Genetic Control of Radical Cross-linking in a Semisynthetic Hydrogel

Austin J. Graham, Christopher M. Dundas, Alexander Hillsley, Dain S. Kasprak, Adrianne M. Rosales,* and Benjamin K. Keitz*

ABSTRACT: Enhancing materials with the qualities of living systems, including sensing, computation, and adaptation, is an important challenge in designing next-generation technologies. Living materials address this challenge by incorporating live cells as actuating components that control material function. For abiotic materials, this requires new methods that couple genetic and metabolic processes to material properties. Toward this goal, we demonstrate that extracellular electron transfer (EET) from Shewanella oneidensis can be leveraged to control radical cross-linking of a methacrylate-functionalized hyaluronic acid hydrogel. Cross-linking rates and hydrogel mechanics, specifically storage modulus, were dependent on various chemical and biological factors, including S. oneidensis genotype. Bacteria remained viable and metabolically active in the networks for at least 1 week, while cell tracking revealed that EET genes also encode control over hydrogel microstructure. Moreover, construction of an inducible gene circuit allowed transcriptional control of storage modulus and cross-linking rate via the tailored expression of a key electron transfer protein, MtrC. Finally, we quantitatively modeled hydrogel stiffness as a function of steady-state mtrC expression and generalized this result by demonstrating the strong relationship between relative gene expression and material properties. This general mechanism for radical cross-linking provides a foundation for programming the form and function of synthetic materials through genetic control over extracellular electron transfer.

KEYWORDS: living materials, synthetic biology, extracellular electron transfer, hydrogels

INTRODUCTION

Nature uses hierarchical and genetically encoded instructions to construct functional materials with specific self-assembly, regulatory, healing, and morphological properties.1 Inspired by such processes, engineered living materials (ELMs) employ the autonomy of living cells to synthesize and control material structures across multiple scales with user-designed functions that are directly coupled to gene expression.2–5 Living materials containing microbes, including biofilms, bacterial cellulose, curli fibers, and synthetic gels loaded with bacteria, are of prominent interest due to their potential application in tissue engineering, 3D printing, soft robotics, metabolic engineering, and living sensors.6–10 Bacteria are particularly attractive as ELM components due to their natural sensing capabilities and programmability. For example, engineered bacteria can act as cellular actuators integrated within the ELM, tailoring its synthesis and function through over-expression, mutagenesis, and gene circuitry.

Not surprisingly, the majority of ELMs rely on materials natively produced by the host organism. For example, several amyloid-based materials have been synthesized by genetically tractable bacteria, such as aggregates of CsgA in Escherichia coli11,12 and TasA in Bacillus subtilis.13 Genetic fusions have allowed these fibrous matrices to bind specific molecules, conduct electricity, perform catalytic reactions, and adhere to complex surfaces.14–17 Apart from amyloids, extracellular polymerization of bacterial cellulose has been engineered using quorum sensing circuits and mutagenesis to create sturdy materials for tissue engineering and sensing applications.18,19 Despite these advances, significant drawbacks of natural materials include their limited chemical functionality, robustness, homogeneity, and scalability compared to engineered synthetic materials.20 For example, manufactured soft materials such as polymers and hydrogels are easily functionalized and versatile, facilitating their adoption in diverse environments. However, synthetic materials largely lack the dynamic adaptability and environmental responsiveness found in natural systems. Introducing these qualities to synthetic materials could synergistically enhance ELMs and enable new applications that combine the precision and chemical diversity of engineered materials with the autonomy and evolvability of living cells. However, such designs will require methods for...
bacteria to control synthetic material properties at the genotypic level. Similarly, robust transcriptional control and quantitative prediction of the relationship between gene expression and material properties are needed for ELMs to approach the design precision of engineered materials.

Toward this goal, we recently developed a cell-controlled radical polymerization reaction using extracellular electron transfer (EET) from the organism *Shewanella oneidensis*.21 EET is a form of anaerobic respiration that couples carbon oxidation to the reduction of extracellular metals or metal oxides.22 In *S. oneidensis*, the Mtr protein pathway is the primary source of EET flux in anaerobic *S. oneidensis* metabolism (Figure 1a). Specifically, the outer membrane cytochromes MtrC and OmcA are terminal reductases in this pathway and responsible for direct electron transfer onto extracellular metal species such as Fe and Cu.21,23 MtrF is a homologue to MtrC and can similarly reduce a variety of metals.24 In our previous polymerization platform, we redirected EET flux to a metal catalyst, which generated a radical from a halogenated initiator and cross-links acrylate-based functional groups.21 EET is a form of anaerobic respiration that couples carbon oxidation to the reduction of extracellular metals or metal oxides.22 In *S. oneidensis*, the Mtr protein pathway is the primary source of EET flux in anaerobic *S. oneidensis* metabolism (Figure 1a). Specifically, the outer membrane cytochromes MtrC and OmcA are terminal reductases in this pathway and responsible for direct electron transfer onto extracellular metal species such as Fe and Cu.21,23 MtrF is a homologue to MtrC and can similarly reduce a variety of metals.24 In our previous polymerization platform, we redirected EET flux to a metal catalyst, which controlled polymerization governed by the atom-transfer radical polymerization (ATRP) mechanism. Importantly, we demonstrated that control over polymer production was directly coupled to cell metabolism and genetically encoded through these EET proteins. Since ATRP is a versatile platform for soft material synthesis,25 we hypothesized that EET-powered catalysis could be extended to control radical cross-linking in a synthetic hydrogel. While there are numerous examples of incorporating live cells into polymer networks, network properties such as cross-link density, mesh size, degradation, and elastic modulus have generally been designed independent of cell activity. In addition, previous attempts to incorporate cells as live cross-linking agents in synthetic hydrogels have relied on the activity of glucose peroxidase or extracellular functionalization of cells after growth,26−30 which compromise cell viability through the creation of toxic reactive oxygen species or are not under cellular control. Cell-free gelation systems using bacterial lysates31 or secreted recombinant peptides32 have also been explored, but removing the living component prevents continued material responsiveness. We envisioned that controlling radical cross-linking via EET gene expression would capitalize on the programmability of bacteria and enable the use of stimuli-responsive synthetic biology circuits to control material function.

