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CymA and Exogenous Flavins Improve Extracellular Electron Transfer and Couple It to Cell Growth in Mtr-Expressing Escherichia coli

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Authors

Jensen, Heather M TerAvest, Michaela A Kokish, Mark G <u>et al.</u>

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5	Heather M. Jensen <sup>1,3,#a</sup> , Michaela A. TerAvest <sup>5,#b</sup> , Mark G. Kokish <sup>2,3,#c</sup> , Caroline M. Ajo-
6	Franklin <sup>1,2,4</sup> *
7	
8	<sup>1</sup> Physical Biosciences Division, Lawrence Berkeley National Laboratory, Berkeley, California,
9	United States of America.
10	<sup>2</sup> Materials Sciences Division, Lawrence Berkeley National Laboratory, Berkeley, California,
11	United States of America.
12	<sup>3</sup> Department of Chemistry, University of California, Berkeley, California, United States of
13	America.
14	<sup>4</sup> Synthetic Biology Institute, Berkeley, California, United States of America.
15	<sup>5</sup> California Institute for Quantitative Biosciences, University of California, Berkeley, California,
16	United States of America
17	
18	<sup>#a</sup> Current Institution: Joint BioEnergy Institute, Emeryville, California, United States of
19	America.
20	<sup>#b</sup> Current Institution: Michigan State University, East Lansing, Michigan, United States of
21	America.
22	<sup>#c</sup> Current Institution: Northwestern University, Evanston, Illinois, United States of America.
23	*Corresponding author:

24 E-mail: cajo-franklin@lbl.gov (CMAF)

25

#### 26 Abstract

27 Introducing extracellular electron transfer pathways into heterologous organisms offers the 28 opportunity to explore fundamental biogeochemical processes and to biologically alter redox 29 states of exogenous metals for various applications. While expression of the MtrCAB electron 30 nanoconduit from Shewanella oneidensis MR-1 permits extracellular electron transfer in 31 *Escherichia coli*, the low electron flux and absence of growth in these cells limits their 32 practicality for such applications. Here we investigate how the rate of electron transfer to 33 extracellular Fe(III) and cell survival in engineered E. coli are affected by mimicking different 34 features of the S. oneidensis pathway: the number of electron nanoconduits, the link between the 35 quinol pool and MtrA, and the presence of flavin-dependent electron transfer. While increasing 36 the number of pathways does not significantly improve the extracellular electron transfer rate or 37 cell survival, using the native inner membrane component, CymA, significantly improves the 38 reduction rate of extracellular acceptors and increases cell viability. Strikingly, introducing both 39 CymA and riboflavin to Mtr-expressing *E. coli* also allowed these cells to couple metal reduction 40 to growth, which is the first time an increase in biomass of an engineered E. coli has been 41 observed under Fe<sub>2</sub>O<sub>3</sub> (s) reducing conditions. Overall, this work provides engineered E. coli 42 strains for modulating extracellular metal reduction and elucidates critical factors for engineering 43 extracellular electron transfer in heterologous organisms.

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- 46 Keywords: synthetic biology, dissimilatory metal-reducing bacteria, bioelectrochemical systems,
- 47 energy conservation, multi-heme cytochrome c

#### 48 Introduction

49 Some metal-reducing bacteria, such as those from the *Geobacter* and *Shewanella* genera, 50 have extracellular electron transfer pathways that can route electrons across the cell membrane to 51 alter the redox state of exogenous metals. This extracellular metal reduction by microorganisms 52 plays a key role in microbial-driven mineral transformations (1) and can be used to drive nanoparticle synthesis under mild conditions (2, 3) or alter the phase of metals for 53 54 bioremediation (4) or biomining (3). The ability to modulate extracellular electron transfer then 55 offers the opportunity to dissect and control these processes. For example, systematically varying 56 the rate of extracellular electron transfer should allow synthetic control over the number and size 57 of nanoparticles and should elucidate the role of redox kinetics on mineral transformation. 58 However, the sparse availability of genetic tools in metal-reducing bacteria and the multiple 59 overlapping pathways for extracellular electron transfer make it challenging to modulate electron 60 transfer and/or introduce other functions of interest. In response to this challenge, we (5-7) and 61 others (8-10) have taken the complementary approach of engineering portions of extracellular 62 electron transfer pathways into the well-studied industrial microbe *Escherichia coli* (11). While 63 this work has shown that the Mtr pathway can route electrons to extracellular metal oxides (6), 64 the low electron flux and the inability of cells to maintain biomass using solid minerals as the 65 terminal electron acceptor has hampered the use of these strains for applications such as nanoparticle synthesis and bioremediation (7). 66

67 Since the extracellular electron transfer pathways of *Shewanella oneidensis* MR-1 have 68 been extensively studied at the molecular level (*12, 13*), they have provided the basis for these 69 engineering efforts (**Figure 1A**). When *S. oneidensis* is grown under metal-respiring conditions, 70 reducing equivalents from oxidation of electron donors are directed to the menaquinol pool and 71 then to the inner membrane tetraheme cytochrome c CymA (14). When extracellular metals or 72 electrodes are present, electrons from CymA are passed to the periplasmic decaheme cytochrome 73 c MtrA. The re-oxidation of CymA completes the Q-cycle, increasing the proton motive force 74 (15) and allowing the cell to conserve energy. Notably, it is debated whether CymA directly 75 reduces MtrA (9, 16) or whether FccA, STC (also known as CctA), or another redox protein, 76 serve as an intermediate to pass electrons from CymA to MtrA (17-19). MtrA reduces MtrC, a 77 decaheme cytochrome c located on the extracellular face of the outer membrane, as part of the 78 MtrCAB complex. This complex is proposed to form a porin cytochrome complex (12) that 79 spans the outer membrane and allows MtrA to directly contact MtrC (20). There are 80 approximately 70,000 MtrC proteins per S. oneidensis MR-1 cell (21), and these proteins can 81 either directly reduce a mineral or electrode surface or use a flavin-dependent process to 82 indirectly reduce the solid electron acceptor. While the mechanism underpinning flavin-mediated 83 electron transfer is still being elucidated (22-24), it is clear that ~80% of the electron transfer 84 occurs through a flavin-dependent process and  $\sim 20\%$  of the electron transfer occurs directly (25-85 27). Electron transfer through MtrCAB does not affect the proton motive force; however, it 86 indirectly contributes to energy conservation by re-oxidizing CymA.

*E. coli* strains expressing combinations of MtrA and CymA can reduce soluble Fe(III), but not extracellular Fe(III) oxides (8, 9). In contrast, expression of the MtrCAB electron nanoconduit in *E. coli* confers the ability to reduce Fe<sub>2</sub>O<sub>3</sub> (s) (6). However, this strain reduces Fe(III) several orders of magnitude more slowly than *S. oneidensis* MR-1. Likely as a result of this slow reduction rate, this *E. coli* strain is unable to conserve energy and its biomass sharply decreases, which poses a particular challenge for further applications. Introduction of extracellular electron transfer pathways into another industrial microbe, *Pseudomonas putida*,

94 has not translated into cell growth (28-30), indicating this may be a common challenge. Our 95 previous work identified re-reduction of MtrA as a rate-limiting step in Fe(III) reduction (6). 96 This strain also lacks CymA, extracellular flavins, and has fewer Mtr electron conduits (30,000 97 (5) vs. 70,000) (Figure 1A). Here we systematically test how co-expression of CymA, changing 98 the number of electron conduits per cell, and introduction of flavins affect the flux of electrons to 99 extracellular iron and E. coli survival. We found that co-expression of CymA with the Mtr 100 nanoconduit and introduction of exogenous flavins significantly increased the electron transfer 101 rate over the original strains. We also found that E. coli strains expressing CymA coupled these 102 improvements in extracellular reduction to enhanced cellular viability. Importantly, the 103 combination of CymA and riboflavin allowed Mtr E. coli cells to couple Fe(III) reduction to 104 growth under  $Fe_2O_3(s)$ -reducing conditions.

