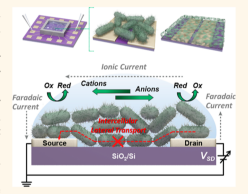


# Nanoelectronic Investigation Reveals the **Electrochemical Basis of Electrical Conductivity** in Shewanella and Geobacter

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Supporting Information

ABSTRACT: The electrical conductivity measured in Shewanella and Geobacter spp. is an intriguing physical property that is the fundamental basis for possible extracellular electron transport (EET) pathways. There is considerable debate regarding the origins of the electrical conductivity reported in these microbial cellular structures, which is essential for deciphering the EET mechanism. Here, we report systematic on-chip nanoelectronic investigations of both Shewanella and Geobacter spp. under physiological conditions to elucidate the complex basis of electrical conductivity of both individual microbial cells and biofilms. Concurrent electrical and electrochemical measurements of living Shewanella at both fewcell and the biofilm levels indicate that the apparent electrical conductivity can be traced to electrochemical-based electron transfer at the cell/electrode interface. We further show that similar results and conclusions apply to the Geobacter spp. Taken together, our study offers important insights into



previously proposed physical models regarding microbial conductivities as well as EET pathways for Shewanella and Geobacter spp.

KEYWORDS: electrogenic bacteria, microbial electrochemistry, bioelectrochemical system, bioelectronics, nanoelectronics, extracellular electron transfer, microbiome

issimilatory metal-reducing bacteria, such as Shewanella and Geobacter spp., harness energy through the metabolic oxidation of electron donors (organic sources/fuels) and subsequent electron transfer to insoluble electron acceptors (minerals). The electron transfer from Shewanella and Geobacter spp. to solid-state minerals or electrodes outside the cell, referred to as extracellular electron transport (EET), plays an important role in global biogeochemical cycles<sup>2</sup> and serves as the fundamental working principles for microbial biofuel productions<sup>3-6</sup> and microbial fuel cell (MFC) technologies.<sup>3,7-13</sup> Despite considerable interest, the mechanism for EET remains unresolved, with various possible pathways proposed. 9,14,15 Among them, one hypothesis suggests that long-range electron transport occurs through the electrically conductive biofilm, allowing multiple layers of cells in the biofilm to interact with electrode for respiration. This pathway is based on the unusual electrical conductivity measured in Shewanella and Geobacter spp., which is intriguing but controversial. To date, studies of the conductive mechanism of Shewanella have been inconclusive as most measurements were carried out under ex situ nonphysiological conditions. 16,17 For Geobacter, electrical measurements have been performed in living biofilms, although the accuracy and interpretation of the data are the subject of ongoing debate. 18-24 Two conflicting models have been suggested to explain the measured electrical conductivity in Geobacter nanowires and biofilms. One hypothesis suggests that electron conduction is redox activity (also referred to as multistep electron hopping or a superexchange mechanism), which occurs through electron hopping between adjacent cofactors in the form of a series of redox exchange reactions,

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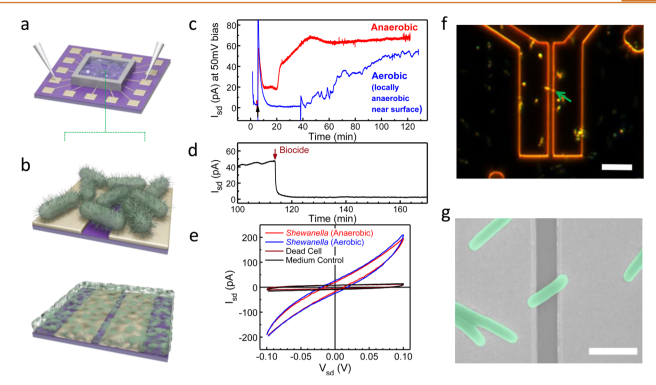


Figure 1. Electrical measurements of Shewanella oneidensis MR-1 under physiological conditions. (a, b) Schematic illustration of the nanoelectronic measurement setup with on-chip pair electrodes bridged by either individual MR-1 cells or biofilms. (c) Representative source—drain current  $(I_{sd})$  of living MR-1 as a function of incubation time on the device, under either aerobic or anaerobic conditions. The arrow indicates the injection of MR-1 cells into the measurement chamber. (d) Source—drain current  $(I_{sd})$  of dead MR-1 cells, the arrow indicates the biocide addition. (e) Representative  $I_{sd}$ — $V_{sd}$  transport characteristics of individual MR-1 cells, acquired after 2 h on-chip incubation. (f) In situ optical microscope (OM) image (dark-field mode) of MR-1 (indicated by green arrows) attached to the electrodes and bridging the gap. (g) Ex situ SEM image of MR-1 bridging the gap between pair electrodes (MR-1 cells are false colored for better visualization). Scale bars are 10 and 2  $\mu$ m in (f) and (g), respectively.