Here, we demonstrate that EET from *S. oneidensis* can be used to control radical cross-linking of a semisynthetic methacrylated hyaluronic acid (MeHA) hydrogel (Figure 1a). First, we show that EET is required for gelation, and that organisms without this metabolic capability (i.e., *E. coli*) are unable to cross-link gels on a comparable time scale or in a controllable manner. Gels did not form unless a constant source of electron flux, radical initiator, and metal catalyst were present. Additionally, the facultative metabolic capability of *S. oneidensis* enabled cross-linking under benchtop conditions without dedicated oxygen removal. Analysis of cell motility and metabolic activity revealed that bacteria remain viable and responsive in the gels for a minimum of 1 week and that the
degree of cross-linking by EET affected cell movement. Next, we found that cross-link density was a strong function of bacterial genetics, as cytochrome knockout strains synthesized gels more slowly and with decreasing stiffness correspondent to the number of removed EET genes. Finally, transcriptional circuits based on controlling the expression of mtrC with the LacI repressor enabled tunable cross-linking rates and hydrogel mechanical properties. We found that hydrogel storage modulus fit well to inducible gene expression models, directly linking steady-state gene expression to a quantifiable and macroscopic material property. Overall, our results suggest that transcriptional control over EET can be used to predictably interface the properties of living systems with potentially any material amenable to radical cross-linking.

■ RESULTS

Extracellular Electron Transfer from Live S. oneidensis Controls Aerobic Radical Cross-linking. To initially validate our hypothesis that EET-controlled ATRP could be used to form a cross-linked hydrogel, we first synthesized a 65% methacrylated hyaluronic acid (MeHA) macromer using an established protocol (Figure 1b, Figure S1). Hyaluronic acid is a common naturally derived biomaterial platform that is attractive for our application due to its biocompatibility and chemical versatility. The high density of functional groups was chosen to increase likelihood of successful cross-linking and to minimize the effect of radical scavenging by oxygen. In initial experiments, MeHA was dissolved at 3 wt % in Shewanella basal medium (SBM) supplemented with 0.05% w/v casamino acids (Tables S2−3), and the dissolved macromer was mixed with a radical initiator, 2-hydroxyethyl 2-bromoisobutyrate (HEBIB, 500 μM), a copper ATRP catalyst with Tris(2-pyridylmethyl)amine ligand (Cu-TPMA, 10 μM), and inoculated with anaerobically pregrown S. oneidensis MR-1 cells (initial OD600 = 0.2 in the gel, ca. 10⁸ CFU·mL⁻¹). Lactate (20 mM) was the electron donor and fumarate (40 mM) was the primary electron acceptor. Gels were initially tested in anaerobic environments to promote EET metabolism and eliminate any effect of oxygen quenching. After mixing, the solution was placed in a humidified anaerobic chamber and monitored via inversion testing. After 2 h, solutions containing S. oneidensis cross-linked to form polymer networks,

Figure 2. Living hydrogel materials cross-linked by S. oneidensis MR-1 can be chemically or biologically tuned. (a) Storage moduli of hydrogels cross-linked without various components, measured by rheology after 2 h of cross-linking. Many gels did not form and could not be characterized. (b) Storage moduli of hydrogels cross-linked for 2 h with varying concentrations of Cu-TPMA (catalyst); one-way ANOVA p = 0.018. (c) Storage moduli of hydrogels cross-linked for 2 h with varying concentrations of HEBIB (radical initiator); one-way ANOVA p = 0.0027. (d) End-point storage moduli at 2 h and (e) in situ rheology of hydrogels cross-linked with varying optical densities of S. oneidensis MR-1 in the reaction, and therefore varying total electron transfer; one-way ANOVA p = 0.0002. Gels did not form using OD₆₀₀ = 0.05 cells in either experiment. (a−d) Data are shown as mean ± SEM, n = 3 biological replicates.
whereas solutions containing E. coli did not (Figure 1b,c). Consistent with our previous results,21 these data suggest that electron flux to the metal catalyst from EET-based metabolism is required for radical generation and cross-linking.

After successful preliminary cross-linking reactions, we more thoroughly investigated the mechanical properties of the gels using shear oscillatory rheology. Gel precursor solutions were inoculated with cells and placed between two hydrophobically treated glass slides with a silicone spacer, allowed to cross-link, and swollen overnight at room temperature in 1X PBS. Gel mechanical properties were then quantified on a rheometer by measuring storage and loss moduli. Storage modulus ($G'$) is a measure of the elastic response to applied shear, wherein the network recovers. In contrast, loss modulus ($G''$) is a measure of viscous, nonrecoverable dissipation of applied shear. The storage modulus of biological tissues and common biomaterial platforms, such as extracellular matrix mimics, typically range from 0.01 to 10 kPa.35 Storage and loss moduli for EET-cross-linked gels were determined from the linear viscoelastic regime as determined by strain and frequency sweeps (Figure S2) and calculated using a 0.01 to 100 Hz frequency sweep at 0.1% strain. Gels formed in anaerobic environments exhibited a storage modulus of $\sim$2500 Pa and a loss modulus of $\sim$20 Pa (Figure S3) and were predominately elastic, as indicated by significantly larger $G'$ compared to $G''$. Taking advantage of its facultative metabolism, we next tested radical cross-linking of hydrogels by S. oneidensis under ambient as opposed to anaerobic conditions. Using the same reagent concentrations as above with aerobically pregrown cells (henceforth, standard conditions), S. oneidensis formed cross-linked networks at 30 °C without dedicated oxygen removal. Gels formed in both aerobic and anaerobic environments yielded comparable mechanical properties, indicating insignificant oxygen inhibition under ambient conditions (Figure S3). Aerobic gels prepared by S. oneidensis were also mechanically similar to acellular gels cross-linked using UV light and the photoinitiator lithium phenyl-2,4,6-trimethylbenzoylphosphinate (LAP, 500 μM) (Figure S4). In addition, controls lacking (a) EET-active bacteria, (b) radical initiator, (c) metal catalyst, or (d) methacrylate functional group did not form measurable gels within 2 h (Figure 2a, Figure S5). Methacrylate functional group conversion in the hydrogels was also measured by $^1$H NMR spectroscopy after hyaluronidase digestion, confirming the cross-linking mechanism (Figure S6). Together, these results demonstrate that EET-cross-linked hydrogels can be synthesized under ambient or anaerobic conditions using electroactive bacteria to form mechanically robust networks that are typical of this macromer.

