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Mechanistic stratification in electroactive biofilms of *Geobacter sulfurreducens* mediated by pilus nanowires

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Electricity generation by *Geobacter sulfurreducens* biofilms grown on electrodes involves matrix-associated electron carriers, such as *c*-type cytochromes. Yet, the contribution of the biofilm's conductive pili remains uncertain, largely because pili-defective mutants also have cytochrome defects. Here we report that a pili-deficient mutant carrying an inactivating mutation in the pilus assembly motor PilB has no measurable defects in cytochrome expression, yet forms anode biofilms with reduced electroactivity and is unable to grow beyond a threshold distance ($\sim 10~\mu m$) from the underlying electrode. The defects are similar to those of a Tyr3 mutant, which produces poorly conductive pili. The results support a model in which the conductive pili permeate the biofilms to wire the cells to the conductive biofilm matrix and the underlying electrode, operating coordinately with cytochromes until the biofilm reaches a threshold thickness that limits the efficiency of the cytochrome pathway but not the functioning of the conductive pili network.

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he ability of Geobacter bacteria to completely oxidize organic compounds to CO2 with an electrode poised at a metabolically oxidizing potential shows promise for the conversion of renewable biomass into electricity, hydrogen and/or liquid fuels in bioelectrochemical systems¹⁻³. Energy recoveries in these devices depend greatly on the electroactivity of the electrode-associated biofilms⁴, yet the mechanism of biofilm conductance is not fully understood. Genetic studies in the model representative Geobacter sulfurreducens have identified genes encoding the biofilm exopolysaccharide (EPS) that anchors c-type cytochromes to the biofilm matrix⁵. Although the genome of G. sulfurreducens contains over 100 genes annotated as c-type cytochromes, some of which are required for efficient extracellular electron transfer^{6–8}, only one, OmcZ, is essential for biofilm electroactivity^{8,9}. Other *c*-type cytochromes may be required, but their genetic identification is challenging because of compensatory effects observed when the encoding genes are mutated^{7,10}.

In situ electrical conductivity measurements on living biofilms of G. sulfurreducens grown on an electrode demonstrated the thermal dependence of charge transport, which is consistent with the incoherent redox conductivity mediated by the haem groups of the matrix-associated c-type cytochromes¹¹. The role of cytochromes as electron carriers in the biofilms is further supported by the studies showing their reversible oxidization or reduction as a function of the anode potential 12,13. However, while it is possible to reduce all of the biofilm cytochromes of thin (<10 µm) biofilms with a positive voltage, only half are reduced in thicker ($\sim 20 \,\mu\text{m}$) biofilms¹². Furthermore, redox potentials decrease sharply as the biofilms grow to or beyond a 20-µm thickness¹⁴ and a redox gradient is established in these spatial scales that results in the accumulation of reduced cytochrome species further away from the electrode¹⁵. Source-drain experiments of biofilms grown across 10-µm wide gaps between electrodes indicate that the redox potential already decreases significantly in biofilm regions 10 µm away from the oxidizing electrode¹⁶. Thus, a redox gradient is established in the biofilms whereby more cytochrome-associated electrons are concentrated in the upper, electron-acceptor-limited biofilm stratum and decrease progressively towards the electrode-attached layers. The redox gradient has been proposed to provide the driving force for electron transport across the biofilms¹⁶, in a manner analogous to how electrons diffuse through redox polymers hopping among immobilized redox cofactors¹⁷

The progressive decline in the rates of cytochrome oxidation and reduction in biofilm layers 10-20 μm away from the electrode indicates that a rate-limiting step exists when biofilms grow to or beyond this thickness. Yet, despite this limitation, G. sulfurreducens can grow electroactive biofilms tens of micrometres from the electrode surface while generating current proportionally to the biofilm thickness¹⁸. Furthermore, cells in the upper biofilm layers are metabolically active and continue to oxidize acetate while contributing to current production¹⁹. Consistent with this, acetate kinase, which serves as a proxy for acetate oxidative metabolism, is uniformly abundant across the biofilm²⁰. By contrast, the expression of the outer membrane c-type cytochrome OmcB increases fourfold in cells located in the upper stratum²⁰, where the redox potential is lowest¹⁴. This has lead to the proposal that cells in these upper, electron-acceptor-limited zones compensate for the lower rates of cytochrome-mediated redox reactions and maintain optimal rates of respiration by expressing 'extra electron-transfer machinery'20.

One obvious candidate for discharging respiratory electrons and alleviating the electron acceptor limitation of cells in the upper biofilm regions is the conductive pili that *G. sulfurreducens* produces to reduce Fe(III) oxides²¹ and uranium²². The pili are

also required to grow multilayered biofilms on iron-coated and glass surfaces²³, plastic surfaces²⁴ and on electrodes poised at an oxidizing potential¹⁸. The need to express pili to form biofilms, and the immunodetection of the outer membrane c-type cytochrome OmcS on chemically fixed pili produced by planktonic cells²⁵, suggested that the pili may serve as a scaffold for the matrix-associated cytochromes 26. However, scanning tunnelling microscopy studies did not reveal spectroscopic haem signatures along cell-associated, untreated pili²⁷. The significance of OmcS in electroactive biofilms is also questionable, because deletion of the omcS gene had no effect in biofilm growth and maximum catalytic current⁹. Furthermore, omcS transcripts are less abundant in current-harvesting biofilms than in biofilms grown on the same electrode but operated in open circuit and fed the soluble electron acceptor fumarate⁸. A mere structural role for the pili is also incongruent with in vitro conductivity measurements, which show that the pilus fibres are protein nanowires capable of transporting charges at rates several orders of magnitude higher than the rates of cellular respiration²⁸. The conductivity measured along individual pilus fibres and thermal dependence of their differential conductance at biologically relevant voltages²⁸ is also consistent with the incoherent redox conductivity of living biofilms¹¹. Further, a structural model of the pilus fibre refined via molecular dynamics identified transversal and axial multistep charge hopping paths involving aromatic residues (phenylalanines and tyrosines) of the pili required for optimal biofilm electrochemical activity²⁹. The pili thus have the attributes necessary to function as biofilm electron carriers and could promote the discharge of respiratory electrons from cells in the electron-acceptor-limited upper regions to the oxidized cytochromes below. This is in accordance with the prediction that electron transfer across electroactive biofilms requires the expression of both cytochromes and pili to promote short-distance, electron-transfer reactions³⁰.

