament preparation from fumarate-grown cells (Figure S4A)

G. sulfurreducens cells grown in fumarate were used for filament purification (Ing et al., 2017). Briefly, G. sulfurreducens (DSMZ strain 12127) was grown anaerobically in 1 L cultures of sterilized and degassed NBAF medium (0.04 g/L calcium chloride dihydrate, 0.1 g/L magnesium sulfate heptahydrate, 1.8 g/L sodium bicarbonate, 0.5 g/L sodium carbonate, 0.42 g/L potassium phosphate monobasic, 0.22 g/L potassium phosphate dibasic, 0.2 g/L ammonium chloride, 0.38 g/L potassium chloride, 0.36 g/L sodium chloride, vitamins and minerals, using 20 mM acetate as the electron donor and 40 mM fumarate as the electron acceptor). Resazurin was omitted and 1 mM cysteine was added as an electron scavenger. All chemicals were obtained from Fisher Scientific unless otherwise noted.

Upon reaching stationary phase, cells were pelleted by centrifugation at 5000 x g for 20 min at 4°C. Cell pellets were removed and extracellular filaments were purified from the supernatant through dropwise additions of 1 M aqueous MgCl2, which was added to a final ratio of 1:10 for aqueous MgCl2:supernatant. Supernatants were stored at 4°C overnight and centrifuged at 50,000 x g for 1 h at 4°C to precipitate extracellular filaments. Pellets of extracellular filaments were resuspended with ultrapure water adjusted to pH 4.3 with HCl and passed through 0.2 μm filters. Additional contaminants were removed from the resuspended pellets using 50 kDa dialysis tubing. After two 1 L exchanges of pH 4.3 water, the dialyzed sample was removed from dialysis tubing and stored at 4°C.

For analysis of filaments from fumarate-grown cells (Figure S4A), the purified filament samples were analyzed for protein content by 12.5% PhastGel (GE Healthcare) and stained with Coomassie. There were two prominent protein bands around 47 kDa, and these were cut from the gel, digested by trypsin overnight, and then analyzed by mass spec at the University of Virginia core facility. The LC-MS system consisted of a Thermo Electron Q Exactive HFX mass spectrometer system with an Easy Spray ion source connected to a MS system. An acetonitrile/0.1 M formic acid gradient at a flow rate of 0.3 mL/min over 2.0 hours. The nanospray ion source was operated at 1.9 kV. The digest was analyzed using the rapid switching capability of the instrument acquiring a full scan mass spectrum to determine peptide molecular weights followed by product ion spectrums (10 HCD) to determine amino acid sequence in sequential scans. This mode of analysis produces approximately 60,000 MS/MS spectra of ions ranging in abundance over several orders of magnitude.

Prior to use, purified filaments were characterized by MALDI-TOF. For MALDI-TOF analysis, filaments were denatured with 10% OG overnight and then diluted with ultrapure water to a final OG concentration of 2% prior to the addition of matrix solution (α-Cyano-4-hydroxycinnamic acid dissolved in a 2:1 solution of ultrapure water; acetonitrile and 0.2% trifluoroacetic acid). The final sample for MALDI was a 12:7:5 ratio of matrix solution to ultrapure water to diluted and denatured protein. Mass spectra were collected in positive ion mode.

Cryo-EM sample preparation conditions

For samples used to build the atomic model, holey carbon coated Quantifoil grids (R 2/2 m/2 μm) were used. Prior to use, the grids were floated in 0.05% Triton 100X solution (Cheung et al., 2013). Cryo-EM specimens were prepared with a FEI Vitrobot Mark IV at 22°C with 100% humidity. 3 μL of sample solution containing WT filaments were dropped on the grids and spread gently by pipette tip before loading to the Vitrobot, blotted for 5.5 s and plunge-frozen in liquid ethane.

Cryo-EM data collection conditions

Micrographs were acquired by FEI Titan Krios electron microscopy performed at 300kV equipped with a Gatan K2 summit camera. Quantum energy filter with a slit width of 20 eV was applied to remove inelastically scattered electrons. Movies were collected using super-resolution imaging mode with a physical pixel size of 1.05 Å and an exposure rate of 6.8 electrons per pixel per second. A total exposure time of 9.75 s was fractioned into 30 frames and 1447 movies were generated using serial EM auto-collection. The data collection and processing parameters are summarized below:

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For analysis of filaments from fumarate-grown cells (Figure S4A), sample (4 µL, ~1 µg/µl) was applied to discharged lacey carbon grids and plunge-frozen using a Vitrobot Mark IV (FEI, Inc.). Grids were imaged in a Titan Krios at 300 keV and recorded with a Falcon III direct electron detector at 1.4 Å per pixel. Micrographs were collected using a defocus range of 1.25–2.25 µm, with a total exposure of 2 s (amounting to ~60 electrons/Å²) distributed into 24 fractions. All the micrographs were first motion corrected (ignoring the first fraction) using MotionCorr (Li et al., 2013) version 2 and then used for CTF estimation by the CTFFIND3 program (Mindell and Grigorieff, 2003).