105

#### 106 **Results and Discussion**

## 107 The *cymA*, *cymAmtrA*, *and cymAmtr* strains express the full-length and redox active 108 cytochromes c.

109 To systematically explore the effect of CymA on soluble iron and solid iron oxide 110 reduction, we created three separate plasmids for expression of cymA, cymAmtrA, and 111 cymAmtrCAB (Figure S1); in these plasmids, the RBS sequences were designed to be 112 approximately the same as the corresponding native RBS strength in S. oneidensis (31). The 113 cymA, cymAmtrA, and cymAmtrCAB plasmids were co-transformed with the cytochrome c 114 maturation (ccm) plasmid (32) into E. coli to create the cymA, cymAmtrA, and cymAmtr strains, 115 respectively (Table 1). The *mtrA* strain, co-expressing *ccm* and *mtrA*, and the *mtr* strain, which 116 expresses *ccm* and *mtrCAB*, (**Table 1**), were used as previously described (6).

117 We next characterized the expression and localization of each cvt c to validate each 118 strain. Proteins from whole cell extracts were resolved by SDS-PAGE, and heme c containing 119 proteins were identified by enhanced chemiluminescence (ECL) analysis. These whole cell 120 extracts show that the MtrA and MtrC proteins in all the strains containing *mtrA* and *mtrC*, i.e. 121 the cymAmtrA, mtrA, mtr, and cymAmtr strains, are full-length (35 kD and 76 kD, respectively; 122 Figure 1B). The cymA strain also clearly shows a band corresponding to full length CymA (18 123 kD, Figure 1B). While difficult to discern by eye, densitometry analysis of the ECL 124 demonstrates the CymA band is also present in the two other strains containing cymA, the 125 cymAmtrA and cymAmtr strains (Figure 1C). ECL analysis of fractionated cells shows CymA, 126 MtrA, and MtrC in *E. coli* are localized (Figure S1) as they are in *S. oneidensis* (33-35). Lastly, 127 diffuse reflectance spectroscopy of all these strains under oxidizing and reducing conditions 128 showed characteristic changes in the  $\alpha$ - and  $\beta$ - bands, indicating that the cyt c proteins are redox 129 active. Thus, each strain expresses its respective redox active, full length and correctly localized 130 cyt c.

131 To characterize the concentration range of cyt c that could be expressed, we increased the 132 promoter activity (as defined in (5)) in *mtr* and *cymAmtr* strains via IPTG induction (SI Table 1) 133 and measured the total heme c content in these cells using diffuse reflectance as described 134 previously (5). In brief, diffuse reflectance measures absorbance of visible light with limited 135 scattering, thus allowing quantitative measurement of the  $\alpha$ -bands in whole cell suspensions 136 (Figure S2). From these spectra, we can determine the concentration of heme c, and thus infer 137 relative abundance of electron conduits in whole cell suspensions. From the 0 to 0.37 promoter 138 activity, the cyt c concentration in the mtr and cymAmtr strains both increase ~5-fold (Figure 139 **1D**). The maximum heme c measured in *mtr* and *cymAmtr* strains are 140 and 106  $\mu$ M heme c

140  $OD_{600nm}^{-1}$ , respectively (**Figure 1D**), which is about half the total heme *c* content of *S. oneidensis* 141 when grown under the same conditions (228  $\mu$ M heme *c*  $OD_{600nm}^{-1}$ ). ECL analysis shows that the 142 relative abundance of each cyt *c* within a strain is very similar across different induction levels 143 (**Figure S3**), indicating that induction boosts the abundance of each cyt *c* rather than a single cyt 144 *c*. Thus, tuning the promoter strength via induction allows us to vary the number of electron 145 transfer proteins in the *mtr* and *cymAmtr* strains.

#### 146 Fe(III) citrate reduction and cell survival are improved by co-expression of CymA.

147 Previous work by Schuetz et al. (9) has shown that co-expression of CymA with MtrA 148 from genomically-integrated constructs can improve Fe(III)-NTA reduction in E. coli over 149 expression of CymA alone, strongly suggesting that CymA would improve Fe<sub>2</sub>O<sub>3</sub> reduction in 150 the cymAmtr strain. However, co-expression of CymA alongside other S. oneidensis MR-1 cyts c 151 dramatically reduces its expression level in our strains (Figure 1B-D), and we have previously 152 observed that subtle changes in expression levels can profoundly impact extracellular electron 153 transfer capabilities (5). Thus, we first tested cell suspensions of the cymA, mtrA, and cymAmtrA 154 strains for the ability to reduce soluble Fe(III) citrate in anaerobic conditions with D,L-lactate as 155 a carbon source to determine the effect of limited expression of CymA. Since lysed E. coli cells can reduce  $Au^+$  (36), we also included a homogenized sample of the *mtrA* strain to determine if 156 157 cell lysis contributed to Fe(III) reduction (Figure S4). The lysed mtrA sample reduced at a 158 similar rate as the *ccm* strain (Figure S4), demonstrating cell lysis is not a major contributor to 159 Fe(III) citrate reduction in this system. While the *mtrA* strain reduced Fe(III) slightly faster than 160 the *ccm* control, the *cymA* and *cymAmtrA* strains increased the Fe(III) reduction rate over the 161 *mtrA* strain by a substantial ~3-fold and ~4-fold, respectively (Figure 2A).

In these iron reduction assays, the E. coli strains have only D,L-lactate as a carbon 162 163 source, and the only available terminal electron acceptor for respiration is Fe(III) citrate. 164 Therefore, we hypothesized that aspects which increase Fe(III) reduction would also improve 165 energy conservation and cell survival. Interestingly, while the cell density of the slowest Fe(III)-166 reducing strains, the *ccm* and *mtrA* strains, sharply decreased, the cell density of the fastest 167 Fe(III)-reducing strains, the cymA and cymAmtrA strain, stayed the same or slightly increased 168 (Figure 2B). Indeed, a direct comparison between the rate of Fe(III) citrate reduction and the 169 linear rate of change of the cell density in CymA-expressing strains (Figure S5) reveals that the 170 iron reduction rate and strain fitness are strongly, positively correlated (Pearson correlation 171 coefficient, R = 0.93). This strong positive correlation strongly suggests that increasing the iron 172 reduction rate in these E. coli strains also improves energy conservation under our experimental 173 conditions. The correlation also suggests that a minimum iron reduction rate ( $\sim 0.4$  mM day<sup>-1</sup>, 174 Figure S5) is necessary to maintain or increase cell biomass. Taken together, these observations 175 show that even low levels of CymA co-expressed with MtrA are sufficient to improve Fe(III) 176 citrate reduction and strongly suggest that increasing the Fe(III) reduction rate helps maintain 177 biomass in *E. coli* by increasing energy conservation.

#### 178 $Fe_2O_3(s)$ reduction and cell survival are improved by co-expression of CymA.

179 We next set out to probe the effects of CymA co-expression on solid Fe<sub>2</sub>O<sub>3</sub> reduction and 180 cell viability in the ccm, cymAmtr, and mtr strains. In the solid iron oxide reduction assays, the E. 181 *coli* strains have only D,L-lactate as a carbon source, and the only available terminal electron 182 acceptor for respiration is solid  $Fe_2O_3$ . To survey a wide set of conditions, we varied the number 183 of electron conduits with a range of promoter activity (Figure 1D, Table S1), and we 184 additionally examined the effect of exogenous riboflavin. The promoter activity is indicated in superscript throughout. Due to the relatively slow reduction of solid  $Fe_2O_3$  by these engineered *E. coli* in comparison to *S. oneidensis*, many strains required 21 day incubations before statistically significant measurements of Fe(II) by ferrozine could be attained. In contrast, changes in colony forming units (cfu) were apparent over much shorter time scales than changes in Fe(II) concentration.

190 We first summarize the effects of CymA co-expression with the Mtr electron nanoconduit 191 in E. coli. The ccm only strain, which accounts for basal Fe<sub>2</sub>O<sub>3</sub> (s) reduction, reduced only  $62 \pm$ 19  $\mu$ M Fe(II) over 21 days (Figure 3A). The best reducing *mtr* strain, *mtr*<sup>0.37</sup>. showed a 192 193 statistically significant ~2-fold improvement over the *ccm* strain (Figure 3A). Interestingly, of all the strains and induction conditions tested, the uninduced *cymAmtr* strain, *cymAmtr*<sup>0.001</sup>, 194 195 reduced the greatest amount of  $Fe_2O_3$ , ~3.5 fold more  $Fe_2O_3$  (s) than the ccm strain. The  $cymAmtr^{0.001}$  strain reduced statistically more Fe<sub>2</sub>O<sub>3</sub> (s) than the  $mtr^{0.37}$  strain, demonstrating that 196 197 co-expression of CymA improves the reduction of solid Fe<sub>2</sub>O<sub>3</sub> (Figure 3A) as well as soluble 198 Fe(III) (Figure 2A).