and is often accompanied by a redox gradient (a concentration gradient of oxidized or reduced cofactors). Another hypothesis posits an intrinsic metallic-like conductivity of Geobacter biofilms. In this model, electron transport occurs through delocalized electronic states along microbial nanowires, nanofilaments made of pilin proteins, with  $\pi$ - $\pi$  stacking of aligned aromatic moieties in pili. These two contrasting models imply significantly different EET mechanisms and entail distinct compositional and structural architectures of microbial nanowires and extracellular matrix involved in long-range electron transport. Therefore, a comprehensive understanding of the charge transport characteristics is critical for elucidating the functional and structural details and the fundamental conduction mechanism(s) of long-range electron transport in these electrogenic microbial systems.

In order to address this issue and to decipher the origins of the apparent electrical conductivity observed in the *Shewanella* and *Geobacter* systems, we investigated extracellular electron transport in detail at the nanoscale under physiologically relevant conditions. Standard on-chip nanoelectronic testing approaches offer ideal platforms for this purpose. However, the implementations of micro/nanoelectrode arrays to date have focused on measurements of either microbial nanowires under dry (nonphysiological) conditions<sup>16,17</sup> or electrochemical activity of actively respiring microbial systems. <sup>25,26</sup> Interdigitated electrode arrays have been employed for electrical measurements of living *Geobacter* biofilms, but with indirect electrochemical bipotentiostat measurements. <sup>21,22,24</sup> Here, we report systematic on-chip electrical transport studies of

Shewanella oneidensis MR-1 and Geobacter sulfurreducens PCA under physiological conditions using standard nanoelectronic approaches along with concurrent on-chip electrochemical measurements. These more comprehensive measurements elucidate the electrochemical origin of the apparent conductivity in Shewanella and Geobacter.

# **RESULTS AND DISCUSSION**

Figure 1a,b schematically illustrates the measurement setup and electrode layout used for probing the electrical transport in microbial cells and biofilms. Electrode pairs (source and drain) were prepatterned on a Si/SiO<sub>2</sub> wafer using electron-beam lithography (EBL), and a poly(dimethylsiloxane) (PDMS) chamber was employed to control exposure of electrodes to the microbial samples in the appropriate medium. To obtain better control of the local environment during measurements and to enable independent current recordings from multiple electrodes, an insulating and electrochemically inert polymer layer was employed to define a testing window through a secondary step of EBL. The confined window also enables the precise control of the exact electrode area exposed to the microbial samples, which is an important parameter that has not been thoroughly investigated in previous studies. The gap between the electrode pairs was set to be 1  $\mu$ m, which is smaller than the typical length of a MR-1 or PCA bacterium. This setup enables measurements of electron transport at the single-cell level, where an individual cell bridges the electrode pairs. We have also used relatively long electrode pairs (up to 3000  $\mu$ m) to maximize the numbers of bridging cells and thus to amplify the

