Metatranscriptomics Supports the Mechanism for Biocathode Electroautotrophy by “Candidatus Tenderia electrophaga”


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ABSTRACT Biocathodes provide a stable electron source to drive reduction reactions in electrotrophic microbial electrochemical systems. Electroautotrophic biocathode communities may be more robust than monocultures in environmentally relevant settings, but some members are not easily cultivated outside the electrode environment. We previously used metagenomics and metaproteomics to propose a pathway for coupling extracellular electron transfer (EET) to carbon fixation in “Candidatus Tenderia electrophaga,” an uncultivated but dominant member of an electroautotrophic biocathode community. Here we validate and refine this proposed pathway using metatranscriptomics of replicate aerobic biocathodes poised at the growth potential level of 310 mV and the suboptimal 470 mV (versus the standard hydrogen electrode). At both potentials, transcripts were more abundant from “Ca. Tenderia electrophaga” than from any other constituent, and its relative activity was positively correlated with current. Several genes encoding key components of the proposed “Ca. Tenderia electrophaga” EET pathway were more highly expressed at 470 mV, consistent with a need for cells to acquire more electrons to obtain the same amount of energy as at 310 mV. These included cyc2, encoding a homolog of a protein known to be involved in iron oxidation. Mean expression of all CO2 fixation-related genes is 0.27 log2-fold higher at 310 mV, indicating that reduced energy availability at 470 mV decreased CO2 fixation. Our results substantiate the claim that “Ca. Tenderia electrophaga” is the key electroautotroph, which will help guide further development of this community for microbial electrosynthesis.

IMPORTANCE Bacteria that directly use electrodes as metabolic electron donors (biocathodes) have been proposed for applications ranging from microbial electrosynthesis to advanced bioelectronics for cellular communication with machines. However, just as we understand very little about oxidation of analogous natural insoluble electron donors, such as iron oxide, the organisms and extracellular electron transfer (EET) pathways underlying the electrode-cell direct electron transfer processes are almost completely unknown. Biocathodes are a stable biofilm cultivation platform to interrogate both the rate and mechanism of EET using electrochemistry and to study the electroautotrophic organisms that catalyze these reactions. Here we provide new evidence supporting the hypothesis that the uncultured bacterium “Candidatus Tenderia electrophaga” directly couples extracellular electron transfer to CO2 fixation. Our results provide insight into developing biocathode technology, such as microbial electrosynthesis, as well as advancing our understanding of chemolithoautotrophy.

KEYWORDS biocathode, electroautotroph, metatranscriptomics, microbial electrochemical system
Biocathodes are components of some bioelectrochemical systems (BES) in which microbial electrode catalysts use the electrode as an electron donor to drive cellular metabolism. Over the last decade, biocathodes have been explored for improving energy recovery in microbial fuel cells (MFCs) and electrode-driven bioremediation and, more recently, to produce chemicals in a process known as microbial electrosynthesis (reviewed in references 1 and 2). Despite widespread interest in biocathodes for biotechnology applications, little is understood about the underlying mechanisms of extracellular electron transfer (EET) for biocathode microorganisms. Although biocathode EET has been demonstrated for a variety of microorganisms, including acetogens (3) and a methanogenic archaeon (4), mechanistic studies aimed at identifying EET conduits from the electrode to cells are limited (5–7).

The *Marinobacter-Chromatiaceae-Labrenzia* (MCL) biocathode is a self-regenerating, self-sustaining, microbial community and has served in our laboratory as a model system for the exploration of aerobic electroautotrophic microbial communities using an omics approach (8, 9). MCL forms a heterogeneously distributed biofilm with cellular aggregates up to 20 μm thick (9) and reproducible electrochemical features following inoculation of a portion of the biofilm into a new reactor. It is proposed that MCL reduces O₂ with electrons supplied solely by the cathode, directing a portion of the acquired energy and electrons for autotrophy. Cyclic voltammetry (CV) revealed a sigmoid-shaped dependency for current associated with catalytic O₂ reduction (turnover current) dependent upon electrode potential (10). It is proposed that this dependency reflects Nernstian behavior of the heterogeneous electron transfer reaction (across the biofilm/electrode interface) that is mediated by a redox cofactor, which is fast, reversible, and not the rate-limiting step in direct electron uptake from the cathode by the biofilm (9, 11, 12). Electroautotrophic growth is assumed for MCL based on an increase in biomass correlated to increasing current, a lack of organic carbon in the bioelectrochemical reactor, and identification of an active Calvin-Benson-Bassham (CBB) cycle (13). Electroautotrophic growth of isolates at potentials greater than −100 mV versus the standard hydrogen electrode (SHE) has thus far been demonstrated only for the aerobic Fe(II)-oxidizing bacteria *Mariprofundus ferrooxydans* PV-1 and *Acidithiobacillus ferrooxidans* (14, 15), while other autotrophs require supplemental energy from light or hydrogen for initial growth on a cathode (6, 16). However, recent reports indicate that communities enriched on high potential O₂-reducing biocathodes reproducibly contain large populations of *Gammaproteobacteria* (17, 18).