The cross-linking kinetics and mechanical properties of polymer networks strongly depend on a variety of chemical factors such as catalyst and ligand identity, rate of initiation, and initiator structure.36 Thus, we next explored the tunable polymer networks strongly depend on a variety of chemical factors such as catalyst and ligand identity, rate of initiation, and initiator structure.36 Thus, we next explored the tunable radical production. Second, we varied the concentration of the initiator over the range of 250 to 1000 μM, which should in turn also tailor radical concentration. As expected, increasing gel stiffness was a function of increasing initiator concentration (Figure 2c). An additional advantage of ATRP over hydroxyl radicals is the potential for using structurally well-defined radical initiators. Consistent with this expectation, we found that a PEG-based initiator, poly(ethylene glycol) bis(2-bromoisobutyrate) ($M_{\text{avg}} = 700$ g/mol), also successfully cross-linked EET-controlled gels at a variety of concentrations (Figure S8). MeHA macromers of varying methacrylate percentages also cross-linked as expected, exhibiting cross-link density saturation around 65% functionalization (Figure S9). Overall, EET-controlled hydrogels exhibited a modulus range of about 1–6 kPa for the conditions tested, which is typical of chain-growth cross-linked MeHA hydrogels and within range for a variety of applications, such as biofilm and tissue mimetics.33,35,38 These results also indicate that traditional approaches to tuning hydrogel mechanics through chemical mechanisms are still applicable when using EET-controlled cross-linking.

Next, we investigated the role of S. oneidensis cell density, and thus aggregate EET flux, on hydrogel modulus. Below a certain critical inoculum (ca. initial OD$_{600} = 0.1$), the rate of bacterial oxygen consumption was not fast enough to overcome oxygen diffusion and radical quenching. As expected, hydrogels formed with sufficient cell density, and stiffness strongly correlated with OD$_{600}$ (Figure 2d). Based on these results, we predicted that cross-linking rate would also be coupled to EET and inoculating cell density. To confirm this, we performed rheological measurements in situ, which provided real-time measurement of mechanical properties during cross-linking, at 1 Hz and 0.1% strain. Gels formed at higher initial cell concentrations were not only stronger, but formed more quickly (Figure 2e). Consistent with end-point experiments, in situ rheology measurements also confirmed that a critical concentration of cells was necessary for oxygen depletion. Together, these results demonstrate that cells play a direct role in cross-linking, and overall stiffness and cross-linking rate can be controlled by cell inoculum. They also suggest that genetic and metabolic manipulations to tune EET flux could be used to influence gel mechanics.

S. oneidensis Remains Viable and Metabolically Active in the Network. For a living material to maintain adaptability, it is critical that the actuating components (i.e., cells) remain viable and encased in the network. The various components of our system, including the Cu catalyst, initiator, and presence of radicals could affect cell viability. Thus, we assessed cell viability and activity after cross-linking. EET-cross-linked gels formed in standard conditions were swollen overnight in 1X PBS after modulus measurements and stained using live/dead fluorescent dyes. Even after mechanical stresses induced by swelling and rheometer measurements, cells maintained approximately 100% viability 5 days after cross-linking (Figure S10). In addition, cells exposed to cross-linking conditions but released from the gel surface during swelling could successfully inoculate new cultures in fresh growth media, indicating viability; these cultures were also able to cross-link new hydrogels with identical properties (Figure S11).

Since new cultures could be inoculated using cells released during swelling, we next quantified escape or leakage of bacteria from the gels after cross-linking. At the functional group density and cross-link molecular weight of our material, the mesh size of a fully converted gel should be on the order of...
10−50 nm, allowing for catalyst and electron shuttle diffusion. However, because this is considerably smaller than average bacterial dimensions, there should be minimal cell movement. To test this prediction, cross-linked gels were prepared at standard conditions, swollen in 1 mL of 1× PBS, and the optical density of the surrounding media was measured. An initial, low optical density of cells was detected immediately upon swelling. We hypothesized that this was due to an instantaneous egress of cells on the periphery of the gels and not contained in the network. After washing gels three times with 1 mL PBS to remove this outer layer of cells, no increase in optical density was detected. Furthermore, colony counting confirmed that escaped cells after 24 h of swelling accounted for <0.005% of the inoculating density (Figure S12), suggesting embedded cells do not escape the network in significant numbers.

Real-time responses within the network and design of new functions require an understanding of spatiotemporal cell behavior within the gels during synthesis. Therefore, we next visualized the relationship between genotype, cross-link density, and cell movement during gelation. We constructed an inducible sfGFP expression plasmid under the control of the LacI repressor protein and its cognate promoter, P\text{lac}. Cells were transformed with this vector such that sensing of isopropyl β-D-1-thiogalactopyranoside (IPTG) would induce a fluorescent response indicative of metabolic activity. We developed two reporter strains by transforming both \textit{S. oneidensis} MR-1 and ΔmtrCΔomcAΔmtrF (an EET-deficient knockout) with this construct. Strains were grown overnight in 1000 μM IPTG, washed, and inoculated into standard gelation mixtures. The solution was then pipetted onto a glass slide and sealed under a coverslip, such that cross-linking occurred in the sealed layer. Bacterial movement was monitored by time-lapse imaging using GFP fluorescence. Cells were uniformly dispersed within the network throughout gelation. For both \textit{S. oneidensis} MR-1 and ΔmtrCΔomcAΔmtrF, a significant degree of bacterial motion was visible upon inoculation, both by convective flow of the reaction mixture and by flagella-based swimming.41 Minutes after inoculation, cell movement and swimming were arrested in the \textit{S. oneidensis} MR-1 sample (0 h, Movie S1; 2 h, Movie S2). Contrastingly, movement both from flow and swimming were still perceptible after 2 h in the ΔmtrCΔomcAΔmtrF sample, indicating that minimal cross-linking occurred (0 h, Movie S3; 2 h, Movie S4). Quantification of cell movement revealed that average cell displacement over 5 s was significantly greater for the knockout strain at both 0 and 2 h (Figure 3a, Figure S13).42 A decrease in cell movement for the ΔmtrCΔomcAΔmtrF mutant was also observed at 2 h, which is likely a result of background cross-linking (described below). Movement was not significantly different between the two strains in nonfunctionalized hyaluronic acid solution, suggesting the observed motility differences were due to cross-linking, even at early times immediately following inoculation (Figure S14, Movies S5–8). These results confirm that cells become trapped in the polymer network as it forms and suggest that the bacterial genotype encodes control over bulk and microscopic properties such as cross-link density and mesh size, affecting flow, diffusion, and cell movement within the material.

Although the cells remained viable for days in the cross-linked gels, we wished to assess their continued sensing and metabolic capabilities over long periods after cross-linking.