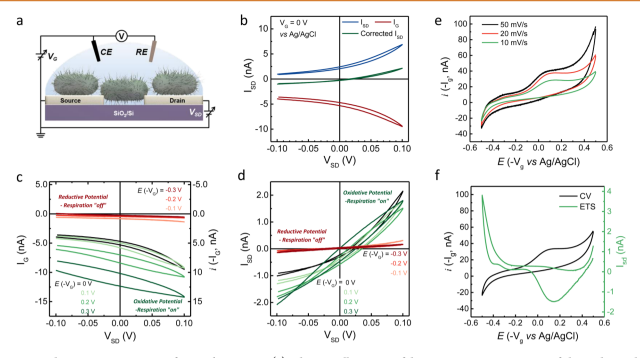


Figure 2. On-chip ETS measurements of S. oneidensis MR-1. (a) Schematic illustration of the concurrent measurements of electrochemical  $(I_{\rm g})$  and electrical transport  $(I_{\rm sd})$  characteristics of MR-1. (b) Representative  $I_{\rm sd}-V_{\rm sd}$  (blue) and  $I_{\rm g}$  (red) behavior of living MR-1 cells at 0 V gate voltage (vs Ag/AgCl), green curve represents the  $I_{\rm sd}$  curve after gate current correction. (c, d) Representative  $I_{\rm g}$  (c) and  $I_{\rm sd}-V_{\rm sd}$  (d) curves of living MR-1 at different gate voltages (from -0.3 to 0.3 V vs Ag/AgCl). Note that in terms of electrochemistry, the sign of the corresponding electrochemical potential (E) and Faradaic current (E) of MR-1 is opposite to the applied  $V_{\rm G}$  and measured  $V_{\rm G}$  as the source electrode (working electrode) is set at ground. (e) In device CV of MR-1. (f)  $V_{\rm g}-V_{\rm g}$  (CV, black curve) and corresponding  $V_{\rm g}-V_{\rm g}$  (ETS, green curve) characteristics of a typical MR-1 device.

relatively small current signals. The details of device fabrication and measurement setup are described in the Methods section.

The measurement of S. oneidensis MR-1 was conducted under both aerobic and anaerobic conditions. Typically, the MR-1 cell culture was injected into the PDMS chamber after a stable baseline was established in media solution. The sourcedrain current  $(I_{sd})$  was continuously recorded at the rate of 2 Hz with a constant bias voltage  $(V_{sd})$  of 50 mV. The current time (I-t) characteristics of S. oneidensis MR-1 in aerobic and anaerobic conditions are shown in Figure 1c. As expected, increased I<sub>sd</sub> over time was observed under both conditions after introduction of the MR-1 culture. The *I*-*t* characteristics of MR-1 were slightly different in these two different environments. In anaerobic measurements, a jump in current was observed immediately, followed by a faster increase. For measurements under aerobic conditions, the current remained negligible (i.e., the same as that in media background) for ca. 40 to 60 min before it started to increase gradually (note that the sharp current "spike" just after the injection of the MR-1 culture is not correlated to the MR-1, but is due to the disturbance and reformation of the electrical double layer on the electrodes). The delayed current increase is likely due to the presence of dissolved oxygen. Assuming  $I_{sd}$  is correlated to the EET process of MR-1, either oxygen acts as an electron acceptor in the EET process, or in the presence of oxygen, EET is not involved in MR-1 metabolism. When oxygen is consumed near the electrode surface, a local anaerobic environment is established for the subsequent electron transfer to electrode and thus current generation. For both conditions, the current reached steady state after about 2 h of MR-1 exposure, indicating a similar (locally anaerobic) condition of MR-1 cells attached onto the electrodes. This result is further

confirmed by the similar current-voltage (I-V) characteristics of MR-1 measured at steady state under both aerobic and anaerobic conditions (Figure 1e). By "steady state" we mean here that the currents have become stable and do not vary with time. These  $I_{\rm sd}$ - $V_{\rm sd}$  characteristics offer valuable insights into the charge transport dynamics of actively respiring S. oneidensis MR-1. After the  $I_{\rm sd}$  current was observed, an injection of 0.25% glutaraldehyde as a biocide<sup>26</sup> completely quenched the current (Figure 1d), verifying that the conducting current measured in this system originates from living MR-1 cell(s). Furthermore, in situ dark-field optical imaging during the electrical measurements was performed, MR-1 was observed to swim to the chip surface and was immobilized gradually on the electrode (some also landed and stayed between the source and drain electrodes, as shown in Figure 1f). This result confirmed that the observed  $I_{\rm sd}$  is correlated with the attachment of MR-1 cells onto the electrodes. The scanning electron microscope (SEM) images of the device after the measurements (Figure 1g) show individual MR-1 cells attached onto the electrodes and bridging the gap. Bacterial nanowires were not observed during the measurements, as determined by both in situ optical microscopy and ex situ SEM characterization (Figure 1f,g). Our results indicate that apparent charge transport is observed in MR-1 cells even without conductive bacterial nanowires. 16,17 When the device was exposed to MR-1 for a longer period of time (12 to 24 h), the  $I_{\rm sd}$  further increased to another plateau, and ex situ SEM measurements confirmed the formation of a thin biofilm layer (with a typical thickness between 1 to 3  $\mu$ m). Similar I-Vcharacteristics with larger amplitude (as compared to the 2 h sample) were observed for the Shewanella biofilm (see Supporting Information Figure S1).