199 CymA also significantly enhanced the survival of *cymAmtr* in the iron reduction assay conditions. The *ccm* strain decreased to 7% of its initial cfu  $mL^{-1}$  over 4 days (Figure 3B), 200 201 consistent with the inability of the native E. coli strain to conserve energy when  $Fe_2O_3(s)$  is the sole terminal electron acceptor available. The *mtr*<sup>0.37</sup> strain maintained 27% of the initial cfu mL<sup>-</sup> 202 203 <sup>1</sup>, indicating that *mtr* improved survival somewhat over native *E. coli*. The most robust strain was the  $cymAmtr^{0.001}$  strain, which increased its cfu mL<sup>-1</sup> 5% above its initial value (Figure 3B). 204 205 More broadly, comparing the amount of Fe(II) produced to the relative cell density after 21 days 206 (Figure S5), shows there is a strong, positive correlation between the degree of reduction and 207 cell survival in CymA-containing strains (Pearson correlation coefficient, R = 0.74). Taken 208 together, these observations indicate that these CymA-expressing *E. coli* strains use metal 209 reduction to maintain biomass under both Fe(III) citrate and Fe<sub>2</sub>O<sub>3</sub>(s) reducing conditions.

## The expression of CymA significantly increases extracellular electron transfer by enabling catalytic turnover of Mtr.

212 Since CymA can reduce MtrA in vitro, we hypothesized that CymA co-expression 213 increased  $Fe_2O_3$  (s) reduction and cell viability by more rapidly reducing the MtrCAB complex 214 than native E. coli inner membrane quinone dehydrogenases. To directly investigate this, we 215 turned to bioelectrochemical techniques that can monitor the redox turnover of MtrCAB. Since 216 these techniques use a carbon felt electrode as the terminal electron acceptor instead of  $Fe_2O_3(s)$ , we first monitored current production from the *ccm*,  $mtr^{0.18}$ , and *cymAmtr*<sup>0.18</sup> strains in 217 218 bioelectrochemical reactors. The working electrode chamber contained cell suspensions in M1 219 media supplemented with 40 mM D,L-lactate, and the counter electrode chamber contained 50 220 mM PIPES buffer (pH 7.4), and was separated by a cation exchange membrane. As expected, the 221 *ccm* strain, which does not express any *S. oneidensis* cyts *c*, produced very low current (Figure **4A**). The  $mtr^{0.18}$  strain did not produce significantly more current than the *ccm* strain, while in 222 contrast, the  $cymAmtr^{0.18}$  strain produced an average of ~4-fold more current than the  $mtr^{0.18}$ 223 224 strain (Figure 4A). This increase between these two strains is comparable to the increase in 225  $Fe_2O_3$  (s) reduction, suggesting that the cause of improved extracellular electron transfer routes 226 in these strains is independent of the identity of the terminal electron acceptor. Next, we 227 monitored reduction of the Mtr pathway by intracellular reducing equivalents using turnover cyclic voltammetry, scanning at 2 mV/s. Both the *ccm* and  $mtr^{0.18}$  strains exhibited an 228 229 irreversible oxidation peak of unknown origin, but did not show a catalytic wave, i.e. an s-shaped 230 peak, indicative of re-reduction. This indicates that the catalytic turnover of the Mtr pathway is

extremely slow in the absence of CymA. In contrast, the turnover cyclic voltammogram of the  $cymAmtr^{0.18}$  strain (**Figure 4B**) showed a clear catalytic wave, centered just below 0.0 V<sub>Ag/AgCl</sub>, which is consistent with previous measurements of the formal potential of the Mtr pathway in *S*. *oneidensis* MR-1 (*37*). The presence of this wave indicates that the Mtr pathway can be rereduced in the presence of CymA under turnover conditions. This rapid turnover additionally shows that there are properly assembled electron conduits in the engineered *E. coli*, however we have not yet determined what fraction of the electron conduits are correctly assembled.

238 Taken together, these data support our initial observation that re-reduction of MtrA is a 239 rate-limiting step for Fe(III) reduction in *mtrA E. coli* (6) and show that CymA increases the rate 240 of Fe(III) reduction in both MtrA- and MtrCAB-expressing E. coli by increasing the rate of re-241 reduction of MtrA. The ~15 nm thick periplasm in E. coli (38) is too thick to allow CymA to 242 directly contact MtrA within an MtrCAB complex, however, we observe a significant amount of 243 MtrA in the periplasmic fraction (Figure S6). If MtrA diffuses in the periplasm with a diffusion constant similar to comparably sized *E. coli* proteins, i.e.  $\sim 2 \mu m^2 s^{-1}$  (39), it would only require 3 244 245 us on average to diffuse from the inner membrane to the outer membrane. This timescale is 3 246 orders of magnitude faster than the ms required for electron transfer across the MtrCAB conduit 247 (20). Coupled with the observation that CymA can reduce MtrA in vitro (40), the simplest 248 explanation of these data is that, in E. coli, CymA directly reduces MtrA and MtrA diffuses 249 across the periplasmic space to shuttle electrons to MtrC.

# The number of electron nanoconduits does not significantly affect solid Fe<sub>2</sub>O<sub>3</sub> reduction and cell survival.

Using the data we gathered on Fe(III) reduction for *cymAmtr* and *mtr* strains at different induction levels, we next probed the relationship between the number of electron conduits in the

254 *mtr* and *cymAmtr* strains and iron reduction. We use [heme c] as a measure of the number of 255 electron conduits because the stoichometry of cyts c is unchanged with different induction levels 256 (Figure S3) in our *E. coli* strains. Pearson correlation analysis shows a strong correlation 257 between [heme c] and [Fe(II)] in both strains, with the *mtr* strain showing a positive correlation 258 and the *cymAmtr* showing a negative correlation (Pearson correlation coefficients  $R_{mtr} = 0.95$  and  $R_{cymAmtr} = -0.84$ , Figure S7). However, the slope of these correlations are fairly small ( $m_{mtr} =$ 259 0.31 [Fe(II)] [heme c]<sup>-1</sup> and  $m_{cvmAmtr} = -0.52$  [Fe(II)] [heme c]<sup>-1</sup>): for example, the Fe(III) 260 261 reduction rate in the *mtr* strain changes only ~1.5-fold when the number of *mtr* electron conduits changes ~5-fold. This indicates that, for either strain, the number of conduits is not a strong 262 263 determinant of the reduction of solid iron oxide.

264 The finding that the number of electron conduits in engineered E. coli is not a strong 265 determinant of  $Fe_2O_3$  (s) reduction suggests that electron transfer through the Mtr complex is not 266 a rate-limiting step in iron reduction in these strains. To further test this idea, we estimated what the rate of iron reduction in E. coli would be if transport through the MtrCAB conduit were rate-267 limiting. Using our diffuse reflectance data of our best performing *cymAmtr*<sup>0.001</sup> strain (Figure 268 **1D**) and assuming that 90% of the heme c is in MtrCAB complexes (Figure 1B), we estimate the 269 number of Mtr electron nanoconduits per E. coli cell is 50,000 complexes cell<sup>-1</sup>. This is 270 comparable to the number of MtrC in S. oneidensis, which is ~70,000 cell<sup>-1</sup> (21). Together with 271 estimates on the maximal rate of electron transfer through the Mtr electron nanoconduit ( $10^3 \text{ e-s}^-$ 272 <sup>1</sup> complex<sup>-1</sup>) (20), we can approximate the maximal rate of iron reduction per  $cymAmtr^{0.001} E$ . 273 *coli* cell would be ~8 x  $10^{-17}$  M Fe(II) s<sup>-1</sup> cell<sup>-1</sup> (Supplemental Calculation 1). This theoretical 274 maximum rate is ~2000-fold higher than what we observe (~4 x  $10^{-20}$  M Fe(II) s<sup>-1</sup> cell<sup>-1</sup>). Thus, 275

we conclude that electron transfer through the Mtr conduit is not rate-limiting for extracellularelectron transfer in these strains of engineered *E. coli*.