Much of the difficulty in assessing the contribution of the pili to biofilm electroactivity stems from the fact that pili-deficient mutants reported thus far also have defects in cytochromes required for extracellular electron transfer^{22,24,31}. Here we investigate the effect of several pili-inactivating mutations in the expression of cytochromes and identify one, a deletion in the gene encoding the pilin polymerization motor PilB, that has no measurable effects in cytochrome expression. Using the pilB mutant, we demonstrate that pili expression is required for optimal biofilm electroactivity in thin ($\sim 10 \,\mu m$) biofilms and to grow the biofilms beyond this threshold distance, which otherwise limits the rates of redox reactions involving the biofilm cytochromes. We also show that the pilB defects are similar to Tyr3, a mutant that carries alanine replacements in the three tyrosines of the pilus multistep hopping pathway²⁹ and produces pili with reduced conductivity. The results thus indicate that pili, like cytochromes, are electron carriers in anode biofilms, but their ability to discharge respiratory electrons becomes progressively more critical as redox reactions involving cytochromes become rate-limiting in the upper zones. This strategy confers on Geobacter cells a metabolic advantage over other bacteria relying solely on cytochromes or diffusible electron carriers, as it maximizes energy generation through extracellular electron transfer even in cells positioned at tens of micrometre distances from the electrode.

Results

Inactivation of PilB does not affect cytochrome expression. The assembly of type IVa pili on the bacterial inner membrane is mediated by a protein apparatus involving three interacting subcomplexes: the pilus subcomplex (major pilin and minor

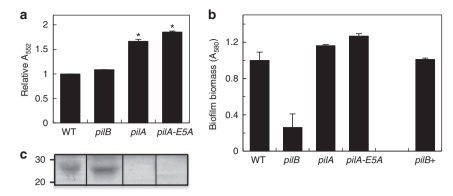


Figure 1 | Phenotypic characterization of pili-deficient mutant (pilB, pilA and pilA-E5A) biofilms. (a) Relative cytochrome content in biofilm extracts of the pili-deficient mutants in reference to WT (average and s.e. of two biological replicates for each). Significant differences in cytochrome content (*P<0.01) relative to the WT were identified in t-test pairwise comparisons. (b) Biofilm biomass on plastic surfaces estimated from the OD₅₈₀ of the biofilm-associated crystal violet solubilized with acetic acid (shown are averages and s.d. of six biofilm replicates of the WT, pili-deficient mutants and the genetic complemented pilB+ strain). (c) OmcZ_S band among haem-stained proteins isolated from the biofilm matrix of the WT and pili-deficient strains (full gels are shown in Supplementary Fig. 2c). Lanes were loaded with 20 µg of protein. Numbers at left are relative molecular masses of protein standards in kDa.

pilins, if present), the motor subcomplex (containing several proteins, including the PilB and PilT ATPases that power pilus polymerization and retraction, respectively) and the alignment subcomplex (which properly aligns the assembled pilus through the outer membrane PilQF secretin complex)³². As mutations that render the pilus and motor subcomplexes inoperative also prevent pili expression^{32,33}, we constructed mutants of *G. sulfurreducens* carrying deletions in the genes encoding the PilA pilin subunit (*pilA* mutant) and the PilB ATPase (*pilB* mutant), respectively. In addition, we constructed a pilideficient mutant (*pilA-E5A*) carrying an alanine substitution in the conserved glutamic E5 amino acid required for proper alignment and assembly of pilins^{32,34}.

Two of the pili-inactivating mutations, pilA and pilA-E5A, had pleiotropic effects in cytochrome expression. Differential redox spectroscopy of reduced-minus-oxidized ultraviolet-visible spectra (Supplementary Fig. 1) revealed, for example, a greater cytochrome content in the pilA and pilA-E5A biofilms than in the wild type (WT; Fig. 1a). Similar increases were observed in planktonic cells of the two mutant strains (Supplementary Fig. 2), ruling out any influence of the physiological state of the cells (planktonic or biofilm) in the phenotype. The gene encoding the outer membrane c-type cytochrome OmcB was also upregulated (Supplementary Fig. 3), suggesting that some of the pleiotropic cytochrome effects observed in the pilA and pilA-E5A mutants were transcriptional in nature. By contrast, transcript levels for the gene encoding the outer membrane *c*-type cytochrome OmcZ were similar in the WT, and the pilA and pilA-E5A mutants (Supplementary Fig. 3). Yet, the processed form of this cytochrome (OmcZ_S, ~30 kDa), which is released to the biofilm matrix⁵ and is required for optimal biofilm redox activity^{8,24}, was absent in the matrix of the mutant biofilms (Fig. 1c). OmcZ_S was however detected in haem-stained preparations of planktonic cells (Supplementary Fig. 2), suggesting that the inability of the pilA and pilA-E5A mutant biofilms to secrete this cytochrome was related to the unique physiology of cells living within a surfaceattached community.

Pili-deficient mutants have been reported to build thin biofilms on glass surfaces²³, presumably because the pili provide the structural support needed to stack cells on the surface. Consistent with this, 24-h biofilms of the *pilA* and *pilA-E5A* strains that formed on plastic surfaces were less dense than the WT (Supplementary Fig. 4); however, the mutant biofilms reached WT biofilm biomass levels after 48 h (Fig. 1b; Supplementary Fig. 4). The upregulation of the *xapD* gene, encoding the ATP-

dependent exporter of the biofilm EPS⁵, in these mutants suggests that EPS synthesis could have compensated for the pili deficiency during the early stages of biofilm formation, as reported for other bacteria³⁵. But once the mutant biofilms reached WT biomass levels at 48 h, the amount of EPS produced per cell was similar to the WT (Supplementary Fig. 5).

In contrast to the pleiotropic nature of the pilA and pilA-E5A mutations, preventing pilin assembly through the inactivation of the PilB motor (pilB mutant) had no measurable effects on the cytochrome content of the biofilms (Fig. 1a). Furthermore, the transcription of genes in the pilin operon (pilA and GSU1497) or those encoding the outer membrane c-type cytochromes OmcB and OmcZ was not affected either (Supplementary Fig. 3). Moreover, OmcZ_S was secreted in the pilB biofilm matrix (Fig. 1c), and xapD transcripts (Supplementary Fig. 3) and biofilm EPS produced per cell (Supplementary Fig. 5) were similar in the pilB and WT biofilms. As with other pili-deficient mutants²³, the *pilB* biofilms that formed on plastic surfaces were less dense than the WT even after 48 h of incubation (Supplementary Fig. 4), as indicated by the reduced biofilm biomass stained with crystal violet (Fig. 1b) and the lower cell protein content extracted from the *pilB* biofilms (75 \pm 1% of the WT, average and s.e. of two independent experiments). Yet, the pilB biofilm defect was rescued in the genetically complemented pilB+ strain, which restored pili production through the expression of the pilB gene in trans from a medium-copy plasmid (Fig. 1b). Hence, the pilB strain provides the elusive genetic tool needed to assess the contribution of Geobacter conductive pili in the growth and electrochemical activity of anode biofilms.

Thin ($\sim 10 \,\mu\text{m}$) biofilms require both cytochromes and pili.