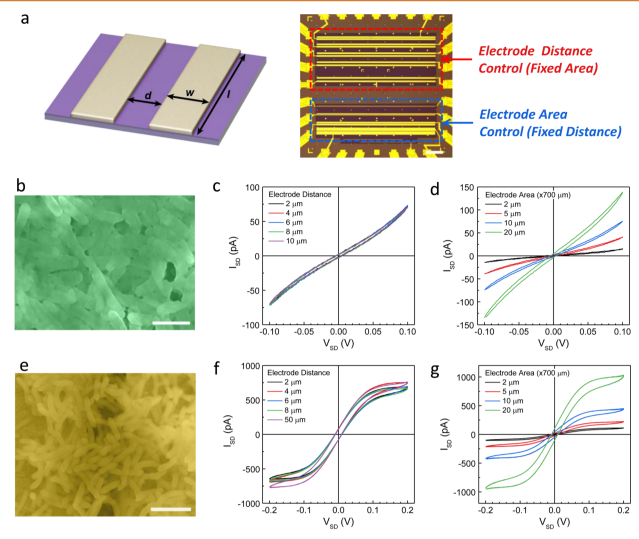


Figure 3. Nanoelectronic investigation of the "electrical conduction" of Shewanella and Geobacter biofilms using electrode pairs with varying gaps and areas. (a) Schematic illustration of the biofilm measurements using two sets of pair electrodes with either varying gaps (with fixed areas) or varying electrode areas (with fixed gaps). (b) Representative SEM image of a Shewanella biofilm. (c, d)  $I_{\rm sd}-V_{\rm sd}$  behavior of living Shewanella biofilms with different pair electrode distances (c) and different electrode areas (d). (e) Representative SEM image of a Geobacter biofilm grown under anaerobic conditions. (f, g)  $I_{\rm sd}-V_{\rm sd}$  behavior of living Geobacter biofilms with different electrode distances (f) and different electrode areas (g). Scale bars in (b) and (e) are 2  $\mu$ m.

The above electrical transport measurements show apparent electrical conductivity in S. oneidensis MR-1 similar to previous studies in electrogenic microbial systems.  $^{16,17,20-22}$  To gain further mechanistic insight and to elucidate the origins of such electrical conductivity, we used electrical transport spectroscopy (ETS) $^{27}$  to probe the microbial system under physiological conditions. The ETS approach introduces an additional reference electrode (RE) and a counter electrode (CE) into the nanoelectronic on-chip measurement platform (Figure 2a). Analogous to a conventional field-effect transistor (FET), the counter electrode functions as a gate electrode, and the reference electrode (Ag/AgCl) is used to establish the electrochemical potentials of MR-1. This platform enables concurrent measurements of the gate current ( $I_{\rm g}$ ) and the electrical transport current ( $I_{\rm sd}$ ) in microbial samples (see Methods).

Our measurements show that the electrical transport current  $(I_{\rm sd})$  (blue line in Figure 2b) and gate current  $(I_{\rm g})$  (red line in Figure 2b) exhibit comparable amplitudes, but with opposite polarities. While  $I_{\rm sd}$  is considered the electrical transport

current, I<sub>o</sub> is generally recognized as "leakage current" in a typical FET measurement. Such leakage currents are undesirable for both solid-state FETs and electrolyte-gated FETs, as the electrical transport  $(I_{sd})$  is expected to be altered only by the electrical field across a dielectric layer for a solid-state FET, or an electrolyte double layer for an electrolyte-gated FET; negligible current should be observed through the gating channel. The considerable  $I_{\rm g}$  observed in MR-1 can be attributed to the Faradaic current (i.e., redox current) originating from electron transfer at the cell/electrode interface.<sup>27</sup> This conclusion is also supported by the negligible  $I_{\sigma}$  observed in the free medium control (see Supporting Information Figure S4). The comparable  $I_{\sigma}$  and  $I_{sd}$  from living MR-1 indicate that electrochemical current  $(I_g)$  contributes at least partly to the electrical transport current  $(I_{sd})$ . With an  $I_g$  of comparable amplitude, the measured  $I_{sd}$  is convoluted with  $I_{g}$ . We have previously shown in our ETS studies that the contributions of  $I_g$  to  $I_{sd}$  can be properly deducted by considering the equivalent circuit (see Supporting Information Figure S5) to obtain  $I_{sd}$  accurately (green curve in Figure 2b).<sup>27</sup>

Together, our results provide the first example of direct and concurrent measurements of the electrochemical  $(I_{\rm g})$  and electrical transport  $(I_{\rm sd})$  currents of *S. oneidensis* MR-1 under physiological conditions, enabling a more comprehensive and conclusive analysis of the system.