#### 278 Flavins couple solid Fe<sub>2</sub>O<sub>3</sub> reduction to transient growth in the *cymAmtr* strain.

279 Lastly, we considered how addition of 5  $\mu$ M riboflavin affected iron reduction and cell 280 survival in the engineered E. coli strains. We found that riboflavin increased the total amount of reduced iron by 2 and 2.5 fold in the  $mtr^{0.18}$  and  $cymAmtr^{0.18}$  strains over those strains without 281 282 riboflavin added, respectively (Figure 5A, Figure S8). To quantitatively assess the fraction of 283 electrons that reduce via direct contact or flavin-dependent mechanisms, we performed 284 experiments in M1 medium, a trace medium with no riboflavin or casein added (See 285 Supplemental Information), with and without supplementation of riboflavin. We assumed that 286 cultures without riboflavin reduce iron solely by direct contact, while cultures supplemented with 287 riboflavin reduce iron by both direct contact and through a riboflavin-mediated mechanism. This 288 allowed us to calculate the percentage of flavin-dependent iron reduction by comparing the iron 289 reduction in the supplemented and non-supplemented cultures. The resulting percentage of iron reduced in a flavin-dependent process in the *mtr*<sup>0.18</sup> and *cymAmtr*<sup>0.18</sup> strains was 72% and 77%, 290 291 respectively (Supplemental Calculation 2). These percentages closely match the ratio observed 292 in S. oneidensis MR-1 (25, 26). This indicates that, like in the native organism, MtrC is able to 293 utilize solid metal oxides as terminal electron acceptors via both direct contact and through 294 flavin-mediated mechanisms.

Interestingly, while riboflavin had similar relative effects on iron reduction in both the mtr and cymAmtr strains, it had strikingly different effects on cell growth in these strains. The addition of riboflavin did not statistically change the survival of the  $mtr^{0.18}$  strain (**Figure S8**). This observation indicates that riboflavin by itself does not support growth under metal reducing

conditions. In the case of the cymAmtr<sup>0.18</sup> strain, supplementation with riboflavin allowed the 299 cells to grow, increasing the cfu mL<sup>-1</sup> 10-fold over the original suspension after 4 days (**Figure** 300 301 **5B**). This cfu increase is temporally associated with an initial period of rapid reduction of Fe(III) in the  $cvmAmtr^{0.18}$  strain but not the *ccm* strain (Figure 5C, Figure S9). Following this period. 302 303 the cfu decrease and the increase in Fe(II) concentration slows (days 4-10). After 10 days, the 304 cell density drops below its initial value and the Fe(II) concentration remains constant, even 305 though a large excess of Fe(III) is still present. Given the correlations between iron reduction and 306 cell viability, we suggest that the initial high rate of Fe(III) reduction conferred by riboflavin and 307 CymA is enough to allow the E. coli to conserve energy and grow. However, as the Fe(III) 308 reduction rate in this strain slows, perhaps due to a depletion of energy stores and slow oxidation 309 of lactate, the E. coli can no longer conserve energy and the cell density decreases 310 correspondingly. Eventually, the cell density is so low that there is no detectable change in Fe(II) 311 concentration. This scenario would be consistent with observations in E. coli, Shewanella sp., 312 and other microbes that the respiratory rate is positively correlated with the growth rate (37, 41). The cvmAmtr<sup>0.18</sup> E. coli strain generates 200 µM Fe(II) over the course of 10 days, which 313 corresponds to a cell normalized reduction rate of  $\sim 8 \times 10^{-20}$  M Fe(II) s<sup>-1</sup> cell<sup>-1</sup>. Also, this strain 314 315 can maintain cell density at or above its initial level over this same period, which marks the first

time an engineered *E. coli* strain has been shown to grow under solid metal-reducing conditions.
These are significant improvements over previously described *E. coli* strains capable of solid
metal reduction. Additionally, this work strongly suggests that further improvements in
extracellular metal reduction will also boost cell growth in these strains.

#### 320 **Opportunities for additional engineering of metal-reducing** *E. coli.*

321 The fastest iron reduction rate in the engineered *E. coli* cultures described here is still  $\sim$ 50 times slower than S. oneidensis MR-1 ( $\sim 8 \times 10^{-20}$  M Fe(II) s<sup>-1</sup> cell<sup>-1</sup> vs.  $\sim 4 \times 10^{-18}$  M Fe(II) s<sup>-1</sup> cell<sup>-1</sup>) 322 323 (42). We attribute the strain's inability to sustain growth for long periods of time to this slower 324 rate and suggest further work is still required in strain engineering for demanding applications. 325 While our cyclic voltammetry data indicates that a significant fraction of the MtrCAB electron 326 nanoconduits are correctly assembled in the outer membrane of E. coli, we cannot rule out that 327 some fraction of MtrCAB complexes are misassembled and non-functional in E. coli. Assessing 328 and potentially improving this assembly may improve extracellular metal reduction. 329 Additionally, the difference between our cymAmtr E. coli strains and S. oneidensis MR-1 may 330 reflect our still-evolving understanding of electron transfer in S. oneidensis. While this work was 331 in preparation, new data generated by Sturm et al. (19) and Alves et al. (17) strongly suggested 332 that Stc and/or FccA may be involved in shuttling electrons across the periplasm. Thus, co-333 expression of Stc and/or FccA may help extracellular electron transfer in Mtr E. coli.

334 The difference between the electron transfer rates in engineered E. coli and S. oneidensis 335 MR-1 may also arise from the fact that that efficient extracellular electron transfer relies on 336 many processes besides transport of electrons from inner membrane to extracellular acceptors 337 (43). Specifically, extracellular electron transfer also depends on the import and oxidation of an 338 electron donor, transfer of the electrons via intracellular redox carriers to the transmembrane 339 pathway, and adhesion of the bacterium to a metal oxide. Several of these processes are slow or 340 disrupted in the E. coli background used here. For example, the reaction rates of the L- and D-341 lactate dehydrogenases in E. coli are ~10 and ~20-fold slower, respectively, than their 342 counterparts in S. oneidensis (44). Additionally, since the Mtr E. coli are grown aerobically 343 before being introduced into anaerobic metal-reducing conditions, these cells may lack the

344 dehydrogenases and quinones needed to efficiently direct reducing equivalents to the Mtr 345 pathway. Lastly, while *Shewanella sp.* can rapidly attach to  $Fe_2O_3$  surfaces (45, 46), the 346 BL21(DE3) derivatives used in this work are disrupted in their ability to initiate attachment to 347 surfaces because they are non-motile (47). Future work will focus on testing these multiple 348 possibilities to increase the Fe(III) reduction rate and cell growth rates in Mtr-expressing *E. coli*.

#### 349 Conclusions

350 This work shows that replicating certain features of the electron transfer pathway of S. 351 oneidensis MR-1 are critical for boosting extracellular electron transfer in E. coli, while others 352 are not. Specifically, we found that increasing the number of Mtr complexes did not increase 353 extracellular electron transfer, but the presence of CymA and riboflavin had a significant impact 354 on extracellular electron transfer and viability. Additionally, we show that as the rate of metal 355 reduction in CymA-expressing *E. coli* increases, these strains gradually transition from rapidly 356 losing biomass to transiently growing under metal-reducing conditions. This knowledge will 357 allow metal-reduction to be more readily introduced into new heterologous hosts.

358 Despite the relatively modest iron reduction in these newest E. coli strains relative to 359 Shewanella sp. or Geobacter sp., the ability of the cymAmtr E. coli strain described here to maintain biomass or, in the presence of riboflavin, grow over limited timescales makes a useful 360 361 new tool for both basic and applied studies. Since E. coli uses many different electron donors 362 compared to Geobacter or Shewanella, it may provide a more versatile tool for bioremediation or 363 biomining. Also, it is well known that both cell surface structures and metal reduction play a role 364 in formation and transformation of metal-containing solids such as metallic nanoparticles and 365 metal-oxide minerals. The ability of our E. coli strains to reduce solid and chelated Fe(III) with a 366 significantly different cell surface chemistry than Shewanella sp. or Geobacter sp. offers an

- 367 opportunity to dissect these separate effects (48). Thus, new *E. coli* strains described herein and
- those enabled by the design rules described herein, will fuel both basic and applied studies.