Figure 3. Cell motility, synthesis rate, and cross-link density within living hydrogels are governed by \textit{S. oneidensis} genetics and EET machinery. (a) Average cell displacement within the gels measured by microscopy over 5 s time-lapses at both 0 and 2 h into cross-linking; student t test \(p < 0.0001\) between strains at \(t = 0\) and 2 h. (b) \textit{In situ} rheology measurements of hydrogels cross-linked by \textit{S. oneidensis} strains with various EET genes knocked out. \textit{E. coli} was included as an EET-deficient control; student t test \(p < 0.0001\) for MR-1 compared to other strains at \(t = 1\) and 2 h. Data are shown as mean ± SEM, (a) \(n = 33\) tracked cells or (c) \(n = 3\) biological replicates.

Gels synthesized at standard conditions with sfGFP plasmid-carrying \textit{S. oneidensis} MR-1 were swollen in 1× PBS for varying lengths of time after cross-linking, then induced for 24 h with 1000 μM IPTG. Significant fluorescence was detected by microscopy in induced samples up to 1 week after cross-linking.
but was not detectable in uninduced samples (Figure S15). Together, these results indicate that the bacteria remain viable, trapped, and maintain transcriptional and translational capabilities for extended periods after cross-linking.

**Bacterial Genetics Govern Cross-link Density.** Understanding the genetic link between EET and cross-linking is critical for biologically controlling hydrogel structure and function. Toward this goal, we employed various EET knockout strains in cross-linking reactions: \( \Delta \text{mtrC} \), \( \Delta \text{omcA} \Delta \text{mtrF} \), and \( \Delta \text{Mtr} \). The knockout \( \Delta \text{Mtr} \) refers to a strain with a large number of EET genes removed from the genome, and should provide minimal electron flux to the catalyst (Table S1).

E. coli MG1655 was also included as an EET-deficient control. We measured in situ cross-linking kinetics using these strains and compared cross-linking rates and density. Both cross-linking rate and hydrogel storage modulus strongly corresponded with bacterial genotype, where decreasing number of EET genes led to decreased cross-linking rates and weaker moduli (Figure 3b). Although MtrC is the primary terminal reductase for many metal substrates, our results show that MtrF exhibits compensatory reduction of Cu in the \( \Delta \text{mtrC} \Delta \text{omcA} \Delta \text{mtrF} \) knockout compared to the \( \Delta \text{mtrC} \Delta \text{omcA} \Delta \text{mtrF} + \text{mtrC} \) knockout. The strong similarity between gels formed by \( \Delta \text{mtrC} \Delta \text{omcA} \Delta \text{mtrF} \) and \( \Delta \text{Mtr} \) knockouts further demonstrates that outer membrane cytochromes are primarily responsible for electron transfer to the Cu catalyst and subsequent cross-linking activity. The minimal, delayed cross-linking activity of \( \text{E. coli} \) suggests that background radical generation or nonspecific Cu reduction can produce weak gels at extended times. In separate experiments, we corroborated these in situ results using end-point, swollen gel measurements after 1 and 2 h of cross-linking (Figures 3c and S16). 

\[ y = 1.1x - 0.15 \]

Figure 4. Modeling gene expression allows quantitative prediction of living hydrogel properties. (a) Genetic circuits utilized in this study placing either \( \text{sfgfp} \) or \( \text{mtrC} \) under inducible control of IPTG via the LacI repressor and \( \text{Ptac} \) promoter. (b) In situ rheology of hydrogels cross-linked by \( \text{S. oneidensis} \) with steady-state induced \( \text{mtrC} \) levels at various IPTG concentrations. \( \Delta \text{mtrC} \Delta \text{omcA} \Delta \text{mtrF} + \text{empty} \) and \( \Delta \text{mtrC} \Delta \text{omcA} \Delta \text{mtrF} + \text{mtrC} \) with 0 \( \mu \text{M} \) IPTG were also tested but did not form gels on the time scale shown. (c) Hill function analysis of \( \text{sGFP} \) fluorescence, denoted as relative expression units by normalization to fluorescence at maximum induction, as a function of IPTG concentration. (d) Hill function analysis of hydrogel storage modulus after 2 h of cross-linking as a function of IPTG concentration. The right y-axis is storage modulus normalized to average modulus at maximum induction. (e) Normalized hydrogel stiffness plotted as a function of relative expression units for corresponding IPTG concentrations. The fit was determined by performing linear regression on the paired data (\( R^2 = 0.80 \)). (c−e) Data are shown as mean ± SEM, \( n = 3 \) biological replicates.
material properties can be controlled through more sophisticated regulation of EET.

Transcriptional Regulation of Extracellular Electron Transfer Yields Tunable Cross-linking Activity. For ELMs to emulate the adaptability of biological materials, the actuating components should continually sense and respond to their environment. Environmental stimuli should then induce a transcriptional response and impart control over material properties. Toward this goal, we constructed an inducible mtrC expression plasmid using the same genetic circuit outlined before, but replacing sfgfp with mtrC (Figures 4a, S17, and S18). We transformed the ΔmtrCΔomcAΔmtrF strain with this plasmid, such that IPTG would sequentially activate mtrC expression, electron transfer, and cross-linking activity. Upstream of the mtrC gene, a computationally predicted weak synthetic ribosome binding site was employed to optimize control over EET and minimize leaky expression. SDS-PAGE and heme staining of total protein from induced and uninduced cell lysates validated inducible MtrC protein production after overnight growth in IPTG-containing medium. High molecular weight bands corresponding to the size of the MtrCAB complex were observed in induced ΔmtrCΔomcAΔmtrF + mtrC samples and a wild-type control, but not in uninduced and empty vector ΔmtrCΔomcAΔmtrF samples (Figure S19). Functional steady-state expression of mtrC in response to IPTG was further validated by measuring Fe³⁺ reduction with the ferrozine assay. After 2 h of reduction, Fe²⁺ concentration increased with the presence of inducing molecule, indicating functional MtrC activity and no leaky EET response over an uninduced control (Figure S20). Next, we verified tailored cross-linking activity in response to varying transcriptional activation. In situ gelation kinetics were assessed after overnight growth in media containing a range of inducing molecule concentrations. Cross-linking activity was a strong function of IPTG concentration, spanning orders of magnitude in storage modulus (Figure 4b). Cross-linking kinetics also corresponded to inducer presence, indicating that both synthesis rate and final material modulus can be customized through differential steady-state gene expression. Both an induced empty vector control and a complemented strain with no IPTG did not form measurable gels in 2 h. Thus, transcriptional regulation over EET gene expression in response to an environmental signal imparts programmable control over hydrogel stiffness.