The above discussions indicate that electron transport within MR-1 is closely correlated to electrochemical processes. More detailed analyses of the results further support this hypothesis. When the gate voltage is set negative ( $\nu$ s a Ag/AgCl reference), the actual electrochemical potentials of MR-1 are more oxidizing, leading to enhanced digestion (oxidation) of electron donors and and increase in  $I_{\rm g}$  (absolute value) (Figure 2c respiration "on" state). With a positive gate voltage ( $\nu$ s Ag/AgCl), MR-1 cells are pinned at a more reducing electrochemical potential. This situation suppresses metabolic oxidation and significantly reduces  $I_{\rm g}$  (Figure 2c, respiration "off" state).

The  $I_{\rm sd}$ - $V_{\rm sd}$  characteristics show similar trends to that of  $I_{\sigma}$ (Figure 2d). Specifically, when MR-1 is pinned at an oxidative potential, the device shows considerably increased  $I_{\rm sd}$  (larger slope in  $I_{sd}$ – $V_{sd}$  curve), demonstrating that the electrical transport current depends strongly on the electrochemical states of MR-1. Hence, a small change in  $V_{\rm sd}$  can produce a large modulation in  $I_{\rm sd}$  and a large  $I_{\rm sd}$ - $V_{\rm sd}$  slope. In contrast, when the MR-1 is set at a reductive potential and respirationoff state, the modulation of  $I_{\rm sd}$  by  $V_{\rm sd}$  is much smaller, and thus the  $I_{\rm sd}$ - $V_{\rm sd}$  slope is smaller (Figure 2d). Given that  $V_{\rm sd}$  actually modulates the electrochemical potentials  $(V_{\rm g})$ , this behavior is consistent with our hypothesis that the overall electron conductive current  $(I_{sd})$  originates at least partially from the electrochemical current. Another approach for ETS investigation is to perform on-chip cyclic voltammetry (CV) of MR-1 while monitoring the corresponding change of their conductivity (by measuring  $I_{sd}$  at a constant  $V_{sd}$ ). As shown in Figure 2e,f, typical CV characteristics of MR-1 in electrondonor-rich media were observed on-chip (Figure 2e), but unlike a conducting material system that has distinguishable ETS  $(I_{\rm sd}$ - $V_{\rm g})$  characteristics in response to the electrochemical processes, <sup>27</sup> MR-1 shows an ETS current  $(I_{sd})$  that is inversely proportional to the corresponding CV current  $(I_{\sigma})$ , which provides additional evidence for the electrochemical origins of the conducting current measured from MR-1.

Note that electrochemically generated current has not been previously considered responsible for the electrical transport current observed in such microbial systems. A major reason is that it is relatively difficult to separate out the electrochemical current, especially under physiological conditions. Our on-chip ETS approach enables separate and concurrent measurement of Faradaic current and electrical transport current from living MR-1 and is thus well suited for such investigations.

To test our hypothesis more conclusively and quantitatively, inspired by the ETS measurements of actively respiring MR-1, we investigated the dependence of electrical conduction ( $I_{\rm sd}-V_{\rm sd}$ ) on the electrode gap distance and the area of the electrodes. These parameters provide key insight into the conduction mechanism in typical nanoelectronic studies. In metal- or semiconductor-based conductive pathways (as suggested in recent debates on the studies of Geobacter), the measured currents should be inversely proportional to the distance between the conducting channels and be independent of the electrode area, whereas electrochemically related currents are expected to have the opposite correlation, *i.e.*, to be proportional to the electrode area and independent of electrode

distance. Therefore, the  $I_{\rm sd}$ - $V_{\rm sd}$  dependence on the pair electrode distance and area can be used to probe the role of electrochemical current directly in the electron conduction measurement and to elucidate unambiguously the quantitative contribution from the electrochemical current to the overall electrical transport current. To this end, we have designed electrode arrays with varying electrode areas and electrode distances (Figure 3a) and conducted systematic electrical transport studies. Surprisingly, I<sub>sd</sub> from MR-1 (either individual cells or biofilms) exhibits no dependence on the pair electrode distance but shows strong correlation with the electrode area (Figure 3c,d and Supporting Information Figure S6). This observation offers direct evidence that the electrical conduction current (I<sub>sd</sub>) from MR-1 largely originates from the electrochemical current that is generated at the cell/electrode interface.