369

370 Methods

#### 371 **Plasmids and strains.**

372 The primers, plasmids, and strains used in this study are listed in **Tables S2**, S3, and 373 Table 1, respectively. The *ccm* (pEC86), *mtrA* (I5024), and *mtrCAB* (I5023) plasmids were 374 described previously (6). The cymA (I5040), cymAmtrA (I5052), and cymAmtrCAB (I5049) 375 plasmids were constructed for this work. In brief, these plasmids were constructed using PCR 376 amplification of the genes from genomic DNA of Shewanella oneidensis MR-1 using Pfx 377 Platinum polymerase (Invitrogen), digestion of the pSB1ET2 plasmid and PCR fragment(s) with 378 restriction endonucleases (New England Biolabs), ligation of these fragments with T4 DNA 379 ligase (Roche) and, in some cases, site directed mutagenesis (QuikChange II, Agilent 380 Technologies). Detailed descriptions of the assembly of I5040, I5052, and I5049 can be found in 381 the Supporting Information. After sequence verification, the resulting cymA plasmid and 382 cymAmtrA were co-transformed with ccm into BL21(DE3) to make the cymA and cymAmtrA 383 strains, respectively. In contrast, the *mtrCAB* and *cymAmtrCAB* plasmids, were co-transformed 384 with pEC86 into the E. coli strain C43(DE3) (Lucigen, Middleton, WI) to make the mtr and 385 cymAmtr strains, respectively.

#### **Growth conditions and medium composition.**

All strains, unless otherwise specified, were grown in 2xYT medium at 30 °C with 50  $\mu$ g mL<sup>-1</sup> kanamycin; strains containing the pEC86 plasmid were grown with an additional 30  $\mu$ g mL<sup>-1</sup> chloramphenicol. Glycerol stocks were used to inoculate 5 mL 2xYT medium, and cultures were grown overnight at 37 °C with 250-rpm shaking. Then, 500  $\mu$ L of overnight cultures were back-diluted into 50 mL 2xYT medium and grown with 200-rpm shaking for 16 h at 30 °C. When indicated, strains were induced with IPTG at an  $OD_{600nm}$  of 0.5-0.7. IPTG concentration is displayed here as promoter activity defined in Goldbeck, et al (5).

Cell suspensions used for iron reduction assays were resuspended in anaerobic defined M1 medium supplemented with 40 mM D,L-lactate and 0.2% casamino acids. The composition of the M1 medium can be found in the Supporting Information.

#### 397 Subcellular Fractionation.

398 Periplasmic and membrane preparations were performed as previously described (6).
399 Membrane samples were solubilized in a solution of 5% (w/v) Triton X-100, 50 mM HEPES pH
400 7.4, and 200 mM NaCl.

### 401 Visible spectra of cytochrome samples by diffuse reflectance.

The concentration of cytochromes in whole cells was determined by diffuse reflectance as previously described by Goldbeck et al. (5). The reduced samples were chemically reduced with sodium dithionite crystals (Sigma, St. Louis, MO). The extinction coefficients at 552 nm of  $104 \text{ mM}^{-1} \text{ cm}^{-1}$  (personal communication, Prof. Julea Butt) and 280 mM<sup>-1</sup> cm<sup>-1</sup> (35) were used to determine the CymA and MtrA concentrations, respectively.

### 407 ECL to detect *c*-type cytochromes in whole cell lysates.

Cell pellets from 1.5 mL of culture was resuspended in 0.1 mL Bacterial Protein Extraction Reagent (B-Per, ThermoScientific, Grand Island, NY). The cells were frozen at -20 °C immediately after growth and then thawed immediately before analysis. Cells were lysed for 30 minutes at room temperature with 6  $\mu$ g mL<sup>-1</sup> chicken egg white lysozyme (Sigma), 1  $\mu$ g mL<sup>-1</sup> DNAase, 3.9 mM MgSO<sub>4</sub>, 0.96 mM EDTA, and 0.98 mM phenylmethylsulfonyl fluoride. Total protein of the cell lysates was determined by BCA Protein Assay Kit (ThermoScientific). Cells were diluted in 100 mM HEPES, pH 7.4, to normalize to equal total protein concentration. 415 Samples were prepared with NuPAGE 4x Sample Buffer (Bio-Rad) and heated at 95 °C for 5 416 minutes. A total of 8  $\mu$ g protein was loaded and run in a 4-20% Tris-HCl polyacrylamide gel 417 (Bio-Rad) at 200 V for 1 hour. The gel was rinsed twice in water and then equilibrated in cold 418 Pierce Western Transfer buffer (ThermoScientific) for 15 minutes. The proteins were transferred 419 to a 0.45  $\mu$ m nitrocellulose membrane (Bio-Rad, Hercules, CA) in Pierce Western Transfer 420 buffer at 30 V for 100 minutes. Ponceau S staining was used to confirm uniform transfer across 421 all lanes.

The nitrocellulose membrane was incubated for 5 minutes in 10 mL of Pierce Pico West Enhanced Chemiluminescence substrate (ThermoScientific), a 1:1 mixture of Pico West Peroxide Solution and Luminol Enhancer solution. The chemiluminescent signal was detected using the ChemiDoc<sup>TM</sup> XRS system. The chemiluminescent signal and molecular weights were quantified using ImageJ (*49*).

#### 427 Assaying soluble Fe(III) citrate reduction and cell density of strains.

428 Fe(III) reduction assays were performed as previously described (6). Briefly, cultures 429 grown aerobically were pelleted, washed, and resuspended to an  $OD_{600 \text{ nm}}$  of 0.5 in anaerobic M1 430 medium supplemented with 40 mM D,L-lactate as the sole carbon source. All anaerobic media 431 and buffers were sparged with nitrogen and anaerobic conditions were maintained in an 432 anaerobic chamber (Coy Laboratory Products, Grass Lake, MI) with an atmosphere of 2% H<sub>2</sub> 433 balance N<sub>2</sub>. Fe(III) citrate (Sigma, St. Louis, MO) was added to a final concentration of 10 mM. 434 The Fe(II) concentration was determined with the ferrozine assay, adapted from Stookey (50). 435 The concentration of Fe(II) in each culture was subtracted by abiotic iron reduction observed in 436 sterile medium-only controls at each time point. Additionally, a lysed mtrA control was included 437 to determine the extent of non-metabolic Fe(III) reduction (Supporting Information). Cell density 438 was determined by measuring the optical density at 600 nm and subtracting the scattering of 439 abiotic media with 10mM Fe(III) citrate. Error bars represent the standard deviation across three 440 biological replicates. The rate of Fe(III) citrate reduction and linear rate of change in optical 441 density were calculated by best linear fit for each individual biological replicate.

442 Bulk Fe<sub>2</sub>O<sub>3</sub> (*s*) reduction assay.

Fe<sub>2</sub>O<sub>3</sub> (*s*) reduction assays were performed as previously described (*6*). Briefly, cultures grown aerobically were pelleted, washed, and resuspended to an  $OD_{600 \text{ nm}}$  of 1.0 in anaerobic M1 medium supplemented with 40 mM D,L-lactate as the carbon source, 6.0 mg mL<sup>-1</sup> Fe<sub>2</sub>O<sub>3</sub> (*s*) (Sigma) as the sole terminal electron acceptor, and IPTG, where indicated. Where indicated, the media was supplemented with 5  $\mu$ M riboflavin.

The colony forming units (cfu) and Fe(II) concentration for each culture was measured after initial anaerobic inoculation and at time points following. Cfu's were determined with kanamycin selection on LB plates grown aerobically at 37 °C. The Fe(II) concentration was determined with the ferrozine assay, adapted from Stookey (*50*). The concentration of Fe(II) in each culture was subtracted by any abiotic iron reduction observed in sterile media-only controls at each time point. Error bars represent standard deviation of triplicate cultures.