Cryo-EM image analysis

For filaments of fumarate-grown cells (Figure S4A), images were extracted using the e2helixboxer program within EMAN2 (Tang et al., 2007) from the dose-weighted fractions 2-10 (amounting to ~20 electrons/Å²), after the images were corrected for the CTF through multiplication by the theoretical CTF (a Wiener filter in the limit of a very poor SNR). A total of 17,800 overlapping 384-px long segments (with a shift of 55 pixels, ~1.5 times the axial rise per subunit) were generated. The determination of the helical symmetry was unambiguous given the 1/(47 Å) meridional layer-line and the 1-start helix of ~20 Å pitch. A reconstruction was generated using the IHRSR method implemented in Spider (Egelman, 2000), and this reconstruction was subsequently filtered to 20 Å for the starting reference in the reconstruction of the second dataset. To build the atomic model of filaments from electrode-grown cells, a total of 573 images were selected, motion-corrected, dose weighted (amounting to ~20 electrons/Å²) and CTF-corrected in the same manner as for filaments of electrode-grown cells. About 2,000 long filaments were extracted using e2helixboxer and then 384-pixel long 28,293 overlapping segments with a shift of 60 pixels were generated. The final volume from the IHRSR reconstruction was estimated to have a resolution of ~3.7 Å based on the model:map FSC (Figure S3). It was filtered to 3.6 Å and sharpened with a negative B-factor of 100. Micrographs with all fractions and boxing coordinates were also imported into Relion for 3D reconstruction. A similar final reconstruction to the one from SPIDER was generated in Relion using the same helical symmetry, starting with the SPIDER volume filtered to 10 Å. Interestingly this reconstruction was estimated to have a resolution of 3.2 Å resolution based on a “gold standard” map:map FSC.

Model building of Omcs filaments

First, about 400 protein C atoms and six heme cofactors corresponding to a single subunit within the filaments were built manually (Figure 2B) in Coot (Emsley and Cowtan, 2004). Then the density corresponding to a single subunit was segmented from the experimental filament density using Chimera (Pettersen et al., 2004). Using the proteins identified in the MS/MS results and real space fitting to the cryo-EM density, the sequence of the filament was unambiguously determined to be cytochrome Omcs (Figures 2 and S2). The full length Omcs protein as well as six heme molecules were then built manually in Coot. Then the Omcs/heme was rebuilt with the RosettaCM protocol (Wang et al., 2015) to remove bad geometries. A total of 3,000 full-length models were generated and the top 20 models were selected based on Rosetta’s energy function, Ramachandran plots and overall fit to the map. These 20 models were then combined into one model by manual editing in Coot using the criteria of the local fit to the density map and the geometry statistics of the model. To better refine heme interacting areas at this resolution, bond/angle restraints for the heme molecule itself, His-Fe, and Cys-heme thioester bonds were created based on high resolution cytochrome c3 crystal structures Nrb (PDB 2PDB) and NrbHA (PDB 2J7A). Then a filament model was generated and further refined using Phenix with additional heme area restraints, and MolProbit (Williams et al., 2018) was used to evaluate the quality of the filament model. The refinement statistics are given in Table 1.