Previous metagenomic and proteomics studies of MCL have led to the identification of putative EET and CO₂ fixation mechanisms (13, 19), although efforts at cultivation have not yielded the proposed electroautotroph, “*Candidatus Tenderia electrophaga*” (20). Other work has resulted in isolates from aerobic biocathode enrichments, but these have not been autotrophic (21, 22). Metaproteomic analysis suggested that proteins that may be involved in EET, including a homolog of Cyc2, a cytochrome known to be involved in Fe(II) oxidation in *A. ferrooxidans* (13), are present at high levels in the biofilm. Subsequent metaproteomic analysis of the biofilm at two different electrode potentials showed that some components of the electron transport chain (ETC) are differentially expressed, including an ortholog of Cyc1, thought to be involved in Fe(II) oxidation in *M. ferrooxydans* PV-1 (23), and a hypothetical protein homologous to the terminal oxidase cytochrome cbb₃ subunit CcoO (19). However, methodological limitations meant that the nine proteins that were detected significantly more often at one potential than at another were likely to represent a small fraction of the response to changing potential in “Ca. Tenderia electrophaga” (19). More-precise quantification of the changes in relative gene expression obtained using RNA sequencing (RNA-seq) (24) can be used to understand the molecular mechanisms used for growth on the cathode.

Building upon our previous work (9, 13, 19), we applied RNA-seq to MCL to compare the levels of gene expression of proposed EET and CO₂ fixation pathways for “Ca. Tenderia electrophaga” at two different potentials. Adjusting the applied potential from 310 mV versus SHE (optimal for growth) to 470 mV (suboptimal for growth) results in
a decrease in the ΔG° value for the reduction of O₂ by about one-third. From results of previous metaproteomic experiments run under identical conditions, the change in potential of the electron donor is expected to result in changes in gene expression to compensate for the change in energy availability. Our metatranscriptomic results support previously hypothesized roles of some protein complexes (13, 19, 23) and reveal possible EET roles for other proteins. We also provide further evidence that “Ca. Tenderia electrophaga” is the keystone species in this community and is strongly associated with electron uptake rates. The results from the metatranscriptomics analyses provide insight into the molecular mechanisms involved in electroautotrophic growth on a cathode, enabling the development of biocathodes for possible future applications, including synthesis of value-added compounds or biofuels as well as potential ET components for microbial bioelectronics.

RESULTS

Biocathode-MCL metatranscriptome. Eight BES were grown under previously described standard conditions at an applied electrode potential of 310 mV SHE. Seven of the eight biological replicates reached the maximum amplitude of current density described standard conditions at an applied electrode potential of 310 mV SHE. Seven of the eight biological replicates reached the maximum amplitude of current density within 5 days (Table 1; see also Fig. S1 in the supplemental material). Replicate S2A took 2 days longer. The midpoint potential (E_m) of each BES measured by CV once steady-state current was achieved was similar to that previously reported (9, 10, 13, 19) at ca. 440 mV versus SHE (Table 1). Following CV, the electrode potential was either returned to 310 mV or adjusted to 470 mV, a potential at which we expect the biofilm to recover less energy per electron based on previous experiments. After 52 h, the reactors were disconnected and biofilm samples were scraped off the electrodes and preserved in RNA later within the span of 2 min. RNA was extracted from all eight samples on the same day to reduce batch effects.