#### 454 Cyclic Voltammetry of *mtr* and *cymAmtr* strains.

455 Cells from 50-mL cultures were pelleted and washed twice with M1 medium 456 supplemented with 40 mM D,L-lactate. Pellets were resuspended to an  $OD_{600nm}$  of 0.7 in the 457 working chamber of 2-chambered bioelectrochemical reactors. The working chamber contained 458 M1 medium supplemented with 40 mM D,L-lactate as the sole carbon source, the counter 459 electrode chamber contained 50 mM PIPES buffer (pH 7.4), and the two chambers were 460 separated by a cation exchange membrane (CMI-7000, Membranes International, Ringwood,

461 NJ). The working electrode was a 25 x 25 mm square piece of graphite felt (GF-S6-06, 462 Electrolytica, Amherst, NY) and was connected to the potentiostat via a piece of Pt wire. The 463 counter electrode was a piece of Pt wire and the reference electrode was a pre-made Ag/AgCl 464 reference (CH Instruments, Bee Cave, TX). During current stabilization, the working chambers 465 were stirred at ~200 rpm with magnetic stir bars, and stirring was turned off during cyclic 466 voltammetry analysis. Cyclic voltammograms were recorded in 2 cycles from -0.5 V<sub>Ag/AgCl</sub> to 467 +0.5 V<sub>Ag/AgCl</sub> at a scan rate of 2 mV/s. Cyclic voltammograms were recorded before and 468 immediately after inoculation. After initial cyclic voltammetry, the working electrodes were held 469 at +0.2 V<sub>Ag/AgCl</sub> overnight to allow cells to associate with the electrode surface. Cyclic 470 voltammograms were recorded again after the overnight period.

471

#### 472 Supporting Information.

473 SI Figure 1. Schematic of plasmids and localization of MtrC, MtrA, and CymA.

474 SI Figure 2. Diffuse reflectance spectra show the spectral signatures characteristic of 475 cytochromes *c* in whole cell suspensions.

476 SI Figure 3. ECL analysis shows that the relative abundance of each cyt c within a strain is 477 similar across different induction levels

478 SI Figure 4. Homogenized MtrA *E. coli* reduce Fe(III) citrate at the same basal rate as *ccm*.

- 479 SI Figure 5. Fitness is strongly correlated to iron reduction rate.
- 480 SI Figure 6. A scaled schematic of the gram-negative double membrane and Mtr pathway.
- 481 SI Figure 7. Correlation analysis of iron reduced against heme *c* concentration.
- 482 SI Figure 8. Riboflavin mildly improves Fe<sub>2</sub>O<sub>3</sub> reduction, but not survival.

483	SI Figure	9.	The	сст	strain	shows	no	temporal	correlation	between	cell	density	and	Fe <sub>2</sub> O <sub>3</sub>
484	reduction.													

- 485 SI Table 1. Relative promoter activity as a function of IPTG concentration.
- 486 SI Table 2. Primers
- 487 SI Table 3. Plasmids
- 488 Supporting Calculation 1. Approximation of maximum iron reduction rate.
- 489 Supporting Calculation 2. Percent Fe<sub>2</sub>O<sub>3</sub> reduced from direct contact or riboflavin-mediated
- 490 mechanisms.
- 491 Supporting Methods.
- 492

#### 493 Abbreviations

494 Colony forming units: cfu. Cytochrome *c*: cyt *c*. Cytochrome *c* maturation: ccm. Enhanced
495 chemiluminescence: ECL. Isopropyl β-D-1-thiogalactopyranoside: IPTG.

496

497 Author Information

498

- 499 Authors' contributions
- 500 HMJ and CMAF conceived the study. HMJ, MAT, MGK performed the experiments, and
- 501 CMAF supervised the work. HMJ analyzed the data. HMJ, MAT, MGK and CMAF drafted the
- 502 manuscript, which was read, revised and approved by all authors.

503

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## 515 **References**

- Fredrickson, J. K., Zachara, J. M., Kennedy, D. W., Dong, H., Onstott, T. C., Hinman, N.
   W., and Li, S.-M. (1998) Biogenic iron mineralization accompanying the dissimilatory reduction of hydrous ferric oxide by a groundwater bacterium, *Geochim. Cosmochim. Acta* 62, 3239 –3257.
- S20 2. Narayanan, K., and Sakthivel, N. (2010) Biological synthesis of metal nanoparticles by
  microbes, *Adv. Colloid Interface Sci. 156*, 1-13.
- 522 3. Zhuang, W.-Q., Fitts, J. P., Ajo-Franklin, C. M., Maes, S., Alvarez-Cohen, L., and
  523 Hennebel, T. (2015) Recovery of critical metals using biometallurgy, *Curr. Opin.*524 *Biotechnol. 33*, 327-335.
- 525 4. Gadd, G. M. (2010) Metals, minerals and microbes: geomicrobiology and bioremediation, *Microbiology 156*, 609-643.
- 527 5. Goldbeck, C. P., Jensen, H. M., TerAvest, M. A., Beedle, N., Appling, Y., Hepler, M.,
  528 Cambray, G., Mutalik, V., Angenent, L. T., and Ajo-Franklin, C. M. (2013) Tuning
  529 Promoter Strengths for Improved Synthesis and Function of Electron Conduits in
  530 *Escherichia coli, ACS Synth. Biol.* 2, 150-159.
- 531 6. Jensen, H. M., Albers, A. E., Malley, K. R., Londer, Y. Y., Cohen, B. E., Helms, B. A.,
  532 Weigele, P., Groves, J. T., and Ajo-Franklin, C. M. (2010) Engineering of a synthetic
  533 electron conduit in living cells, *Proc. Natl. Acad. Sci. U. S. A.* 107, 19213-19218.
- TerAvest, M. A., Zajdel, T. J., and Ajo-Franklin, C. M. (2014) The Mtr Pathway of *Shewanella oneidensis* MR-1 Couples Substrate Utilization to Current Production in *Escherichia coli, ChemElectroChem 1*, 1874-1879.
- 537 8. Gescher, J. S., Cordova, C. D., and Spormann, A. M. (2008) Dissimilatory iron reduction
- in *Escherichia coli*: identification of CymA of *Shewanella oneidensis* and NapC of *E. coli* as ferric reductases, *Mol. Microbiol.* 68, 706-719.