Modeling Gene Expression Enables Predictable Material Properties. Due to successful transcriptional regulation of mtrC, we hypothesized that a material property such as storage modulus could be predicted from inducible gene expression models. Since the sfgfp and mtrC circuits have identical transcriptional regulation, we tested whether both fit to activating Hill function models. First, we measured the response function of the sfgfp circuit in S. oneidensis MR-1 by inducing overnight cultures in a variety of IPTG concentrations. Steady-state fluorescence was quantified using a plate reader and normalized to optical density (Figure 4c). As expected, relative expression (i.e., normalized fluorescence) was a strong function of IPTG concentration and fit well to a Hill function with a hillslope of \( n = 1.57 \) and a half-maximal effective concentration of \( EC_{50} = 98.6 \ \mu M \) (Table S5). These results indicate that our circuit generates a predictable transcriptional response. Next, end-point gel measurements were used to examine storage modulus as a function of steady-state cytochrome (MtrC) expression. Gels were cross-linked for 2 h since our in situ results indicated this time would provide sufficient differentiation between induced cultures at varying IPTG concentrations. Specifically, \( \Delta mtrC\Delta omcA\Delta mtrF \) complemented with LacI-regulated \( mtrC \) was grown overnight in a variety of IPTG concentrations and allowed to react for 2 h at standard gelation conditions. We found that, similar to sfgfp fluorescence, hydrogel storage moduli were also under strong transcriptional control and could be modeled using a Hill function with \( n = 1.40 \) and \( EC_{50} = 96.4 \ \mu M \) (Figure 4d). As the sfgfp signal is effectively a measure of the transcriptional rate at different IPTG concentrations, the similarity between fitted constants for sfgfp expression and hydrogel stiffness suggests that transcriptionally regulated MtrC levels can predictably control hydrogel properties via a linear model (Note S1). To further visualize this relationship, we plotted normalized storage modulus as a function of relative expression units for each corresponding IPTG concentration and indeed observed a linear correlation (Figure 4e). The slope and intercept obtained by linear regression of the paired data are similar to the fit predicted by the individual Hill functions, corroborating the model between steady-state gene expression and hydrogel modulus. Together, these results demonstrate that EET gene expression can be modulated to control ELM properties (e.g., gel stiffness) and that fluorescence-parametrized models for existing and new genetic circuits may be adapted to design, predict, and control more complex macroscopic material outputs.

**DISCUSSION**

We showed that S. oneidensis can genetically control radical cross-linking in a semisynthetic hydrogel via electron transfer to a redox-active polymerization catalyst. Similar to other cross-linking chemistries, storage modulus was dependent on catalyst and initiator concentrations as well as initial cell density. A significant advantage of our system is that it is theoretically amenable to any substrate that can undergo radical cross-linking and support microbial life. We used methacrylate-functionalized hyaluronic acid, but other semisynthetic materials based on functionalized alginate, collagen, and cellulose, as well as completely synthetic substrates, such as PEG, should show similar behavior. In addition to flexibility in macromer structure, our design also allows for a variety of well-defined ATRP initiators to be used as cross-linking agents. For example, we showed two traditional ATRP initiators, HEBIB and bis-brominated PEG, could both form cross-linked hyaluronic acid hydrogels. Similar to other radical cross-linking methodologies, bacteria-controlled cross-linking is also compatible with various biochemical modifications including the installation of integrin recognition motifs (e.g., RGD), orthogonal cross-linking chemistries (e.g., Michael addition), and other common polymer engineering paradigms. The chemical flexibility and general compatibility with a variety of polymer network scaffolds should facilitate the use of our platform in tissue engineering, 3D printing, soft robotics, and drug delivery. While utilizing bacteria for applications requiring mammalian biocompatibility is challenging, potential uses include platforms for studying cell–microbe interactions mediated by the extracellular matrix, such as tissue repair and the gut lumen. In contrast to other biologically driven radical cross-linking methods, most notably hydroxyl radicals generated from glucose oxidase, EET-controlled cross-linking did not negatively impact cell health. Cells remained viable at least 1
week following gelation and transformed cells could express sffgp in response to an external stimulus. We also observed genotypic changes in cell motility and convective flow as a result of EET-dependent cross-linking, implying genetic control over gel microstructure. Overall, our design avoids cell viability concerns associated with other radical cross-linking methodologies and could enable synthetic platforms for studying biofilm formation, trapping, or functionizing cells.

We found that cross-linking activity and overall hydrogel stiffness were governed by EET cytochrome expression. Specifically, S. oneidensis MR-1 with wild-type EET pathways generated stiff gels within an hour while negative controls containing E. coli MG1655, which lacks EET machinery, did not form gels on comparable time scales. At longer time scales (ca. 2–4 h), EET-knockout strains and E. coli showed some cross-linking activity. This background radical generation could be caused by nonspecific copper reduction (e.g., release or secretion of cytosolic reducing agents) or spurious radical activation. To further reduce background cross-linking, decreasing catalyst and/or initiator concentration, lowering cell density, or changing the identity of chemical components could all potentially be tuned. Overall, the strong link between S. oneidensis genetics and cross-linking rate and density lays the foundation for developing more sophisticated EET-based regulation of material properties.

Using in situ and end-point rheology measurements, we showed that hydrogel cross-linking is directly linked to mtrC expression levels, which has a number of implications for adaptable and dynamic materials. Interestingly, placing mtrC under the control of the LacI repressor generated a hydrogel stiffness response function that is characteristic of inducible gene expression. This response function mirrored one generated from sffgp expression in the same construct, indicating robust transcriptional control over both gene expression and material properties. More importantly, our results suggest that previously characterized genetic circuits, including genetic logic gates, designed to express fluorescent reporters could be readily adapted to control mtrC expression and gel stiffness. Although the ultimate stiffness of the hydrogel will depend on the specific cross-linking chemistry, our observation of canonical Hill function responses demonstrates that changes in hydrogel stiffness governed by transcriptional regulation can be partially predicted. Overall, our results suggest that a variety of transcriptional circuits could be extended to control the macroscopic properties of synthetic materials in a predictable and programmable manner. Additionally, robust genetic control over cross-linking should complement other stimuli-responsive hydrogel designs, including integration of biochemical signals, actuators, and complex geometric designs.