We investigated the ability of the pili-deficient mutants to generate current in microbial electrolysis cells (MECs) fed an initial concentration of 1 mM acetate and operated in batch in reference to the WT strain. Under these conditions, all of the strains tested grew reproducibly to a thickness of ca. 10 μ m, but the rates of current production and the maximum current reached were reduced in the three pili-deficient mutants (Fig. 2a). The *pilA* and *pilA-E5A* mutant biofilms, which in addition to the pili deficiency do not secrete OmcZ_S in the biofilm matrix (Fig. 1c), reached low levels of current (0.08 \pm 0.05 and 0.14 \pm 0.08 mA, respectively). As a result, prolonged incubations were required for the mutant biofilms to oxidize all the acetate in the anode chamber (Fig. 2a)

and to reach WT thickness ($\sim 13.5 \pm 3.5$ and $11.6 \pm 1.9 \, \mu m$, respectively, in duplicate MECs). Similar low levels of current $(0.11 \pm 0.07 \, \text{mA})$ for duplicate MECs and lengthy incubation times were recorded in control MECs driven by a null omcZ mutant (Fig. 2a), consistent with the need of the biofilms to concentrate $OmcZ_S$ near and on the electrode for efficient electron transport to the anode surface³⁶. The expression of $OmcZ_S$ in the pilB biofilm matrix (Fig. 1c) improved MEC performance compared with the pilA and pilA-E5A mutants, but did not fully restore WT levels of current (Fig. 2a). Maximum current harnessed from the pilB anode biofilms in triplicate MECs averaged, for example, $0.52 \, (\pm 0.06 \, \text{mA})$, which is about half of the maximum current produced by WT biofilms $(1.06 \pm 0.05 \, \text{mA})$.

The partial restoration of biofilm electroactivity in pilB compared with the other pili-deficient mutants did not result from functional complementation by the PilB homologues MshE and GspE (putative ATPase motors of type II secretion systems) because deleting the pilB and mshE or gspE genes generated double mutants that were phenotypically indistinguishable from the pilB single mutant (Fig. 2a, inset). In addition, the pilB mutant defect could not be complemented in MECs co-inoculated with the pilB and omcZ mutant cells (Fig. 2a, inset). We did observe a delay in current generation in the co-culture-driven MECs, which is the phase when cells colonize the electrode before growing on it while coupling the oxidation of acetate to the reduction of the electrode³⁷. However, once current production was initiated, the performance of the co-culture MECs was similar to the pilB MECs. This suggests that biofilm cells need to express both pili and OmcZ_S for optimal biofilm growth and electroactivity.

The rates of current increase in the *pilB* MECs, which correlate with the exponential phase of biofilm growth³⁷, were also reduced

in half $(0.61 \pm 0.15 \,\text{mA})$ per day compared with the WT biofilms $(1.45 \pm 0.06 \,\mathrm{mA})$ per day). However, coulombic efficiencies in the pilB-driven MECs (96.6 \pm 3.1) were similar to the WT (94.6 ± 4.1) , indicating that, on average, the pilB biofilms converted the same amount of acetate to electricity as the WT biofilms, they just did so at slower rates. Confocal micrographs of WT and pilB biofilms collected at the end of the experiment, when all of the acetate had been depleted, and stained with fluorescent viability dyes showed predominantly live cells and similar biofilm structure (Fig. 2b,c, respectively). Furthermore, the biofilm biomass estimated from the fluorescence emitted by the biofilm cells was similar in both strains (10.05 \pm 1.00 in WT and $9.06 \pm 1.46 \,\mu\text{m}^3 \,\mu\text{m}^{-2}$ in *pilB*). Thus, the *pilB* defects cannot be attributed to a reduction in the number of cells actively contributing to electron transfer or structural variations of the biofilms. Furthermore, the pilB phenotypes in MECs fed with 1 mM acetate (Fig. 2) are similar to those reported for Tyr3, a mutant that produces pili with alanine replacements in the pilin's three tyrosines of the pilus multistep hopping pathway that impair the cell's ability to transfer electrons to extracellular electron acceptors such as Fe(III) oxides²⁹. Hence, the pilB phenotypes are consistent with reductions in the respiratory rate of individual cells resulting from the inability of the mutant cells to discharge electrons using the pili.

The conductivity of the pili is required to grow thick biofilms. By increasing the initial amount of electron donor (acetate) added to the anode chamber of the MECs operated in batch, we were able to increase the thickness of the WT anode biofilms from ~ 10 to $15\,\mu m$ (2 mM acetate) and $20\,\mu m$ (3 mM acetate; Fig. 3a). These are spatial scales that lead to the progressive accumulation

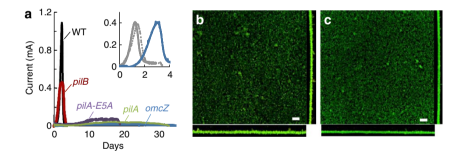


Figure 2 | Current production by anode biofilms of WT and mutants in MECs fed with 1 mM acetate. (a) Representative plots of current generation by WT, pili-deficient strains (*pilB*, *pilA* and *pilA-E5A*) and *omcZ* mutant in MECs operated in batch and fed with an initial concentration of 1 mM acetate. Inset shows current generation (*Y* axis, in mA) over time (*X* axis, in days) by double mutants *pilB* mshE (solid grey), *pilB* gspE (dashed grey) and an equal mixture of *pilB* and *omcZ* cells (blue). (b,c) Confocal micrographs showing top and side views of anode biofilms of the WT (b) and *pilB* (c) strains collected at the end of the MEC experiment, when current had decreased to < 0.1 mA, and stained with the BacLight viability kit (green, live; red, dead). Scale bar, 20 μm.

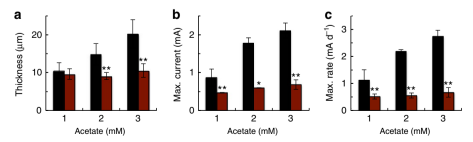


Figure 3 | Thickness and electroactivity of WT and pilB biofilms. Anode biofilm thickness (a), maximum current (b) and maximum rates of current production (c) in batch MECs fed with an initial concentration of 1, 2 and 3 mM acetate and driven by the WT (black) or the pilB mutant (maroon) strains. Shown are average and s.d.'s or errors of four (1 mM) or two (2 and 3 mM) independent MEC experiments, respectively. Significant changes between the WT and pilB values at each acetate concentration are indicated with stars (* $^{*}P < 0.05$; * $^{*}P < 0.005$). Representative plots of current generation used to estimate biofilm thickness and electroactivity at each acetate concentration are shown in Supplementary Fig. 6.