Atomic force microscopy

To visualize individual filaments (Figure 4), 5 µl of buffer solution containing filaments were deposited on mica or on a silicon wafer insulated by a 100 nm silicon dioxide dielectric layer with gold electrodes patterned by e-beam or nanoimprint lithography (Tan et al., 2016a). For nanoimprinting, the substrate was cleaned with a Piranha solution (H2SO4:H2O2 = 3:1) and a diluted Hydrofluoric acid solution before patterning. Two layers of resist (50-nm-thick poly methyl methacrylate and 60-nm-thick UV-curable resist) were then sequentially spin-coated onto the cleaned substrate. Circuit patterns including nanoelectrodes separated by nano-sized gaps, microscale fan outs, and contact pads were transferred from a quartz mold to the UV resist with nanoimprint lithography in a homemade imprint chamber. The residual UV-resist layer and the poly methyl methacrylate underlayer were removed by reactive ion etching with fluorine based (CHF3/O2) and oxygen-based gases respectively. The excess buffer was absorbed with filter paper. The sample was air-dried and was mounted on a metal puck (Oxford Instrument, Cypher ES). Atomic force microscopy (AFM) experiments were performed using soft cantilevers (ASYELEC-01, Oxford Instrument Co.) with a nominal force constant of 2 N/m and resonance frequencies of 70 Hz. The free-air amplitude of the tip was calibrated with the Asylum Research software and the spring constant was captured by the thermal vibration method. The sample was imaged with a Cypher ES scanner using intermittent tapping (AC-air topography) mode. AFM showed that gold electrodes were bridged with individual filaments to facilitate conductivity measurements (Figure 4F, inset). To visualize helical features of filaments, AFM was operated in attractive force imaging mode and both phase and height channel images were reported to visualize axial periodicity (Figure 4A). All lateral and axial height analyses for filaments (Figure 4) were performed using Gwyddion and IGOR Pro software (WaveMetrics Inc).
Direct current conductivity measurements
All direct current (DC) conductivity measurements of filaments were performed under fully hydrated buffer conditions in a 2-electrode configuration inside a triaxially shielded dark box using a probe station (MPI Corp.) connected to a semiconductor parameter analyzer (Keithley 4200A-SCS) equipped with preamplifiers, allowing 0.1 fA current resolution and 0.5 μV voltage resolution. A DC voltage, typically in the range of −0.5V to +0.5V, was applied between the two electrodes and the current was measured over a minimum period of 120 s until the steady state was reached (Figure 4F). The linearity of the I-V characteristics was maintained by applying an appropriate low voltage and the slope of the I-V curve was used to determine the conductance (G). Measurements were performed at low voltages (< 0.5 V) and over longer times (> 100 s) to ensure a lack of electrochemical leakage currents or faradic currents as evidenced by the absence of significant DC conductivity in buffer or filaments of ΔomcS strain that were maintained under identical buffer conditions as filaments of the WT strain (Figures 4F and 4G). All analysis was performed using IGOR Pro (WaveMetrics Inc.).

Conductivity calculations
The conductivity (σ) of filaments was calculated using the relation (Malvankar et al., 2011) \( \sigma = G \cdot (L/A) \) where G is the conductance, L is the length of the filament, and \( A = \pi r^2 \) is the area of cross section of filament with 2r as the height of the filament measured by AFM (Figures 4F and 4G).

QUANTIFICATION AND STATISTICAL ANALYSIS
Quantification and statistical analyses employed in this publication pertain to the analysis on atomic force microscopy and electron microscopy data and the determination of structures by electron microscopy, which are integral parts of existing algorithms and software used.

DATA AND SOFTWARE AVAILABILITY
All data for OmcS filament were deposited in EMDB and PDB with the following entry codes: EMDB: EMD-9046 and PDB: 6EF8.
Supplemental Figures

Figure S1. Biochemical Characterizations of Filament Preparations Show PilA and Omcs Proteins, Related to Figures 1 and 2

(A) SDS-PAGE gel of filaments showing Omcs and PilA at expected molecular weights of ~45 kDa and ~6.5 kDa. LC-MS/MS analysis of metalloproteins in purified G. sulfurreducens filaments from (B) the ~45 kDa band and (C) the ~30 kDa band. EMPAI: Exponentially modified protein abundance index. LC-MS/MS analysis showing matched peptides in (D) Omcs in the ~45 kDa band and (E) PilA in the ~6.5 kDa band. (F) Western immunoblotting using anti-PilA antibody showing PilA in purified filament preparations (left) and cell lysate (right).