The gel stiffness response function was measured after 2 h of gelation since in situ measurements indicated this would be sufficient time to distinguish between differentially induced cells. Because cross-linking is a dynamic process, the hydrogel response function also varies as a function of time, even at steady-state expression levels of MtrC. For example, at early times (<1 h), it is exaggerated since measurable gels do not form at low induction levels (Figure S21). At longer time scales, the response function begins to collapse as background polymerization starts to compete with EET-driven cross-linking. Applications leveraging genetic control over material properties will require both an understanding of how EET influences cross-linking kinetics and the optimization of real-time transcriptional responses. Understanding how cells coordinate transient gene expression to control dynamic outputs such as cell motility, morphogenesis, biofilm structure, or extracellular matrix construction are ongoing challenges in developmental and systems biology. Similarly, we are currently investigating how to match gene expression to the polymerization kinetics in our system. Overall, continued optimization of MtrC (or other EET protein) expression and material chemistry should allow for actuation of material changes over time scales similar to transcription and translation, as well as predictive models that relate gene expression to material function.

Overall, we found thatextracellular electron transfer from S. oneidensis could power a radical polymerization catalyst and form a semisynthetic hydrogel composed of functionalized hyaluronic acid. A variety of chemical and biological factors controlled cross-linking kinetics and the resulting storage moduli of the gels, demonstrating a tunable and adaptable platform. Most importantly, we found that robust transcriptional control over mtrC expression and metabolic electron flux enabled precise and predictable control over hydrogel mechanical properties. While cells are frequently incorporated into polymer networks, our platform allows for a variety of network properties including cross-link density, mesh size, degradation, diffusion, and elastic modulus to be controlled through cellular metabolism and gene expression. In summary, our results provide a powerful foundation for programming adaptive and responsive behavior into the vast functional space of synthetic materials through the conduit of biological electron transfer.

**MATERIALS AND METHODS**

**Chemicals and Reagents.** Sodium hyaluronate (72 kDa, Lifecore Biomedical), methacrylated hyaluronic acid (Sigma-Aldrich, 94%), copper(II) bromide (CuBr₂, Sigma-Aldrich, 99%), tris(2-pyridylmethyl)amine (TPMA, Sigma-Aldrich, 98%), 2-hydroxyethyl 2-bromoisobutyrate (HEBIB, Sigma-Aldrich, 95%), poly(ethylene glycol) bis(2-bromoisobutyrate) (PEGBIB) (Sigma-Aldrich, 95%), MtrC (or other EET protein) expression and metabolic electron flux enabled precise and predictable control over hydrogel mechanical properties.

*ACS Biomater. Sci. Eng.* 2020, 6, 1375–1386
solution was dialyzed using 6–8 kDa dialysis tubing in DI water while stirring for 2 weeks. The mixture was then frozen and lyophilized. Methacrylate functionalization was quantified by $^1$H NMR spectroscopy and determined to be ~65% from integration of the vinyl group relative to the HA backbone (Figure S1). Functionalized MeHA solution was passed through a basic alumina column immediately prior to cross-linking reactions. Methacrylate functionalized group conversion was quantified by adding 10 μL of 50 mg/mL hyaluronidase solution in 1X PBS to 50 μL hydrogels and degrading overnight at 37 °C in an incubator with 250 rpm shaking. Liquefied solutions were diluted in 550 μL of D$_2$O and the methacrylate peak quantified using $^1$H NMR spectroscopy.

**Bacteria Strains and Culture.** Bacterial strains and plasmids are listed in Table S1. Cultures were prepared as follows: bacterial stocks stored in 20% glycerol at −80 °C were streaked onto LB agar plates (for wild-type and knockout strains) or LB agar with 25 μg/mL kanamycin (for plasmid-harboring strains) and grown overnight at 30 °C for *Shewanella* and 37 °C for *E. coli*. Single colonies were isolated and inoculated into *Shewanella* basal medium (SBM) supplemented with 100 mM HEPES, 0.05% trace mineral supplement, 0.05% casamino acids, and 20 mM sodium lactate (2.85 μL of 60% w/w sodium lactate per 1 mL culture) as the electron donor. Aerobic cultures were pregrown in 15 mL culture tubes at 30 °C and 250 rpm shaking. Anaerobic cultures were pregrown using the same procedure outlined above, but in degassed growth medium in a humidified anaerobic chamber and supplemented with 40 mM sodium fumarate (40 μL/mL of a 1:1 stock) as the electron acceptor. Cultures were washed 3× after pregrowth using SBM supplemented with 0.05% casamino acids (degassed for anaerobic cultures). OD$_{600}$ was measured using a NanoDrop 2000C spectrophotometer and normalized to 2.0 for 10-fold dilution into gel mixtures (5 μL of OD$_{600}$ = 2.0 concentrated cell culture into 45 μL of gel mixture) unless otherwise noted.

**MeHA Hydrogel Cross-linking Using *S. oneidensis***. CuBr$_2$, TPMA, and HEBIB stock solutions were prepared according to previously established protocol. For three 50 μL hydrogel discs that were analyzed by rheology, a reaction mixture was prepared as follows: MeHA was dissolved at 3.76 wt % in SBM with 0.05% casamino acids and aliquoted into an autoclaved microfuge tube (119.2 μL). Solutions of 400 μM Cu-TpPMA (3.75 μL), 69 mM HEBIB (1.09 μL), 60% sodium lactate (0.428 μL), and 1 M sodium fumarate (6 μL) were added to the MeHA solution and mixed. Per 50 μL gel mixture, the remaining 1.5 μL was used for antibiotic and inducing molecule addition where necessary, otherwise 1.5 μL of SBM with casamino acids was added. The final concentrations in solution were 3 wt % MeHA, 10 μM Cu TPpMA, 500 μM HEBIB, 20 mM lactate, and 40 mM fumarate. This solution was distributed into three autoclaved microfuge tubes of 45 μL aliquots to which 5 μL of OD$_{600}$ normalized cells were added. The gel solutions were mixed and dispensed onto hydrophobically treated glass slides with a 0.5 mm silicone spacer separating the two glass layers. Slides were sealed with a binder clip and allowed to react at 30 °C for 2 h unless otherwise noted. Hydrogels were removed from the slides using a razor blade and placed into 3 mL baths of 1X PBS overnight to swell to equilibrium. Hydrogels analyzed by in situ rheology were prepared using the same mixture outlined above, but inoculated with cells and immediately placed on the rheometer for analysis.

**Rheological Analysis.** End-point rheological analysis: swollen hydrogels prepared as outlined above were analyzed by oscillatory shear rheology using a TA Instruments Discovery HR-2 Rheometer with an 8 mm parallel plate geometry. Hydrogels were loaded onto a Peltier plate and excised to 8 mm diameter using a biopsy punch. The geometry gap was then lowered until the measured axial force was above 0.02 N (usually between 500–800 μm, depending on the cross-link density and swelling ratio). Storage and loss moduli were measured using frequency sweeps from 0.01 to 100 Hz at a constant strain of 0.1%. Moduli for a single gel were quantified by averaging the linear viscoelastic region of each frequency sweep.