Due to the paucity of documented electrical measurements of living Shewanella for reference, we have conducted analogous investigations on biofilms of Geobacter sulfurreducens PCA (Figure 3e). Therefore, previously reported results/models on the G. sulfurreducens can be used to cross-reference the data and the conclusions obtained in this study. Similarly, the PCA biofilms were grown in situ on the device by a previously developed anode growth method<sup>20,21</sup> and studied under physiological conditions using the same approach described above. The  $I_{\rm sd}$ - $V_{\rm sd}$  measurement of Geobacter biofilms also shows that current is not dependent on the electrode distance but is only related to electrode area (Figure 3f,g), indicating the same interfacial electrochemistry model that we proposed for Shewanella. These results argue against the previously suggested superexchange model and metal/semiconductor model in which the current should be inversely proportional to the distance between the two electrodes. Also, note that the I-Vcharacteristics observed for the Geobacter biofilms show sigmoidal shapes (distinct from that of Shewanella) with steady-state currents at high voltages. Although the superexchange hypothesis where the current is driven by a redox gradient (first developed to describe the charge transport in electroactive polymers with fixed molecular redox sites)<sup>28,29</sup> can also be used to explain the sigmoidal I-V characteristics of Geobacter biofilms,  $^{21,22,24}$  it does not explain the different I-Vcharacteristics observed for Shewanella or the lack of distance dependence of the Geobacter's I-V behavior.

The lack of distance dependence and the distinct area dependence of the measured current strongly suggest that the electrical conduction from Shewanella and Geobacter are dominated by the electrochemical current generated at the cell/electrode interfaces. The transport current across the source and drain electrodes is attributed to the different electrochemical current (turnover rate) at the source and drain electrodes due to slightly different potentials (resulted from the  $V_{
m sd}$  bias), which leads to an ionic current across the sourcedrain electrodes to balance the charge. As similar nanoelectronic fabrication techniques and investigations have not previously been employed for microbial studies, our study suggests that this approach can elucidate important factors and insights that were missed previously. In particular, the area of electrodes exposed to the living bacteria/biofilm and/or medium environment should be carefully controlled in such investigations, and the current dependence on the electrode distance should be investigated for proposed transport mechanisms.

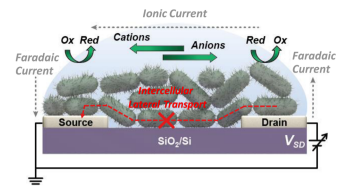


Figure 4. Schematic illustration of the interface electrochemistry model for the electrical conducting current in a typical electrode pair measurement. The transport current is determined by the vertical electron transfer (electrochemical/Faradaic current) at the bacteria/electrode interfaces, whereas lateral (non-Faradaic) electron transport pathway across the biofilm does not exist. Ionic transport (current) toward the electrodes in the lateral direction forms a complete electrochemical circuit.

Figure 4 schematically illustrates the interface electrochemistry model. The transport current across the sourcedrain electrodes is limited by Faradaic electron transfer at the bacteria/electrode interface (electrochemical current) and is coupled with ionic transport current in the lateral direction to form a complete electrochemical circuit. On the other hand, the previously suggested electron transport pathway across the biofilm, either by electron hopping through a network of redox centers or via the intrinsic metallic-like conductivity, does not exist or does not play a significant role in the system we are studying here. Additional results that are consistent with our conclusions are the observation of  $I_{\rm sd}$ - $V_{\rm sd}$  current in the case where cellular bridges have not formed to connect the pair electrodes physically (Supporting Information Figure S6). Once the origins of the conducting current from these microbiomes are clarified, their measurement can be used to elucidate the mechanism of the corresponding EET process. For instance, with the interface electrochemistry model, the long-range electron transport across the electrically conductive biofilm thickness (through either the metallic pili mechanism or the redox matrix mechanism) has been called into question. Therefore, this model weighs in favor of two redox-based EET mechanisms: direct contact (via membrane cytochromes) or "electron shuttling" (via secreted mediators).