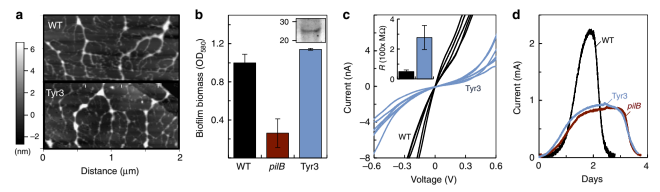


Figure 4 | Role of pilus conductivity in the growth and electroactivity of anode biofilms. (a) AFM topographic images of pili purified from the WT and Tyr3 strains and deposited on a HOPG surface. (b) Crystal-violet-stained biomass of 48-h-old biofilms of the *pilB* and Tyr3 mutants relative to WT (shown are averages and s.d.'s of six biofilm replicates for each strain). Inset: Omc Z_S band in haem-stained proteins isolated from the biofilm matrix of the Tyr3 biofilms (20 μ g of protein loaded per lane; full gel is shown in Supplementary Fig. 2d). Numbers at left are relative molecular masses of protein standards in kDa. (c) Representative current-voltage (*I*-*V*) plots obtained after probing the transversal conductivity of individual WT and Tyr3 pili (a). Inset shows the average and s.d. of the electrical resistance of four or more WT or Tyr3 pili, each probed in at least three positions. (d) Current production by WT, *pilB* and Tyr3 biofilms grown in MECs fed initially with 3 mM acetate.

of reduced cytochromes in the upper biofilm layers, thus limiting the ability of the cells to use cytochromes in these regions as electron acceptors for respiration 12 . Yet, cells in these electron-acceptor-limited layers are actively respiring and contributing to current production. To investigate a potential role for the conductive pili in alleviating the limitations imposed by the reductions of redox potential in the upper biofilm stratum, we grew the $\it pilB$ mutant in MECs fed with 2 and 3 mM acetate. In contrast to the linear increases in WT biofilm thickness with initial acetate feeding, the $\it pilB$ biofilms remained thin ($\sim 10\,\mu m$) under all the conditions tested (Fig. 3a). Similarly, parameters that measure the biofilm electroactivity such as current maxima and rates of current production increased proportionally to acetate availability in the WT MECs, but were unaffected in the $\it pilB$ MECs (Fig. 3b,c).

As a control, we also characterized the Tyr3 mutant strain phenotypically (Fig. 4). The Tyr3 mutant produced pili at WT levels (Fig. 4a), which provided the structural support needed to grow dense biofilms on plastic surfaces when fumarate was provided as the electron acceptor (Fig. 4b). The Tyr3 biofilms also expressed OmcZ_S in the biofilm matrix (Fig. 4b, inset). However, as predicted in molecular dynamics simulations²⁹, the reduced number of aromatic contacts in the Tyr3 pili increased their electrical resistance (\sim 280 M Ω) fivefold over that measured in the WT pili ($\sim 50\,\mathrm{M}\Omega$) (Fig. 4c). Furthermore, Tyr3 anode biofilms in MECs fed with 3 mM acetate remained thin $(13.3 \pm 2.4 \,\mu\text{m}, \text{ for duplicate MECs})$ and reached current levels $(0.81 \pm 0.10 \,\mathrm{mA}, \,\,\mathrm{for} \,\,\mathrm{duplicate} \,\,\mathrm{MECs})$ comparable to those recorded in the pilB-driven MECs and fed the same amount of acetate (Fig. 4d). Thus, the ability of cells to grow thick ($> 10 \mu m$) biofilms on the anode electrode while coupling the oxidation of acetate to current generation depends on the conductivity, rather than the structural support, provided by the pili.

Discussion

The studies described herein provide the elusive genetic evidence, in the form of the *pilB* mutant, needed to assess the contribution of the conductive pili of *G. sulfurreducens* to the electrochemical activity of electrode-associated biofilms. Indeed, the *pilB* mutant had no measurable changes in *c*-type cytochromes (Fig. 1) and was able to form biofilms of WT thickness ($\sim 10 \, \mu m$) in MECs operated in batch and fed an initial acetate concentration of 1 mM (Fig. 2). Yet, despite forming anode biofilms as thick as the

WT under these conditions, the pili deficiency prevented the mutant from generating current at the same rates as the piliated WT strain and reduced current maxima (Fig. 3). Furthermore, the pilB mutant was unable to grow biofilms beyond this threshold thickness when the initial concentration of acetate fed to the MECs was increased from 1 to 2 or 3 mM, which are conditions that allow the WT biofilms to grow ~ 15 - and 20- μ m thick, respectively (Fig. 3). We can rule out mass transport limitations because the biofilms investigated in this study were too thin (<50 µm) to restrict the diffusion of the electron donor, acetate³⁸. Furthermore, the phenotypes of the pili-deficient pilB mutant in MECs were similar to those of a Tyr3 mutant, which produces pili with reduced conductivity (Fig. 4). This finding challenges the proposal that the pili's role is restricted to providing structural support needed for cytochrome organization²⁶ and is instead consistent with the need of the biofilm cells to express conductive pili to transport electrons.

The genetic evidence therefore supports a model in which the conductive pili and the cytochromes work coordinately as electron carriers to maintain optimal rates of electron transfer in thin (~ 10 -µm thick) biofilms (Fig. 2). This is in accordance with the early models that suggested that electron transfer across electroactive biofilms may require the expression of both cytochromes and pili to promote short-distance, electron-transfer reactions³⁰. However, as the biofilms grow further away from the electrode, the cytochromes become progressively more reduced and the ability of the pili to discharge respiratory electrons becomes more critical (Fig. 3). The redox potential decreases significantly already in regions of the biofilm only 10 µm away from an oxidizing electrode¹⁶. When compared with a 10-µm thick WT biofilm, the electrochemical activity of the pilB biofilms (rates of current production and maximum current) was reduced approximately in half (Figs 2 and 3). The pilB defects cannot be attributed to decreases in the number of biofilm cells contributing to electron transfer, because the biofilm thickness and the number of viable cells in the *pilB* biofilms were comparable to the WT. EPS production (Supplementary Fig. 5) and OmcZ_S expression in the biofilm matrix (Fig. 1c) were also similar in the pilB and WT biofilms. The reductions in electrochemical activity of 10-μm *pilB* biofilms are also within the ranges reported for Tyr3 biofilms grown to a similar thickness²⁹. Tyr3 pili are expressed at WT levels, but carry alanine replacements in the three tyrosines of the pilus multistep hopping pathway²⁹ that reduce the conductivity of the pili more than fivefold (Fig. 4). We measured, for example,

an average electrical resistance of $\sim 50 \, \mathrm{M}\Omega$ for WT pili deposited on highly oriented pyrolytic graphite (HOPG), which is within the ranges reported for pili deposited on gold electrodes²⁸, and $\sim 280 \, \mathrm{M}\Omega$ in the Tyr3 pili. Thus, the reduced electrochemical activity of the *pilB* and Tyr3 biofilms results from the inability of the mutant cells to discharge respiratory electrons via the pili.