In situ rheological analysis: hydrogels measured by in situ oscillatory shear rheology were prepared using the mixtures and rheometer outlined above. Immediately after inoculating reaction mixtures with cells, 80 μL of mix was loaded onto the Peltier plate, which was maintained at 30 °C. A 20 mm parallel plate geometry was lowered onto the solution while spinning such that the mixture coated the entire geometry surface and filled the gap (~350 μm gap size). The edges of the geometry and gap were then coated with silicone oil to prevent evaporative losses. In situ cross-linking was monitored using 1 Hz oscillation and 0.1% strain over variable lengths of time (1.5–8 h).

**Microscopy and Cell Tracking.** All microscopy was performed using a Nikon Ti2 Eclipse inverted epifluorescence microscope. Cells assessed for viability by microscopy were cross-linked using standard conditions and the resulting gels swollen in 1X PBS at room temperature for varying lengths of time. The gels were then incubated in the dark in the BacLight Live/Dead stain mix (1 μL/mL Syto9, 2.5 μL/mL propidium iodide in 0.85% NaCl solution) for 30 min. Stained gels were then washed by pipetting 3X in 1 mL PBS to remove unbound dye. Gels were loaded onto glass microscope slides, and a no. 1 coverslip was placed on top. The gel thickness prevented using nail polish to seal the sides, but evaporative losses were not noticeable over the course of the experiment (~30 min). Fluorescence for each stain (green for Syto9, red for propidium iodide) was measured using GFP and Texas Red excitation/emission filter cubes on a Nikon Ti2 Eclipse, as outlined previously. To assess metabolic activity, gels were cross-linked with sfGFP-harbouring strains and allowed to swell in 1X PBS for varying lengths of time. sfGFP fluorescence was assessed before induction to ensure there was no detectable background fluorescence. Gels were then incubated in 0 μM or 1000 μM IPTG in PBS for 24 h and monitored by fluorescence using the GFP channel.

Cells tracked by microscopy during cross-linking were prepared with reaction mixtures as outlined above. Upon cell inoculation, the cross-linking mixture was loaded onto glass slides, covered, and sealed with nail polish. The slides were loaded onto the microscope and cell movement monitored using the Time-lapse function in NIS-Elements. Images were taken every 1 or 5 s with 100 or 300 ms exposure time using the GFP channel. Time-lapse images were edited and quantified using TrackMate in Fiji 1.0. Images were first background subtracted and equally brightened by thresholding. The top 50 highest quality cells were selected, as determined by the TrackMate user interface, and tracked over 5 or 10 s. The highest quality tracks, as determined by the software, were used to quantify average total displacement over the time-lapse. The number of tracks used to calculate the average was the maximum number of tracks from the image with the fewest tracks at each time point (e.g., if *S. oneidensis* MR-1 had 100 high-quality tracks and *S. oneidensis* JGS96 had 50 high-quality tracks at the 0 h time point, the 50 highest quality tracks from each image were used).

**Plasmid Construction.** DNA sequences and plasmid maps for each genetic part and plasmid used in this study are given in the Supporting Information (Figure S17–S18, Table S4). All plasmids were assembled via Golden Gate cloning using enzymes and buffers from New England Biolabs. In addition to T4 Ligase, Golden Gate reactions contained either SapI or mtrC plasmid assembly or BsaI for sfGFP and empty plasmid assembly. The pCD backbone was assembled by PCR amplifying (Phusion, New England Biolabs) regions of pSR8.6 (B0015 and T0 terminators, ColE1 origin of replication), pTREI-loV (kanamycin resistance), and pAL-flp (RP4 origin of transfer). To construct the gene expression unit (instructing terminators, RiboJ ribozyme, and lacI regulation unit), a gBlock was synthesized (Integrated DNA Technologies) and used in Golden Gate cloning. sfGFP and mtrC were PCR amplified from pSR8.6 and purified *Shewanella oneidensis* MR-1 genomic DNA, respectively, with ribosome binding sites and Golden Gate restriction enzyme sites added via oligonucleotide primers. Generally, 10 μL Golden Gate reactions that contained 10 fmol of pCD plasmid backbone and 40 fmol of each gBlock and/or PCR insert (as necessary) were used. In a thermocycler, Golden Gate reactions were cycled 25 times: 90 s at 97 °C followed by 3 min at 16 °C. After the 25 cycles, reactions were incubated at 37 °C for 5 min, 80 °C for 10 min, and then held at 4 °C.

Golden Gate reactions were used to directly transform freshly prepared electrocompetent *S. oneidensis* strains. To prepare
electrocompetent *S. oneidensis*, 5 mL of overnight *S. oneidensis* growth in LB medium at 30 °C was washed 3 times with sterile 10% glycerol at room temperature and concentrated to ~300 μL. A 2 μL portion of Golden Gate reaction was mixed with 30 μL of concentrated electrocompetent *S. oneidensis*, transferred to a 1 mm electroporation cuvette, and electroporated at 1250 V. To recover electroporated cells, 250 μL of LB was immediately added post electroporation and cells were incubated/shaken at 30 °C and 250 rpm. After 2 h of recovery, 100 μL of cell suspension was plated onto LB agar plates containing 25 μg mL⁻¹ kanamycin sulfate and incubated overnight at 30 °C to obtain single colonies (generally 5–100 colonies observed for 1–3 part assemblies). Single colonies were used to inoculate LB liquid medium containing 25 μg mL⁻¹ kanamycin sulfate and incubated/shaken overnight at 30 °C and 250 rpm. These cultures were used to generate 22.5% glycerol stocks, which were stored at −80 °C, and harvest assembled plasmid for Sanger sequencing (DNA Sequencing Facilities, University of Texas at Austin).