540	9.	Schuetz, B., Schicklberger, M., Kuermann, J., Spormann, A. M., and Gescher, J. (2009)
541		Periplasmic Electron Transfer via the c-Type Cytochromes MtrA and FccA of
542		Shewanella oneidensis MR-1, Appl. Environ. Microbiol. 75, 7789-7796.
543	10.	Sturm-Richter, K., Golitsch, F., Sturm, G., Kipf, E., Dittrich, A., Beblawy, S.,
544		Kerzenmacher, S., and Gescher, J. (2015) Unbalanced fermentation of glycerol in
545		Escherichia coli via heterologous production of an electron transport chain and electrode
546		interaction in microbial electrochemical cells, <i>Bioresource Technol 186</i> , 89-96.
547	11.	Ajo-Franklin, C. M., and Noy, A. (2015) Crossing Over: Nanostructures that Move
548		Electrons and Ions across Cellular Membranes, Adv. Mater. 27, 5797–5804.
549	12.	Richardson, D. J., Butt, J. N., Fredrickson, J. K., Zachara, J. M., Shi, L., Edwards, M. J.,
550		White, G., Baiden, N., Gates, A. J., Marritt, S. J., and Clarke, T. A. (2012) The 'porin-
551		cytochrome' model for microbe-to-mineral electron transfer, Mol. Microbiol. 85, 201-
552		212.
553	13.	Shi, L., Rosso, K. M., Clarke, T. A., Richardson, D. J., Zachara, J. M., and Fredrickson,
554		J. K. (2012) Molecular Underpinnings of Fe(III) Oxide Reduction by Shewanella
555		oneidensis MR-1, Front. Microbiol. 3, 50.
556	14.	Marritt, S. J., McMillan, D. G. G., Shi, L., Fredrickson, J. K., Zachara, J. M., Richardson,
557		D. J., Jeuken, L. J. C., and Butt, J. N. (2012) The roles of CymA in support of the
558		respiratory flexibility of Shewanella oneidensis MR-1, Biochem. Soc. Trans. 40, 1217-
559		1221.
560	15.	Simon, J., van Spanning, R. J. M., and Richardson, D. J. (2008) The organisation of
561		proton motive and non-proton motive redox loops in prokaryotic respiratory systems,
562		Biochim. Biophys. Acta, Bioenerg. 1777, 1480-1490.
563	16.	McMillan, D. G. G., Marritt, S. J., Butt, J. N., and Jeuken, L. J. C. (2012) Menaquinone-7
564		is specific cofactor in tetraheme quinol dehydrogenase CymA, J. Biol. Chem. 287,
565		14215-14225.
566	17.	Alves, M. N., Neto, S. E., Alves, A. S., and Fonseca, B. M. (2015) Characterization of
567		the periplasmic redox network that sustains the versatile anaerobic metabolism of
568		Shewanella oneidensis MR-1, Front. Microbiol. 6, 665.
569	18.	Fonseca, B. M., Paquete, C. M., Neto, S. E., Pacheco, I., Soares, C. M., and Louro, R. O.
570		(2013) Mind the gap: cytochrome interactions reveal electron pathways across the
571		periplasm of Shewanella oneidensis MR-1, Biochem. J. 449, 101-108.
572	19.	Sturm, G., Richter, K., Doetsch, A., Heide, H., Louro, R. O., and Gescher, J. (2015) A
573		dynamic periplasmic electron transfer network enables respiratory flexibility beyond a
574		thermodynamic regulatory regime, ISME J. 9, 1802-1815.
575	20.	White, G. F., Shi, Z., Shi, L., Wang, Z., Dohnalkova, A. C., Marshall, M. J., Fredrickson,
576		J. K., Zachara, J. M., Butt, J. N., Richardson, D. J., and Clarke, T. A. (2013) Rapid
577		electron exchange between surface-exposed bacterial cytochromes and Fe(III) minerals,
578		Proc. Natl. Acad. Sci. U. S. A. 110, 6346-6351.
579	21.	Ross, D. E., Brantley, S. L., and Tien, M. (2009) Kinetic Characterization of OmcA and
580		MtrC, Terminal Reductases Involved in Respiratory Electron Transfer for Dissimilatory
581		Iron Reduction in Shewanella oneidensis MR-1, Appl. Environ. Microbiol. 75, 5218-
582		5226.
583	22.	Brutinel, E., D., and Gralnick, J., A (2012) Shuttling happens: soluble flavin mediators
584		of extracellular electron transfer in Shewanella, Appl. Microbiol. Biotechnol. 93, 41-48.

585 23. Edwards, M. J., White, G. F., and Norman, M. (2015) Redox Linked Flavin Sites in 586 Extracellular Decaheme Proteins Involved in Microbe-Mineral Electron Transfer, Sci. 587 *Rep.* 5, 11677. 588 24. Okamoto, A., Nakamura, R., and Nealson, K. H. (2014) Bound Flavin Model Suggests 589 Similar Electron Transfer Mechanisms in Shewanella and Geobacter, ChemElectroChem 590 1, 1808-1812. 591 25. Jiang, X. C., Hu, J. S., Fitzgerald, L. A., Biffinger, J. C., Xie, P., Ringeisen, B. R., and 592 Lieber, C. M. (2010) Probing electron transfer mechanisms in Shewanella oneidensis 593 MR-1 using a nanoelectrode platform and single-cell imaging, Proc. Natl. Acad. Sci. U. 594 S. A. 107, 16806-16810. 595 Marsili, E., Baron, D. B., and Shikhare, I. D. (2008) Shewanella secretes flavins that 26. 596 mediate extracellular electron transfer, Proc. Natl. Acad. Sci. U. S. A. 105, 3968-3973. 597 27. Kotloski, N. J., and Gralnick, J. A. (2013) Flavin electron shuttles dominate extracellular 598 electron transfer by Shewanella oneidensis, mBio 4. 599 28. Nikel, P. I., and de Lorenzo, V. (2013) Engineering an anaerobic metabolic regime in 600 Pseudomonas putida KT2440 for the anoxic biodegradation of 1, 3-dichloroprop-1-ene, 601 Metab. Eng. 15, 98-112. 602 29. Schmitz, S., Nies, S., Wierckx, N., Blank, L. M., and Rosenbaum, M. A. (2015) 603 Engineering mediator-based electroactivity in the obligate aerobic bacterium 604 Pseudomonas putida KT2440, Front. Microbiol. 6, 284. 605 Steen, A., Ütkür, F., Acuña, B.-d. J. M., and Bunk, B. (2013) Construction and 30. 606 characterization of nitrate and nitrite respiring *Pseudomonas putida* KT2440 strains for 607 anoxic biotechnical applications, J. Biotechnol. 163, 155-165. 608 31. Salis, H. M., Mirsky, E. A., and Voigt, C. A. (2009) Automated design of synthetic 609 ribosome binding sites to control protein expression, Nat. Biotechnol. 27, 946-950. 610 Arslan, E., Schulz, H., Zufferey, R., Künzler, P., and Thöny-Meyer, L. (1998) 32. 611 Overproduction of the Bradyrhizobium japonicum c-Type Cytochrome Subunits of the 612 cbb 3 Oxidase in Escherichia coli, Biochem. Biophys. Res. Commun. 251, 744-747. 613 33. Myers, C. R., and Myers, J. M. (1997) Cloning and sequence of cymA, a gene encoding a 614 tetraheme cytochrome c required for reduction of iron (III), fumarate, and nitrate by 615 Shewanella putrefaciens MR-1, J. Bacteriol. 179, 1143-1152. 616 34. Myers, J. M., and Myers, C. R. (2002) Genetic complementation of an outer membrane 617 cytochrome omcB mutant of Shewanella putrefaciens MR-1 requires omcB plus 618 downstream DNA, Appl. Environ. Microbiol. 68, 2781-2793. 619 35. Pitts, K. E., Dobbin, P. S., Reyes-Ramirez, F., Thomson, A. J., Richardson, D. J., and Seward, H. E. (2003) Characterization of the Shewanella oneidensis MR-1 Decaheme 620 621 Cytochrome MtrA, J. Biol. Chem. 278, 27758-27765. 622 36. Wang, V. B., Yantara, N., Koh, T. M., Kjelleberg, S., Zhang, Q., Bazan, G. C., Loo, S. C. 623 J., and Mathews, N. (2014) Uncovering alternate charge transfer mechanisms in 624 Escherichia coli chemically functionalized with conjugated oligoelectrolytes, Chem. Commun. (Cambridge, U. K.) 50, 8223-8226. 625 626 Carmona-Martinez, A. A., Harnisch, F., Fitzgerald, L. A., Biffinger, J. C., Ringeisen, B. 37. 627 R., and Schröder, U. (2011) Cyclic voltammetric analysis of the electron transfer of 628 Shewanella oneidensis MR-1 and nanofilament and cytochrome knock-out mutants, 629 Bioelectrochemistry 81, 74-80.