In vitro charge transport measurements indicate that individual pilus fibres can transport charges at micrometre distances and at rates (ca. 10⁹ electrons per second for a 1-µm-long pilus at 100 mV) that greatly exceed the cellular respiratory rates²⁸. Furthermore, the pili can reach micrometre lengths²¹ and physically interact with extracellular cytochromes²⁵ of the biofilm matrix⁵. This could allow the biofilm pili to directly discharge electrons from the cells to matrix-associated cytochromes such as OmcZ_S. This multihaeme c-type cytochrome preferentially localizes closer to the electrode and on its surface³⁶ and has the wide redox potential working range³⁹ needed to function as an electrochemical gate between the biofilm and the electrode³⁶. Indeed, pilin mutations that resulted in defects in OmcZ_S expression (pilA and pilA-E5A; Fig. 1) reduced the capacity of the biofilms to produce current to levels similar to a control omcZ mutant (Fig. 2). The pilA and pilA-E5A mutant cells also grew very slowly on the anode, taking several days to oxidize the acetate and reach WT biofilm thickness ($\sim 10 \, \mu m$). The expression of OmcZ_S in the *pilB* biofilm matrix (Fig. 1c) alleviated the defect but only partially (Fig. 2). Thus, the pili are required for optimal current production within spatial scales $(\sim 10 \,\mu\text{m})$ where biofilm cytochromes, including OmcZ_S, are available as electron carriers.

The coordinated interaction between pili and cytochrome electron carriers in the biofilm layers closer to the electrode is further supported by the cross-regulation of pili assembly and cytochrome export observed in this and one other study⁴⁰. Earlier studies demonstrated that the PilA pilin protein of G. sulfurreducens is translated as both a short and a long prepilin isoform, which interact to promote the processing and assembly of the short pilin and the export of *c*-type cytochromes such as OmcZ to the outer membrane⁴⁰. Indeed, we showed that mutations that prevented pilin expression or disrupted pilin-pilin interactions (pilA and pilA-E5A, respectively) interfered with the pilin's role in secretion and lead to defects in cytochrome expression and localization (Fig. 1). Similarly, *pilA*-deficient mutants used in earlier genetic studies ^{18,21,41} also had defects in c-type cytochromes required for optimal extracellular electron transfer^{22,24}. The dual role that Geobacter pilins play in pili synthesis and cytochrome export is in accordance with the multiple biological roles reported for type IV pilins from other bacteria. In Pseudomonas aeruginosa, for example, the structural subunit of the pilus fibre (the PilA pilin) interacts with components of the general secretion pathway to promote the export of proteins across the outer membrane⁴². As a result, mutations that inactivate PilA in this bacterium affect the secretion of numerous proteins and produce mutant strains with multiple phenotypes. By contrast, the function of PilB is restricted to pilus biogenesis. Thus, inactivating mutations in the pilB gene of P. aeruginosa prevent pili expression, but do not cause export defects⁴³. Similarly, inactivating the PilB motor of G. sulfurreducens prevents the assembly of the pilins, but does not interfere with pilin interactions needed for cytochrome export. As a result, cytochrome expression in the pilB mutant was as in the WT strain (Fig. 1).

As the biofilms grow further apart from the electrode (beyond a threshold distance of $\sim 10\,\mu\text{m}$), the reversible oxidization or reduction of biofilm cytochromes becomes rate-limiting ¹². Redox potentials decrease sharply beyond this threshold distance ^{14,16} and reduced species progressively accumulate in biofilm layers

further away from the electrode¹⁵. The redox gradient provides the driving force for electron transport across the biofilms ¹⁶, but also limits the ability of cells in the upper layers to discharge respiratory electrons to nearby cytochromes. It has been proposed that cells in these outer regions of the biofilm overcome this limitation by upregulating the expression of electron transport proteins involved in extracellular electron transfer²⁰. Consistent with this model, while it was possible to grow thicker (ca. 15 and 20 µm) WT biofilms in MECs, preventing the assembly of the pilins (pilB mutant) or reducing the conductivity of the assembled pili (Tyr3 mutant) effectively prevented the biofilms from growing > 10 µm away from the electrode (Fig. 4). Hence, the expression and conductive properties of the pili are required to maintain optimal electrical connectivity as the biofilms grow in thickness and the cytochromes become progressively more reduced. These upper layers of the biofilm may be electron acceptor limited, but are also the regions where electron donor (acetate) availability is highest 19. Studies show G. sulfurreducens can continue to metabolize acetate in the absence of an electron acceptor by overexpressing its extracytoplasmic cytochromes and using them as a capacitor to store up to 10⁶ electrons per cell⁴⁴. Indeed, the outer membrane c-type cytochrome OmcB, which is transcriptionally upregulated under conditions of electron acceptor limitation⁴⁵, is four times more abundant in cells from the 10-20-μm upper biofilm stratum than in the biofilm layers closer to the electrode²⁰. This suggests that cells in these upper biofilm regions could continue to metabolize acetate and generate a proton motive force for energy generation, accumulating electrons in the cytochromes of the cell envelope. The pilus apparatus is anchored in the cell envelope of Gram-negative bacteria; hence, it could potentially accept electrons from the extracytoplasmic cytochromes, promoting their discharge from the cell to the biofilm matrix. By growing several micrometres in length²¹ and intertwining to form a complex web of nanowires²⁷, the pili could electronically connect the upper and bottom strata of the biofilm, effectively bypassing the rate limitation imposed by the accumulation of reduced cytochromes in the electron-acceptor-limited regions.

Taken together, the results support a model of mechanistic stratification in electroactive biofilms mediated by the pilus nanowires (Fig. 5). According to the model, the conductive pili network works coordinately with the matrix-associated cytochrome carriers to transfer electrons in the region (<10 µm) closer to the electrode. However, the pili become the primary mechanism for discharging respiratory electrons in the upper (>10 µm) region, where cytochromes are predominantly in a reduced state (Fig. 5). The pili can discharge electrons at micrometre distances and at rates that greatly exceed the cellular respiratory rates²⁸. These attributes could allow the pili network to electronically connect cells in the upper layers of the biofilm to the oxidized cytochromes present in the regions closer to the electrode. As a result, mutants that do not assemble (pilB) or produce poorly conductive (Tyr3) pili cannot grow beyond this threshold distance and have reduced electroactivity compared with WT biofilms of comparable thickness (Fig. 5). The coordinated interactions between pili and cytochrome electron carriers revealed in our genetic studies challenge the proposal⁴¹ that metallic-like pilus conductivity fully accounts for electron flow across the biofilms and is in accordance with the prediction³⁰ that electron transfer across electroactive biofilms involves short-distance, multistep hopping reactions between pili and cytochromes. Furthermore, living biofilms show the thermal dependence of incoherent conductivity predicted for electrontransfer reactions involving haems of c-type cytochromes¹¹ and also demonstrated for charge transport via pili²⁸. Phenylalanine and tyrosine residues cluster during pilin assembly, creating axial

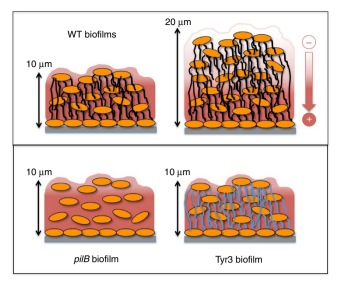


Figure 5 | Model of mechanistic stratification of electroactive biofilms.