**Verification of MtrC Inducible Expression and Functional Activity.** Heme staining was performed by adapting previously described methods.⁶¹ A 5 mL portion of uninduced (0 μM IPTG) and induced (1000 μM IPTG) *S. oneidensis* strains were anaerobically cultured overnight in SBM containing 20 mM lactate and 40 mM fumarate. The total culture was washed once in 1× PBS, concentrated to 500 μL, and lysed by sonication. The cell lysate was centrifuged for 10 min at 10 000 rcf, and the supernatant was transferred to a separate tube. The pellet was resuspended in 100 μL 1× PBS, and the total protein concentration of both lysate fractions was determined by Bradford assay. A 10 μg portion of protein from the supernatant and pellet were loaded into each well of a 12% Bis-Tris SDS-PAGE gel and run for ca. 120 min at 110 V. The gel was stained in a 3:7 mixture of 0.3 M TMBZ in methanol:0.25 M sodium acetate (pH 5.0) for 2 h in the dark. Heme-containing protein bands were visualized upon addition of 30 mM hydrogen peroxide for 30 min.

**Ferrozine assay** was performed to determine functional activity of MtrC-expressing strains. Strains were anaerobically grown overnight in IPTG-free SBM containing 20 mM lactate and 40 mM fumarate. Within anaerobic chamber, cells were diluted 100-fold into 96-well plate wells filled SBM containing 20 mM lactate, 5 mM Fe(III)-citrate, and 1 mg mL⁻¹ ferrozine (final total volume of 250 μL per well). Each well contained either 0 or 750 μM IPTG and Fe(II) standards were also added to the plate. Immediately after addition of cells and Fe(III), the plate was sealed with optically transparent sealing film and a plate cover lined with silicone grease. The plate was then removed from the anaerobic chamber and statically incubated at 30 °C. Absorbance at 562 nm was periodically measured using a BM LabTec CLARIOstar plate reader.

**Quantification and Modeling of Inducible Constructs.** The ΔmtrCΔomcaAΔmtrF + sfgf and + mtrC strains were anaerobically pregrown overnight using the same conditions outlined above, with the addition of 25 μg mL⁻¹ kanamycin and varying IPTG concentrations. Anaerobic conditions were used for pregrowth to optimize expression of Mtr pathway components besides mtrC. Cells were anaerobically washed 5× in degassed SBM with 0.05% casamino acids, normalized to OD₆₀₀ = 2.0, and diluted 10-fold into reaction mixtures (also containing kanamycin and IPTG) prepared in ambient conditions. Prior to measuring sfGFp fluorescence, protein translation was arrested by supplementing a 100 μL aliquot of cell suspension with kanamycin sulfate to a final concentration of 2 mg mL⁻¹. Subsequently, this suspension was shaken aerobically for 1 h at 30 °C to allow for sfGFp maturation. sfGFp fluorescence (488/530 nm) and cell suspension absorbance (600 nm) was measured using a BM LabTec CLARIOstar plate reader to yield fluorescence-absorbance for each sample. For each sample, the background fluorescence-absorbance was subtracted. The background subtracted values were then normalized to the average fluorescence-absorbance value at maximum induction (1500 μM IPTG) to give relative expression units. Cross-linking strains complemented with mtrC were allowed to form gels for 2 h, and the gels allowed to swell overnight in 1× PBS at room temperature. Gels were then analyzed by oscillatory shear rheology as outlined above. A nonlinear fitting algorithm in GraphPad Prism 8.0 was used to fit inducible gene expression and hydrogel storage modulus to the following activating Hill function: \( y = \min + (\max - \min) \left( \frac{1}{1 + \left(\frac{x}{IC_{50}}\right)^n} \right) \). Normalized hydrogel storage modulus and relative expression units were plotted for values at corresponding IPTG concentrations and modeled using a linear regression. Further details on modeling can be found in Note S1. Fitting parameters and “goodness of fit” can be found in Table S5.

**Statistical Analysis.** Unless otherwise noted, data are reported as mean ± SEM of n = 3 biological replicates. Significance was calculated in GraphPad Prism 8.0 using either a two-tailed unpaired student t test or a one-way ANOVA (α = 0.05).

### ASSOCIATED CONTENT

> **Supporting Information**
>
> The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsbiomaterials.9b01773.

Strains used in this study, supplementary quantification of hydrogel properties, viability and microscopy studies, plasmid and promoter maps, and notes on gene expression modeling (PDF)

Movies depicting cell movement and bulk fluid motion under several conditions as mentioned in the text (AVIs)

### AUTHOR INFORMATION

**Corresponding Authors**

**Adrienne M. Rosales** — McKetta Department of Chemical Engineering and Center for Dynamics and Control of Materials, University of Texas at Austin, Austin, Texas 78712, United States; orcid.org/0000-0003-0207-7661; Email: arosales@che.utexas.edu

**Benjamin K. Keitz** — McKetta Department of Chemical Engineering and Center for Dynamics and Control of Materials, University of Texas at Austin, Austin, Texas 78712, United States; orcid.org/0000-0003-3314-0053; Email: keitz@utexas.edu

**Authors**

**Austin J. Graham** — McKetta Department of Chemical Engineering and Center for Dynamics and Control of Materials, University of Texas at Austin, Austin, Texas 78712, United States; orcid.org/0000-0001-8924-181X

**Christopher M. Dundas** — McKetta Department of Chemical Engineering and Center for Dynamics and Control of Materials, University of Texas at Austin, Austin, Texas 78712, United States; orcid.org/0000-0001-9183-8236

**Alexander Hillsley** — McKetta Department of Chemical Engineering, University of Texas at Austin, Austin, Texas 78712, United States

**Dain S. Kasprak** — Department of Biomedical Engineering, University of Texas at Austin, Austin, Texas 78712, United States

Complete contact information is available at: https://pubs.acs.org/10.1021/acsbiomaterials.9b01773

**Author Contributions**

Notes
The authors declare no competing financial interest. Experimental data supporting the findings in this study are publicly available through the Texas Data Repository (https://dx.doi.org/10.18738/T8/YSPW8X).

ACKNOWLEDGMENTS
S. oneidensis knockouts were a generous gift from Prof. Jeffrey Gralnick (U. Minnesota). A.J.G. was supported through a National Science Foundation Graduate Research Fellowship (Program Award No. DGE-1610403). This research was supported by the Welch Foundation (Grant F-1929) and by the National Science Foundation through the Center for Dynamics and Control of Materials: an NSF Materials Research Science and Engineering Center under DMR-1720595. Additional research support was provided by the National Institute of General Medical Sciences of the National Institutes of Health under Award Number R35GM133640 (B.K.K.). The authors acknowledge use of shared research facilities supported in part by the Texas Materials Institute, the Texas Data Repository (https://dx.doi.org/10.18738/T8/YSPW8X).

REFERENCES
(28) Lülsdorf, N.; Pitzler, C.; Biggel, M.; Martinez, R.; Vojicic, L.; Schwaneberg, U. A flow cytometer-based whole cell screening toolbox...