630	38.	Matias, V. R. F., Al-Amoudi, A., Dubochet, J., and Beveridge, T. J. (2003) Cryo-
631		transmission electron microscopy of frozen-hydrated sections of Escherichia coli and
632		Pseudomonas aeruginosa, J. Bacteriol. 185, 6112-6118.
633	39.	Mullineaux, C. W., Nenninger, A., and Ray, N. (2006) Diffusion of green fluorescent
634		protein in three cell environments in Escherichia coli, J. Bacteriol. 188, 3442-3448.
635	40.	Firer-Sherwood, M. A., Bewley, K., D., Mock, JY., and Elliott, S. J. (2011) Tools for
636		resolving complexity in the electron transfer networks of multiheme cytochromes c,
637		Metallomics 3, 344-348.
638	41.	Andersen, K. B., and Meyenburg, K. (1980) Are growth rates of <i>Escherichia coli</i> in batch
639		cultures limited by respiration?, J. Bacteriol. 144, 114-123.
640	42.	Bose, S., Hochella, M. F., Gorby, Y. A., Kennedy, D. W., McCready, D. E., Madden, A.
641		S., and Lower, B. H. (2009) Bioreduction of hematite nanoparticles by the dissimilatory
642		iron reducing bacterium Shewanella oneidensis MR-1, Geochim. Cosmochim. Acta 73,
643		962-976.
644	43.	TerAvest, M. A., and Ajo-Franklin, C. M. (2015) Transforming exoelectrogens for
645		biotechnology using synthetic biology, Biotechnol. Bioeng. in press, DOI:
646		10.1002/bit.25723.
647	44.	Pinchuk, G. E., Rodionov, D. A., Yang, C., Li, X., Osterman, A. L., Dervyn, E.,
648		Geydebrekht, O. V., Reed, S. B., Romine, M. F., Collart, F. R., Scott, J. H., Fredrickson,
649		J. K., and Beliaev, A. S. (2009) Genomic reconstruction of Shewanella oneidensis MR-1
650		metabolism reveals a previously uncharacterized machinery for lactate utilization, Proc.
651		Natl. Acad. Sci. U. S. A. 106, 2874-2879.
652	45.	Bonneville, S., Behrends, T., Cappellen, P., Hyacinthe, C., and Röling, W. F. M. (2006)
653		Reduction of Fe (III) colloids by Shewanella putrefaciens: A kinetic model, Geochim.
654		Cosmochim. Acta 70, 5842-5854.
655	46.	Glasauer, S., Langley, S., and Beveridge, T. J. (2001) Sorption of Fe (hydr) oxides to the
656		surface of Shewanella putrefaciens: cell-bound fine-grained minerals are not always
657		formed de novo, Appl. Environ. Microbiol. 67, 5544-5550.
658	47.	Pratt, L. A., and Kolter, R. (1998) Genetic analysis of <i>Escherichia coli</i> biofilm formation:
659		roles of flagella, motility, chemotaxis and type I pili, Mol. Microbiol. 30, 285-293.
660	48.	De Windt, W., Boon, N., and Van den Bulcke, J. (2006) Biological control of the size and
661		reactivity of catalytic Pd (0) produced by Shewanella oneidensis, Antonie van
662		Leeuwenhoek 90, 377-389.
663	49.	Schneider, C. A., Rasband, W. S., and Eliceiri, K. W. (2012) NIH Image to ImageJ: 25
664		years of image analysis, Nat. Methods 9, 671-675.
665	50.	Stookey, L. L. (1970) Ferrozinea new spectrophotometric reagent for iron, Anal.
666		<i>Chem.</i> 42, 779-781.
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Strain	Parental Strain	Plasmid(s)	Gene(s)	Source	Request Name
сст	BL21(DE3)	pEC86, pSB1ET2	ccmA-H	Jensen 2010	MFe208
mtrA	BL21(DE3)	pEC86, I5024	ccmA-H, mtrA	Jensen 2010	MFe291
cymA	BL21(DE3)	pEC86, I5040	сстА-Н, сутА	this work	MFe431
cymAmtrA	BL21(DE3)	pEC86, I5052	ccmA-H, cymAmtrA	this work	MFe538
сст	C43(DE3)	pEC86, pSB1ET2	ccmA-H	Goldbeck 2013	MFe408
mtr	C43(DE3)	pEC86, I5023	ccmA-H, mtrCAB	this work	MFe409
cymAmtr	C43(DE3)	pEC86, I5049	ccmA-H, cymAmtrCAB	this work	MFe444

670 **Table 1. Strains used in this study.** 

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673 Figure 1. CymA is co-expressed with MtrCAB to provide an extracellular electron transfer 674 pathway that spans both membranes of E. coli. (A) Schematic of the Mtr pathway in S. 675 oneidensis. Electrons present in menaquinone are routed across the inner membrane by CymA 676 (blue box) and are then sequentially transferred to MtrA and to MtrC through the MtrCAB 677 electron nanoconduits (red box). Electrons can then be passed from MtrC to  $Fe_2O_3$  either directly 678 or through a flavin-mediated process (green box). (IM = inner membrane. OM = outer 679 membrane.) (B) Heme staining of whole cell lysates of the cymA, cymAmtrA, and mtrA strains 680 show CymA and MtrA are present in the respective strains. Similarly, heme staining of whole 681 cell lysates of the *mtr* and *cymAmtr* strain shows both strains contain MtrA and MtrC. Heme 682 staining was performed via ECL in (B). (C) Densitometry of the cymAmtrA (green), mtr (red), 683 and cymAmtr (blue) containing lanes shown in (B) show that CymA is present only in the 684 cymAmtr strain. (D) Heme c concentration per cell density for the mtr and cymAmtr strains as a 685 function of relative promoter activity. Heme c concentration was measured by diffused 686 reflectance and relative promoter activity is measured as Goldbeck et al (5).

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Figure 2. Co-expression of CymA with MtrA improves Fe(III) citrate reduction and maintains biomass in *E. coli*. (A) Fe(II) concentration as a function of time for the *ccm* (open black circles), *mtrA* (closed red circles), *cymA* (half-filled green circles), and *cymAmtrA* (halffilled blue circles) strains. The *cymA* and *cymAmtrA* strains reduce Fe(III) citrate ~3x and ~4x faster than the *mtrA* strain. (B) Change in cell density over time for the *ccm* (open black circles), *mtrA* (closed red circles), *cymA* (half-filled green circles), and *cymAmtrA* (half-filled blue circles)
strains. Only the *cymAmtrA* strain maintains constant biomass under Fe(III) reducing conditions.

696 Figure 3. Co-expression of CymA with MtrCAB increases Fe<sub>2</sub>O<sub>3</sub> (s) reduction and 697 maintains biomass in *E. coli*. (A) The concentration of bulk  $\alpha$ -Fe<sub>2</sub>O<sub>3</sub> reduced to Fe(II) by *ccm* and the best performing *mtr* and *cymAmtr* strains, showing that  $cymAmtr^{0.001}$  significantly 698 increased iron reduction over both the *ccm* and the  $mtr^{0.37}$  strains (p = 0.002 and 0.031, 699 respectively). This indicates that the co-expression of CymA with Mtr increases the flux of 700 electrons to iron oxide. (B) Relative cell density of the *ccm*.  $mtr^{0.37}$  and *cvmAmtr*<sup>0.001</sup> strains after 701 702 4 days under  $Fe_2O_3$  (s) reducing conditions, showing that only the *cymAmtr* strain maintains 703 constant biomass. The relative cell density is the cell density after 4 days divided by the initial 704 cell density. Average values (n=3) are plotted and error bars represent standard deviation.

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#### 706 Figure 4. The expression of CymA significantly increases current production by enabling

707catalytic turnover of Mtr. (A) Current production in electrochemical reactors with working708electrodes poised at +0.2  $V_{Ag/AgCl}$  increased significantly when *cymA* was co-expressed with *mtr*.709Average values (n=4) are plotted and error bars represent standard deviation. (B) Turnover cyclic710voltammetry (representative scans plotted) at 2 mV/s reveals a catalytic wave only when *cymA* is711co-expressed with *mtr*, indicating that the inner membrane cytochrome is necessary for catalytic712turnover of the outer membrane cytochromes.

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714 Figure 5. Riboflavin improves  $Fe_2O_3$  (s) reduction and permits growth in the cymAmtr strain. (A) Riboflavin significantly enhanced iron reduction by ~2.5x in the  $cymAmtr^{0.18}$  strain (p 715 716 = 0.001). Assuming direct contact contributes equally in the cultures with and without the 717 supplementation of riboflavin, 77% of the iron oxide was reduced by riboflavin. (B) The survival of the *cymAmtr*<sup>0.18</sup> strain dramatically increased in the presence of riboflavin ( $p = 3x10^{-5}$ ), but not 718 in the  $mtr^{0.18}$  strain (p = 0.13). Average values (n = 3) are plotted and error bars represent 719 720 standard deviation. (C) Concentration of Fe(II) produced and colony forming units as a function of time for the  $cymAmtr^{0.18}$  cultures with riboflavin, showing a that the initial period of rapid 721

Fe(II) production is accompanied by increase in cell density for the *cymAmtr* strain with riboflavin.

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