The pili permeate the biofilm matrix and promote the discharge of respiratory electrons to the oxidized cytochromes (in red, positive charge) in thin (~ 10 - μm thick) biofilms. As the biofilms grow in thickness, reduced cytochromes accumulate (in white, negative charge) and the pili provide the means for the cells to discharge respiratory electrons to the oxidized cytochromes below. In the absence of pili (pilB biofilms) or in biofilms expressing poorly conductive pili (Tyr3 biofilms), biofilm growth is limited to the thickness ($\sim 10~\mu m$), where oxidized cytochromes are available to serve as electron acceptor to the cells.

and transversal pathways with inter-aromatic distances and dimer geometries optimal for multistep hopping²⁹. The tyrosine residues are in close proximity to acidic residues, which could be transiently protonated to promote proton-coupled electron transfer between tyrosine residues²⁹. The simultaneous transfer of a proton and an electron lowers the oxidation potential of the tyrosine residue to the levels needed to enable fast rates of electron transfer in biological systems^{46,47}. The transient protonation of acidic residues of the pili during charge transport also makes the rates of charge transport along the pili pH dependent. Thus, pH gradients that establish across anode biofilms^{14,48} likely tune the rates of charge transport via the pili and interactions with cytochromes, particularly in the biofilm regions closer to the electrode, where the pH is lower.

In addition to pH gradients, redox and electrical gradients form across the biofilms that can influence the physiology of the cells and could control the efficiency of charge transport along the pili and interactions with cytochromes to maintain optimal rates of electron transfer. Nutrient gradients also form in biofilms, which can influence the cell physiology and, therefore, the expression of biofilm electron carriers ¹³,14. Microarray analyses of thin sections of anode biofilms of G. sulfurreducens did not detect changes in the expression of the pilA gene⁴⁹. However, pilA is also transcribed under conditions that do not result in pili assembly⁵⁰. Thus, the levels of cell piliation across the biofilms could be modulated by the local biofilm microenvironment to maintain optimal rates of respiration, as observed for other respiratory components²⁰. The regulatory networks that control the expression and electrochemical activity of electron carriers such as the pili may be responsive to more than one biofilm parameter, such as pH, nutrient availability and redox potential. Biofilms of oxygenrespiring bacteria are, for example, responsive to redox potentials and use redox cues from their microenvironment to alter the biofilm structure, maximize oxygen diffusion and maintain redox homeostasis⁵¹. Similar mechanisms could control the expression

of pili and cytochromes to facilitate their coordinated interactions and redox homeostasis in *G. sulfurreducens* biofilms. Thus, understanding the multiple functions that pili play in *G. sulfurreducens* and their regulation may prove instrumental to improve the performance of *Geobacter*-driven electrochemical systems for applications in bioenergy.

Methods

Bacterial strains and culture conditions. The bacterial strains used in this study are described in Supplementary Table 1. The WT strain, G. sulfurreducens strain PCA (ATCC 51573), was kindly provided by Daniel Bond (University of Minnesota) and was used to construct three knock-out mutants: pilB (carrying a deletion in the pilB gene, GSU1491), pilA (carrying a deletion in the pilin gene pilA, GSU1496) and omcZ (carrying a deletion in the omcZ gene, GSU2076). When indicated, the pilB mutation was complemented in trans by expressing a WT copy of pilB from the pRG5 plasmid (pilB+ strain). The WT strain was also used to construct the pilA-E5A mutant (carrying a single alanine replacement of the glutamic 5 residue in the mature pilin PilA protein) and the Tyr3 mutant (carrying alanine replacements in the pilin's three tyrosine residues Y27, Y32 and Y57). The WT and mutant strains were routinely cultured anaerobically in NB medium supplemented with 15 mM acetate as the electron donor and 40 mM fumarate as the electron acceptor (NBAF) and 0.1% yeast extract and 1 mM cysteine (NBAFYE). For biofilm assays, cells were cultured in modified freshwater FW medium²² with 30 mM acetate and 40 mM fumarate. Cells used to inoculate the anode chamber of MECs were grown in mineral DB medium⁵³ supplemented with 20 mM acetate and 40 mM fumarate (DBAF).

DNA manipulations and mutant construction. Deletions of the pilB, pilA and omcZ genes were constructed using the cre-lox system⁵⁴ with the primers listed in Supplementary Table 2. The general procedure included the PCR amplification of the gentamycin (Gm) resistance cassette (aaaC1) flanked by loxP sites (Gm-loxP) from plasmid pCM351 (ref. 54) using primer set RS21-RS22 and of the upstream/ downstream target chromosomal regions using primer sets RS1-RS2/RS3-RS4 (pilB), RS5-RS6/RS7-RS8 (pilA) and RS9-RS10/RS11-RS12 (omcZ). The upstream region of the target gene, the Gm-loxP cassette and the downstream region of each gene were then fused in that order by overlap extension PCR using the corresponding external forward and reverse primers, and the Herculase II Fusion DNA Polymerase (Agilent Technologies). PCR conditions for this last amplification step were: 2 min of denaturation at 95 °C; 35 cycles of 20 s at 95 °C, 25 s at 54 °C and 2 min at 72 °C; and a final 3-min extension at 72 °C. The PCR products were then separated on an agarose gel, purified using the Zymoclean Gel DNA Recovery kit (Zymo Research), and cloned into the pCR2.1 plasmid (Supplementary Table 2) using the TOPO TA Cloning kit (Life Technologies) for sequence confirmation. After cloning, the constructs were PCR-amplified with the external primers, purified from an agarose gel and electroporated into electrocompetent cells of G. sulfurreducens following a previously published procedure⁵². Selection of recombinant strains was performed on NBAFYE plates supplemented with $5 \,\mu \mathrm{g} \,\mathrm{ml}^{-1}$ of gentamycin. When indicated, the Gm cassette carried by the pilB mutant was excised from its chromosomal location by expressing the Cre recombinase from plasmid pCM158 (ref. 54) and selecting for transformants on NBAFYE plates supplemented with $200 \, \mu g \, ml^{-1}$ of kanamycin. The excision of the marker was confirmed by PCR and the resulting mutant (pilB) was transferred twice in NBAFYE medium without kanamycin and then plated on NBAFYE with and without kanamycin to confirm the loss of the plasmid. The pilB strain was used to construct the double pilB gspE and pilB mshE mutants (Supplementary Table 1) using the general PCR procedure described above but with primer sets RS9-RS12, RS13-RS16 and RS17-RS20 (Supplementary Table 2), respectively.