It has been previously suggested that Shewanella and Geobacter undergo different EET pathways. For Shewanella, extracellular electrons are transferred to the electrode via cellsecreted flavins that diffuse to the electrode surface and act as redox mediators (*i.e.*, the indirect "electron shuttling" mechanism);  $^{25,30-32}$  for *Geobacter*, on the other hand, electrons are directly transferred to the electrode through outer membrane c-type cytochrome (c-Cyts) protein that is in physical contact with the electrode surface (direct contact mechanism).  $^{33,34}$  Interestingly, the distinct conductive I-Vcharacteristics observed for Shewanella and Geobacter in this study can be rationalized with the different EET mechanisms in these two model bacteria. Specifically, the sigmoidal I-Vcharacteristic (Figure 3f) obtained from Geobacter can be explained by the combination of the direct electron transfer mechanism and our interfacial electrochemistry model. With relatively efficient electron transfer kinetics at the cytochrome/ electrode interface, the overall electron transfer rate and,

therefore, the Faradaic current at relatively high bias are limited by the catalytic rate of the enzyme/electrode system or the diffusion-limited supply of the redox species, 35,36 thus leading to a steady-state current that is independent of the bias voltage at relatively higher potentials and sigmoidal I-V behavior of Geobacter (following the Nernst-Monod equation). 35,36 As for Shewanella, the current generation is limited by the relatively slow kinetics of the interface electron transfer (i.e., redox of flavin on the electrode), and thus its I-V behavior follows the Butler-Volmer equation 35,36 and increases with bias voltage without apparent saturation in the measurement range. Therefore, our interface electrochemistry model for the mechanism of microbial electrical conductivity, along with corresponding experimental observations, provide additional evidence and important insight into the EET mechanisms of both Shewanella and Geobacter systems and might also shed light on the fundamental basis of electron transport in other microbial systems.<sup>37</sup> Future investigations of genetically engineered microbial systems (e.g., Shewanella strains lacking specific functional proteins via selective gene deletion) could offer further insight into the charge transport mechanisms in these microbial systems.

# **CONCLUSIONS AND PROSPECTS**

We have employed nanoelectronic fabrication and measurements to elucidate the origins of the electrical conducting current from two model electrogenic microbial systems, S. oneidensis MR-1 and G. sulfurreducens PCA. The electrical characteristics of each strain were obtained under physiological conditions together with in situ optical imaging and other onchip measurements. We have shown that the conductive current from these microbes originates from electrochemical electron transfer at the cell/electrode interface, and no lateral electron transport across the cell or biofilm was observed. The experimental data reported in this study indicate that results and the interpretations made in earlier experimental and theoretical studies should be revisited, particularly paying attention to electrode distance/area dependence (which has not always been measured or reported). Furthermore, our interface electrochemistry model provides valuable insight into the EET mechanisms of both model microbial systems. From a broader perspective, these nanoelectronic testbeds and investigations reveal important factors and insights into microbial studies. These tools will accelerate our basic understanding of earth's microbiomes and harness the capabilities of microbial ecosystems. 38-41

## **METHODS**

Growth of Shewanella oneidensis MR-1 and Geobacter sulfurreducens PCA. S. oneidensis strain MR-1 was grown aerobically in LB broth (20 g/L, 20 mL, Sigma-Aldrich) with gentle shaking (200 rpm) at 30 °C for approximately 12 h. The cells were then redispersed into fresh LB broth at a volume ratio of 1:100. The redispersed culture was shaken (200 rpm) aerobically for approximately 6 h at 30  $^{\circ}$ C (up to an OD<sub>600</sub> of 0.5) before measurements. G. sulfurreducens strain PCA was grown in a continuous flow bioreactor (BioFlo 110; New Brunswick Scientific) with a medium inflow rate of 0.4 mL/min and a working volume of 1 L. Sodium acetate (20 mM) was used as electron donor and sodium fumarate (480 mM) as electron acceptor. In addition, the growth medium contained the following chemicals as nutrients (in mg/L): NH<sub>4</sub>Cl, 200; NaH<sub>2</sub>PO<sub>4</sub>· H<sub>2</sub>O, 69; KCl, 380; CaCl<sub>2</sub>·H<sub>2</sub>O, 40; MgSO<sub>4</sub>·7H<sub>2</sub>O, 200, nitrilotriacetic acid, 15; MnCl·4H<sub>2</sub>O, 10; FeSO<sub>4</sub>·7H<sub>2</sub>O, 5; CoCl<sub>2</sub>·6H<sub>2</sub>O, 2; ZnCl<sub>2</sub>, 1; CuSO<sub>4</sub>·5H<sub>2</sub>O, 0.3; AlK(SO<sub>4</sub>)<sub>2</sub>·12H<sub>2</sub>O, 0.05; H<sub>3</sub>BO<sub>3</sub>, 0.05; Na<sub>2</sub>MoO<sub>4</sub>,