MiSeq mRNA sequencing yielded between 4.5 and 16.2 million reads per replicate sample that passed quality control steps, and the data were used for read alignment to the metagenome, of which 37.7 to 90.0% could be unambiguously aligned (Table 1). The proportion of reads mapping to a metagenome bin normalized by the relative length of the bin was used as a proxy for the relative activity of MCL constituents. Using this metric, MCL constituents previously implicated in key roles in the biocathode—“Ca. Tenderia electrophaga,” Marinobacter sp. strain CP1, and Labrenzia sp. strain CP4—were highly active in all samples at both potentials (Fig. 1). Ten other metagenome bins were active in all eight samples using a cutoff of 0.01% of relative activity. Two of these, Parvibaculum sp. (6.9% mean relative activity) and Kordiimonas sp. (3.5% mean relative activity), had activity similar to that of Labrenzia sp. strain CP4 (7.9% mean relative activity) and Marinobacter sp. strain CP1 (4.8% mean relative activity), indicating that they may also have significant roles in the biocathode community. “Ca. Tenderia electrophaga” was the most active constituent in all eight samples, comprising 53% to 84.3% of activity (Fig. 1). Marinobacter sp., Labrenzia sp., Kordiimonas sp., and Parvibaculum sp. averaged 24.6% of the remaining activity. These five constituents make up the
core of the biofilm community, on average accounting for 93.6% of the community activity. Two-tailed Student’s t tests indicated that no constituent had a significant change in activity between the two potentials tested ($P > 0.09$ for all bins).

**Correlation of “Ca. Tenderia electrophaga” transcriptional activity to current density.** Activity of “Ca. Tenderia electrophaga” was strongly correlated to current density at 310 mV (Fig. 2A) with a Pearson correlation coefficient of 0.982 ($P = 0.018$). The correlation coefficient seen at 470 mV was 0.870, which corresponds to a $P$ value of 0.13. The activity of the four most active heterotrophic constituents, *Labrenzia* sp., *Marinobacter* sp., *Parvibaculum* sp., and *Kordiimonas* sp., was negatively correlated or not correlated with current density (Fig. 2B to E). Previous metagenomic and metaproteomic data indicated that “Ca. Tenderia electrophaga” is likely an autotrophic organism capable of performing direct EET (13, 19), and the correlation of current density to activity supports this theory. For this reason, we concentrated our analysis of metatranscriptomics data on “Ca. Tenderia electrophaga” to focus specifically on how direct EET is linked to CO$_2$ fixation.

**Effect of electrode potential on electron transport gene expression.** Efforts to cultivate a representative isolate of “Ca. Tenderia electrophaga” from MCL have thus far been unsuccessful, suggesting that other community constituents satisfy unknown requirements for its growth. We therefore used metatranscriptomics to obtain higher-resolution analysis of differentially expressed mRNA levels for proteins suspected to be involved in electroautotrophic growth. A total of 240 genes in “Ca. Tenderia electrophaga” were significantly more highly expressed at 470 mV, with a false-discovery rate (FDR) of <0.05, while 140 genes were more highly expressed at 310 mV. This allowed us to test the hypothesis that expression of protein-coding genes involved in respiration and energy flux in the primary electroautotroph is directly influenced by the potential of the electrons supplied by the cathode. A complete list of genes that we propose to be associated with these pathways is presented in Table S1 in the supplemental material and discussed below.

Electrochemical measurements indicate that redox-dependent direct electron transfer occurs between the electrode and “Ca. Tenderia electrophaga” (9, 10); therefore, we searched the metatranscriptome for evidence of an EET conduit whose expression might be affected by the change in electrode potential (Table S1). The cyc2 homolog in “Ca. Tenderia electrophaga” (Tel_03480) was predicted to have a role in EET (13) based on its proposed involvement in iron oxidation and was significantly differentially expressed, with a log$_2$-fold change (LogFC) of 0.69 at 470 mV relative to 310 mV. An undecaheme c-cyt (Tel_16545) and a predicted hexaheme lipoprotein (Tel_04230) were previously identified as a possible route(s) for EET in “Ca. Tenderia electrophaga” due to the large number of predicted heme binding sites, known to be important for EET in *Shewanella* and *Geobacter* spp., and to conservation among other EET-capable organisms (13). The genes for these proteins were not differentially expressed.
It is thought that soluble, periplasmic cytochromes mediate ET between the outer membrane and the cytoplasmic membrane-bound ETC (25, 26). Several genes previously noted from “Ca. Tenderia electrophaga” which may encode proteins involved in transferring electrons across the periplasm to ETC were examined for changes in expression between the two electrode potentials (Table S1). A diheme cytochrome $c\_4$, Cyc1, has been proposed to be a periplasmic electron shuttle in the iron-oxidizing bacterium $M. ferrooxydans$ (23). The gene encoding this protein is found in “Ca. Tenderia electrophaga,” in a region displaying synteny with two contigs from the $M. ferrooxydans$ genome (Fig. S2). It is very highly expressed at both potentials (Table S3) but is not significantly differentially expressed. Another potential periplasmic electron carrier identified from the metatranscriptome that was not previously implicated in MCL EET is a monoheme $c\_1$-type cytochrome (Tel_12755). This gene has the largest change in expression of any annotated cytochrome in “Ca. Tenderia electrophaga” (1.62 LogFC more highly expressed at 310 mV), suggesting that it may be important for electron transfer at this lower potential. It is homologous to genes for proteins in several iron-oxidizing bacteria, including Sideroxydans (45% amino acid identity), Gallionella (42% identity), and Acidithiobacillus (42% identity) species. This may make it an important gene to consider in models of iron oxidation.