We constructed the pilA-E5A and Tyr3 mutants by introducing one (E5A) or three (Y27A, Y32A and Y57A) alanine replacements, respectively, in targeted codons in the pilA gene (GSU1496). The primers used to construct these mutants are listed in Supplementary Table 2. The general procedure was to PCR amplify the pilA gene with its upstream chromosomal region (primers RS23-RS24), the downstream region of pilA (primers RS25–RS26) and a spectinomycin cassette (Sp; aadA gene) from plasmid pRG5 (ref. 10; primers RS27-RS28). The three fragments were then fused by overlap extension PCR (primers RS23 and RS26) to generate a 1684-bp DNA construct (pilA-Sp) containing the pilA gene and its upstream region, the antibiotic cassette (aadA), and the pilA downstream region. The construct was gel-purified and cloned into plasmid pCR2.1-TOPO TA vector (Invitrogen). Targeted nucleotide substitutions in the pilA gene were introduced using the QuikChange Lightning Site-Directed Mutagenesis kit (Agilent Technologies) using primers RS29 and RS30 (E5A substitution). Tyr3 was generated by sequentially introducing Y27A (RS31-RS32), Y32A (RS33-RS34) and, last, Y57A (RS35-RS36), as previously described²⁹. The mutated fragments were confirmed by sequencing before PCR amplification using the external primers, gel purification and electroporation into electrocompetent cells of G. sulfurreducens. Selection of recombinant strains was performed on NBAFYE plates supplemented with 75 μg ml $^{-1}$ of spectinomycin.

Reverse Transcription-quantitative-PCR. Transcripts for key components of the biofilm matrix (pili, EPS matrix and matrix-associated c-type cytochromes) were quantified by reverse transcription-quantitative PCR (RT-qPCR). The gene targets were the pilin-encoding gene pilA (GSU1496) and its downstream gene (GSU1497) in the pilA operon²¹, the ATP-dependent EPS exporter xapD (GSU1501)⁵, the outer membrane c-type cytochrome omcB (GSU2737) (ref. 6) and omcZ (GSU2078), which encodes for the precursor of the matrix-associated c-type cytochrome OmcZ_S (ref. 39). The constitutive gene rpoD (GSU3089) was used as a control. WT, pilB, pilA and pilA-E5A biofilms were grown for 48 h in the wells of 48-well polystyrene plates containing 600 μl of FWAF before decanting the culture broth and resuspending the biofilm cells in 50% (v/v) of ice-cold methanol to stop transcription. The cell suspension was then centrifuged to isolate the cells as a pellet and trizol reagent (Invitrogen) was used to extract their RNA. RNA treatment with RNase-free DNase (Promega) and reverse transcription with random primers (Promega) using the Super Script III Reverse Transcriptase (Invitrogen) were carried out following manufacturers recommendations. RT-qPCR was performed using the rEVAluation qPCR Master Mix (Syzygy), as recommended by the manufacturer, using the primers listed in Supplementary Table 2. The comparative C_T method⁵⁵ was used to calculate the relative expression of each gene using rpoD constitutive expression as an internal control (ΔC_T value or C_T (target) $-C_T$ (tpoD)) for each strain, and the average of the difference between each mutant strain ΔC_T and the WT ΔC_T was used to calculate the $\Delta \Delta C_T$. The relative fold change of expression for each target gene versus the rpoD internal control for each mutant strain versus the WT was then calculated with the formula $2^{-\Delta\Delta CT}$ for four replicate samples. Statistically significant changes in gene expression relative to pilB were determined in pairwise comparisons using the t-test function of the Microsoft Excel software.

Microbial electrolysis cells. The growth and electrochemical activity of the WT and mutant biofilms were assayed in H-type MECs equipped with anode and cathode graphite rod electrodes (Alfa Aesar, 1.27-cm diameter, 99% metals basis, 12 cm²) and a 3 M Ag/AgCl reference electrode (Bioanalytical Systems, Inc.). The MECs were set up, inoculated with cell suspensions harvested from early stationary phase DBAF cultures, filled with DB medium and operated in batch with a poised anode electrode (0.24 V versus reference electrode) as described previously except that the electron donor in the anode chambers was acetate, provided in 1, 2 or 3 mM concentrations, as indicated. Supernatant samples were periodically removed from the anode medium broth, filtered (0.45 µm) and analysed by highperformance liquid chromatography (HPLC), as described elsewhere⁵⁶, to monitor electron donor removal. Current production was recorded with a VSP potentiostat (BioLogic) and biofilm electroactivity was inferred from the rates of linear current production (mA per day) before reaching maximum current and initiation of the deceleration phase. The exponential phase of current production was also fitted statistically ($\hat{R}^2 > 0.98$) to an exponential curve and the exponent of the resulting formula was used to estimate the generation times of the biofilm cells growing exponentially on the anode electrode.

At the end of the MEC experiment, when current had decreased to $<\!0.1\,\mathrm{mA}$, the anode electrodes were removed and the live and dead biofilm cells were differentially stained in green and red, respectively, with the SYTO-9 and propidium iodide dyes of the BacLight viability kit (Invitrogen). The anode electrodes with the stained biofilms were then immersed gently in a Lab-Tek coverglass chamber (Nunc) filled with 3 ml of PBS and examined using a FluoView FV1000 inverted microscope system (Olympus, Center Valley, PA) equipped with an Olympus UPLFLN \times 40 oil immersion objective (numerical aperture, 1.30). SYTO-9 was excited at 488 nm and propidium iodide was excited at 543 nm. Vertical two-dimensional images of the biofilms were collected every 1 $\mu \mathrm{m}$ from \sim 10 random fields (1,024 by 1,024 pixels, 0.31 $\mu \mathrm{m}$ per pixel) per electrode, using a minimum of two biological MEC replicates. The biofilm thickness was then manually analysed by averaging thickness measurements of five representative areas per field. When indicated, the COMSTAT software 57 was used to estimate other biofilm structural parameters, as described previously 53 .