0.9; NiCl<sub>2</sub>, 0.5; Na<sub>2</sub>WO<sub>4</sub>·2H<sub>2</sub>O<sub>7</sub>, 0.2; Na<sub>2</sub>SeO<sub>4</sub>, 1. The culture media were maintained at 25 °C and at pH = 7. A mixture of N<sub>2</sub>·CO<sub>2</sub> (80:20) was constantly flowing into the reactor (0.4 mL/min) to remove any traces of oxygen and to maintain a strictly anaerobic environment. The culture in the reactor was stirred at a constant speed of 50 rpm. The PCA cell culture in the flow reactor was kept at a steady state with a measured OD<sub>600</sub> of 0.6 before measurements.

**Fabrication of the Pair Electrode Device.** Pair electrodes (as source and drain) on the substrate ( $p^{+2}$  silicon wafer with 300 nm thermal oxide) were defined by EBL, followed by evaporation of 20 nm Ti and 50 nm Au. To eliminate the influence of electrolyte and to avoid electrochemical reactions on the metal electrodes, a passivation layer of poly(methyl methacrylate) (PMMA, ~500 nm thick, electrochemically inert) was spin-coated on the device. Windows that only exposed desired areas and electrodes to the samples were opened by EBL.

**Electrical Measurement and** *in Situ* **Optical Imaging of Living Cells.** A source measurement unit (SMU, Agilent B2902a) was used for all microbial measurements. For I-t measurements, sourcedrain current was recorded at a rate of 2 Hz, with a small bias voltage of 50 mV. Typically, the MR-1 cell culture was injected into the PDMS chamber after a stable baseline was established in medium solution. For I-V measurements, the source—drain voltage was swept between -0.1 and 0.1 V at a typical rate of 10 mV/s to minimize the capacitive charging current (appeared as hysteresis in I-V curve). All anaerobic measurements were carried out in an anaerobic hood (Coy Anaerobic Chamber, Type B, Vinyl) with an in-hood probe station. *In situ* optical imaging was carried out with an upright microscope (QImaging Retiga 2000R) operated under dark-field.

**Gating and ETS Measurement.** The two-channel SMU was used for the gate measurement (one as a gate channel and one as a sourcedrain channel). A four-probe configuration was used with a gate channel to function as a potentiostat. The potential of the drain electrode was controlled with respect to the reference electrode ( $V_g$  vs Ag/AgCl), while the current ( $I_g$ ) was collected and measured through the counter electrode. The source—drain channel was used either to supply a small bias potential (50 mV) between source and drain electrodes and collect the corresponding current ( $I_{\rm sd}$ ) during a  $V_g$  sweep or to sweep the source—drain voltage ( $V_{\rm sd}$ ) at an applied gate voltage ( $V_g$ ) to obtain the  $I_{\rm sd}-V_{\rm sd}$  curve. The equivalent circuit of the ETS measurement is illustrated in Supporting Information Figure S5.

# **ASSOCIATED CONTENT**

## S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.6b03655.

Additional I-t,  $I-V_g$ , and I-V measurements (PDF)

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## **Author Contributions**

OThese authors contributed equally. M.D., H.-Y.S., G.W., P.S.W., G.C.L.W., K.H.N., Y.H., and X.D. designed the research; M.D. conducted the electrical measurement with the assistance of H.-Y.S.; H.-Y.S. and S.-H.L. prepared the microbial cultures with the assistance of C.K.L., M.D., G.W., and T.D.Y.; M.D. and X.D. analyzed the data and wrote the paper with the assistance of all the authors; P.S.W., G.C.L.W., K.H.N., Y.H., and X.D. supervised the research.

## **Notes**

The authors declare no competing financial interest.

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