Expression was higher for three triheme cytochromes (Tel_16515, Tel_16520, and Tel_16530) at 310 mV, but the results were not statistically significant ($P > 0.4$). Overall,

![Graphs showing activity of top five biocathode-MCL constituents versus current density at the optimal and suboptimal electrode potentials. (A) “Ca. Tenderia electrophaga” has positive correlations with increasing negative current at both the optimal (circles) and suboptimal (squares) potentials. (B to E) Other active members of the community, Labrenzia sp. strain CP4 (B), Marinobacter sp. strain CP1 (C), Kordiimonas sp. (D), and Parvibaculum sp. (E) had a negative or insignificant correlation at both potentials. Linear regression lines are shown with $R^2$ values.](http://msystems.asm.org/)
these genes are among those most highly expressed in “Ca. Tenderia electrophaga,” indicating their importance for growth at the cathode. They also appear to be cotranscribed with a gene encoding a tetratricopeptide repeat domain (Tel_16525; Table S3), which is known to be involved in protein:protein interaction and may be involved in aligning cytochromes in a bridge configuration as proposed for other EET-capable bacteria (27). Peptides from this tetratricopeptide repeat protein were previously found to be significantly more abundant at the suboptimal potential (19), although in the metatranscriptomic data, this group of four genes was slightly more highly transcribed at 310 mV.

Regardless of the pathway used, the e− need to reach the cytoplasmic membrane-bound ETC, which is the main mechanism for conservation of energy via chemiosmotic gradient formation (28). Metatranscriptomic analysis of the ETC components does not indicate large changes in the expression of most of these genes, which suggests that the same ETC components are used at both electrode potentials. Small (LogFC < 0.4) changes in the ratios of expression for whole pathways may represent subtle adjustments in the ratio of electrons going down each path (Table S1). Whether this is in addition to or instead of the pathway being selected by the potential of the electron donor in a manner reminiscent of the results of recent work on electron acceptors in Geobacter sulfurreducens remains to be determined (29, 30).

Most genes encoding components of the ETC were not significantly differentially expressed, although many had small but consistent changes in expression between the two potentials. For example, the genes encoding the alternative complex III (ACIII) and the NADH:ubiquinone oxidoreductase complex (NUOR) were not differentially expressed. However, hierarchical clustering based upon the slight changes of expression among these complexes (Fig. 3) suggests that they are functionally linked. This supports the concept of a role as the reverse electron transport (RET) chain.

Only one operon in “Ca. Tenderia electrophaga” contains all four of the canonical genes for a bacterial cytochrome cbb3 oxidase (ccoNOPQ), but it has relatively low expression under both sets of conditions. However, the genome also encodes several
proteins homologous to terminal oxidases (complex IV; Table S1) that could potentially reduce O$_2$ and generate proton motive force (PMF). Of these, the genes encoding complex IV-2 had some of the highest total expression of any genes. Furthermore, hierarchical clustering places this complex in a cluster that contains the ACIII and NUOR complexes, along with genes for three groups of potential electron transport proteins or complexes reported above (Cyc1, an undecaheme cytochrome, and three paralogous triheme cytochromes), thus linking EET to generation of membrane potential and the reduction of NAD(P)$^+$ for CO$_2$ fixation (Fig. 3). The orthologous Cyc1 from _M. ferrooxydans_ is thought to be responsible for transferring electrons from the outer membrane to the cytochrome cbb$_3$ oxidase (23).