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Matched peptides in Omcs shown in bold:

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51. SATAQFTTGP MLLQGATQSS SCLNCQHAG DTGPSYSIH TAEADMPAGT
101. APLQMTPGGD FGWVKKTYTW NVRGNTSEG ERKGHNIVAG DYNVADTTL
151. TTAPGGTYPN NQHCSCHD PHGYRRFVD GIATTGLPI KNSGSYQNSN
201. DPTAWGAVGA YRILGGTYQ PKSLSGSYAF ANQVPAAVAP STRNTEATT
251. QTRVAYGQGM SEWCAHCNTD IHNSAYPTNL RHPANGKAFL GATIAGLYNS
301. YKSGDLTGT QASAYLSLAP FEEGTADTV LGKHAADDDT ALTGADATSN
351. VNCLSCRRAH ASGDFSMTRF NLAYEFTTIA DASGNSIYGT DPNTSSLQGR
401. SVNEMTAAYY GRTADKFAPY QRALCNKCHA KD Filament Cell

Matched peptides in PilA shown in bold:

1. FTIELLIVVAILIGLAIAIPQFSAYRK
31. AYNSAASSDLRNKLTALESFAADDQTYPDES

Figure S1. Biochemical Characterizations of Filament Preparations Show PilA and Omcs Proteins, Related to Figures 1 and 2

(A) SDS-PAGE gel of filaments showing Omcs and PilA at expected molecular weights of ~45 kDa and ~6.5 kDa. LC-MS/MS analysis of metalloproteins in purified G. sulfurreducens filaments from (B) the ~45 kDa band and (C) the ~30 kDa band. EMPAI: Exponentially modified protein abundance index. LC-MS/MS analysis showing matched peptides in (D) Omcs in the ~45 kDa band and (E) PilA in the ~6.5 kDa band. (F) Western immunoblotting using anti-PilA antibody showing PilA in purified filament preparations (left) and cell lysate (right).
Figure S2. De Novo Atomic Model Building of OmcS Filaments, Related to Figures 1 and 2
(A) Sequence-based alignment of OmcS and two other c-type cytochromes detected by mass spectrometry, with the same highlighted colors used in Fig 2A. Regions indicated by red, orange, yellow, green, cyan, blue and purple arrowheads are shown in (B)–(G), respectively, with OmcS atomic model fit into the cryo-EM map. Both OmcT and GSU2501 have deletions or insertions at those regions and cannot fit into the cryo-EM map.
Figure S3. Illustration of Cryo-EM Map Quality, Related to Figures 1 and 2
(A) The model-map FSC (Fourier Shell Correlation) calculation using a 0.38 criterion, which is sqrt (0.143), estimates the Spider map to have a resolution of 3.7 Å. 
(B) A view of amino acids 325-350 fit into the Spider map used in (A) after filtering the map to 3.6 Å.  (C) The Relion map:map “gold standard” FSC using the 0.143 criterion estimates the Relion map to have a resolution of 3.2 Å.  (D) A view of the same region as (B) but with the Relion map which has been filtered to 3.2 Å, shows that the Relion estimate of resolution is considerably over-optimistic.  (E) Fitting of six heme co-factors into the corresponding cryo-EM densities.  (F) The Histidine 51 interaction with heme 2 shown along with surrounding residues fit into the cryo-EM density.
Figure S4. Power Spectra of Purified Filaments from Fumarate-Grown Strain and Cell-Attached Filaments Are Similar to Power Spectra from Filaments of the Electrode-Grown Strain Used to Build the Atomic Model, Related to Figures 1 and 2

(A) OmcS filaments (black arrow) from the fumarate-grown strain. Some flagellar filaments (white arrow) are present in this strain. Scale bar, 500 Å. (B) The averaged power spectrum of the OmcS filaments from fumarate-grown cells (left) shows similar layer lines and helical symmetry as that from filaments of electrode-grown cells (right). (C) Power spectra generated from previously published images (Leang et al., 2013) of intact filaments attached to G. sulfurreducens cells. Left, the power spectrum of a bundle of filaments showing the meridional layer line (blue arrow) at 1/47 Å. Because of the bundling, a strong sampling is seen on the near-equatorial layer line. Right, averaged power spectra from 13 isolated segments to avoid the sampling. These power spectra show that the cell-attached filaments have symmetry similar to that as observed in cryo-EM images of purified OmcS filaments, with a rotation of 84 degrees and a rise of 47 Å per subunit.
Figure S5. The Thinner Filament Observed in Cryo-EM Imaging, Related to Figure 1

(A) The cryo-EM images frequently showed a thinner filament (white arrow) compared to OmcS filaments (black arrow). Scale bar, 200 Å. (B) The averaged power spectrum of the thinner filaments shows layer lines similar to those from the OmcS filament, but with a somewhat different axial rise of ~57 Å and rotation of ~160°. (C) A helical reconstruction of the thinner filament filtered to 20 Å. (D) The projection of the reconstruction in (C) with a low threshold (left) and a high threshold (right). The high threshold shows strong peaks separated by approximately 10 Å, labeled by yellow dots, which could be hemes.