Static biofilm assays on plastic surfaces. The WT and mutant strains were also grown on plastic surfaces using a soluble electron acceptor (fumarate), essentially as described elsewhere²⁴. Briefly, the strains were grown in FWAF to midexponential phase and inoculated to a final OD_{600} of 0.02 in 150 μl of FWAF dispensed in each of six replicate wells of a 96-well, polystyrene, tissue culturetreated plate (Costar, Corning Life Sciences). Unless otherwise indicated, the plates were incubated for 48 h at 30 °C and the optical density of the culture at 600 nm (OD₆₀₀) measured before discarding the culture broth and staining the biofilms for 30 min with 150 µl of a 0.01% (w/v) aqueous solution of crystal violet. After staining, the biofilms were washed with double-distilled H2O and dried overnight before resolubilizing the biofilm-associated crystal violet with 33% (v/v) acetic acid. The optical density of the solution at 580 nm (OD580) was used to estimate the biofilm biomass (cells and matrix) and the biofilm biomass for each strain was averaged for six replicate samples. Time-course biofilm assays were performed the same way and biofilm growth was expressed as the biofilm biomass that stained with crystal violet (OD580) relative to the OD600 of the planktonic culture.

When indicated, 48-h WT and pilB biofilms grown in the wells of three 48-well plates were collected and treated with SDS, as described earlier²⁴, to lyse the cells and solubilize the total biofilm protein. Protein in the biofilm lysates was estimated with the Pierce BCA protein assay (Thermo Fisher Scientific). We estimated that $\sim 90\%$ of the protein solubilized with SDS was contributed by the biofilm cells, thus serving as a proxy of cell numbers in the biofilms.

Isolation and characterization of the biofilm matrix. The WT and mutant strains were each grown in the wells of two 48-well plates for 48 h to grow biofilms, as described above. The biofilm biomass from the two 48-well plates was then scraped and pooled together, and biofilm lysates were generated by SDS treatment, as described above for the isolation of cell protein. The biofilm matrix was isolated as previously described²⁴, except that the biofilm lysate was generated by repeated passages of the biofilm biomass through an 18-G needle and included several steps of centrifugation and washes to isolate the insoluble biofilm matrix fraction. The sample was lyophilized and its EPS component was acid hydrolysed, as previously described¹. EPS content was calculated as glucose equivalents using a published protocol⁵, except that the glucose content in the acid-hydrolysed sample was estimated by HPLC, using a Waters HPLC instrument (Milford, MA) equipped with the refractive index and ultraviolet detectors, and operated as described elsewhere⁵⁸.

Assays for cytochrome content and profiling. The cytochrome content of 48-h biofilms and planktonic cells of the WT and mutant strains was estimated by redox difference spectroscopy. The biofilms were first grown for 48 h in 96-well plates, as described above for the biofilm assays on plastic surfaces, before discarding the culture broth and storing the plates at -80 °C for a minimum of 24 h to freeze the biofilms. Before redox difference spectroscopy, the plates with biofilms were thawed at room temperature for a minimum of 15 min before suspending the biofilms in FW medium. A total of 96 biofilm samples (that is, 96 wells) were pooled together for each strain and the OD_{600} of the biofilm cell suspension was adjusted to 0.2 with FW medium. SDS (final concentration of 0.1% w/v) was then added to the cell suspension to lyse the cells and fresh 1 mM dithionite was added to the cell extract as a reducing agent anaerobically inside a glove bag (Coy Laboratories). The ultraviolet-visible spectrum from 350 to 650 nm of the reduced cell extract was then collected before and after oxidation with 2 mM ferricyanide using a Shimadzu UV-2401 spectrophotometer. The difference between the reduced and oxidized spectra at 552 nm (the α-Soret band in the reduced state) of the biofilm cell extracts in the WT and mutants (pilB, pilA and pilA-E5A) was used to estimate the overall biofilm cytochrome content (cells and biofilm matrix)⁵⁹. Spectra were also collected from cell extracts obtained from planktonic cells collected from exponential (OD₆₀₀. 0.4-0.5) cultures grown in FWAF at 30 °C. The total haem content of the cells was analysed using the alkaline pyridine haemochrome method⁶⁰. Statistically significant changes in cytochrome content relative to the WT were determined in pairwise comparisons using the t-test function of the Microsoft Excel software.

Cytochromes were also extracted with SDS from the biofilm matrix, isolated as described above and separated electrophoretically by SDS-polyacrylamide gel electrophoresis in 12% Mini-Protean TGX gels (Bio-Rad). Novex Sharp molecular-weight markers (Invitrogen) were used as standards. Haem-containing protein bands were stained with 3,3′,5,5′-tetramethylbenzidine, as previously described⁶¹. Replicate gels were run and stained with Coomassie blue to ensure that the total protein content per lane was comparable. When indicated, SDS-polyacrylamide gel electrophoresis and haem staining were also used to profile all haem-containing proteins within planktonic cells grown in FWAF to mid-exponential phase.

Pili purification and conductivity measurements by CP-AFM. Pili from WT or the Tyr3 mutant strain of G. sulfurreducens were purified as previously described²², except that all buffers contained 1 mM EDTA and all drying steps were carried out under a constant flow of filter-sterilized (0.22 µm) N2 gas. The pili were deposited on the surface of freshly cleaved HOPG for 30 min, then blotted dry, to probe the transversal conductivity of hydrated pili by conductive probe-atomic force microscopy (CP-AFM). The sample was scanned with the AFM tip in tapping mode to image the pili and to identify individual fibres. The CP-AFM tip was then positioned at different points along each pilus filament to measure its transversal conductivity while applying a bias voltage within the $\pm 1 \text{ V}$ range (3 nN force, 1 Hz rate). To ensure reproducibility, two to three current-voltage (I-V) curves were collected at each point, each pilus filament was probed in at least three positions, and four to five pilus fibres were probed for each strain. Furthermore, the tip was periodically moved to the HOPG surface adjacent to the pili to control tip quality. From the linear portion of each I-V curve, we estimated the electrical resistance (R), as described elsewhere 28, and used the average and s.d. of all the resistance values to compare the conductivity of the WT and Tyr3 pili.

Data availability. The authors declare that the data supporting the findings of this study are available within the article and its Supplementary Information files, or from the corresponding author upon request.

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Author contributions

R.J.S., S.L.-P. and G.R. designed the experiments. R.J.S. performed all the experiments except for the WT and Tyr3 pili purification and CP-AFM conductivity measurements, which were performed by S.L.-P. R.J.S. and G.R. analysed the data and co-wrote the paper. All authors discussed the results and commented on the manuscript.

Additional information

Supplementary Information accompanies this paper at http://www.nature.com/ naturecommunications

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Corrigendum: Mechanistic stratification in electroactive biofilms of *Geobacter sulfurreducens* mediated by pilus nanowires

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Two previous studies (Vargas *et al.* 2013, Liu *et al.* 2014) reporting that conductive pili are required for long-range electron transport in *Geobacter sulfurreducens* were inadvertently omitted from the reference list of this Article, and should have been cited in the Introduction section where the possibility that pili function as biofilm electron carriers is discussed.

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