Among the genes significantly more highly expressed at 470 mV, hierarchical clustering reveals a similarity in expression patterns between _cyc2_, a gene encoding a protein that has been proposed to play a role in EET in iron-oxidizing bacteria (31), and a gene for hexaheme lipoprotein previously identified as a potential route for EET (Fig. 3; Table S1) (13). They also have expression similar to that of a cytochrome bd oxidase that could accept electrons from the quinone pool and reduce O$_2$ and that is significantly more highly expressed at 470 mV. A possible nitrite reductase complex (Tel_15450 to Tel_15465) is also found within this cluster. However, this complex lacks genes encoding proteins that could function as analogs of the membrane-associated proton-pumping CcoN. Thus, it is unlikely to make a substantial contribution to PMF. Genes in another set that are also within this cluster encode a potentially interesting complex: a cupredoxin domain protein, a hypothetical tetraheme protein, and two hypothetical diheme proteins. This group of genes is labeled as a putative electron transfer complex, and the three heme-binding proteins are differentially expressed.

All of the genes encoding components of the two ATP synthase operons were more highly expressed at 310 mV, and five of them have an FDR of $<0.05$, indicating that the energy levels in "Ca. Tenderia electrophaga" are dependent upon the electrode potential (Fig. 3; Table S1). A gene encoding a bifunctional cytochrome cbb3 oxidase, which is a fusion of the catalytic subunit CcoN and the proton-pumping CcoO, displays a similar pattern of expression, and, via hierarchical clustering, these three groups of genes are clustered with several other genes that may be involved in electron transfer. One set of genes encodes a polyferredoxin and three other hypothetical proteins (Tel_00495 to Tel_00510) that are within the syntenous regions described above (Fig. S2). The three genes encoding hypothetical proteins are significantly more highly expressed at 310 mV. We also found genes in this cluster encoding two monoheme soluble c-type cytochromes, including Tel_12755, which is significantly more highly expressed at 310 mV.

**Effect of potential on carbon fixation and central carbon metabolism.** The "Ca. Tenderia electrophaga" genome contains all the genes necessary for CO$_2$ fixation via the CBB cycle, including two forms of RuBisCO—one associated with carboxysomes (form IAc) and one not associated with carboxysomes (form IAq) (13). The lower energy yield per electron from a cathode poised at 470 mV would result in less energy available for CO$_2$ fixation, so the central carbon metabolism of "Ca. Tenderia electrophaga" was examined for changes in expression (Fig. 4; Table S3). Genes which could be involved in two or more pathways, for example, the CO$_2$ fixation pathway and the pentose phosphate pathway, were separated where possible by examining colocalization and changes in expression among the eight samples. Genes with ambiguous functional roles that could not be assigned to a single pathway were excluded from this analysis. Most genes encoding enzymes of the central carbon metabolism, including the CBB cycle, were more highly expressed at 310 mV, although the absolute magnitude of the difference was relatively low (Fig. 4; Table S3). The small differences in expression of individual genes for the central carbon metabolism were not statistically significant; however, in examining all genes in a given pathway, they were mostly consistent, indicating that the pathways are likely genuinely more highly expressed at 310 mV. The exceptions are the form IAq RuBisCO genes and the carboxysome genes, which were
more highly expressed at 470 mV. This suggests a switch to a more efficient enzyme under energy limitation conditions.

**DISCUSSION**

Our characterization of the biocathode-MCL community activity by metatranscriptomics supports previous metaproteomic and 16S rRNA amplicon characterizations of biocathode-MCL indicating that there is substantial variability in the abundance of specific bacteria, even between seemingly identical reactors inoculated on the same day with the same inoculum (10, 19). However, the major constituents that are present have been stable for over 6 years, as measured by several methods, including the use of 16S clone libraries, metagenomics, proteomics, 16S amplicon sequencing, and fluorescence *in situ* hybridization microscopy (9, 10, 13, 19, 20), allowing us to use the previously generated metagenome sequencing data to align the metatranscriptomics data. This is important for studies where only limited biomass can be recovered from the electrode surface. As in the previous studies, which examined the abundance of biofilm constituents, changing the electrode potential from 310 mV to 470 mV for 52 h after the community had developed did not result in significant changes in the average relative levels of activity of biocathode constituents between the two potentials. While the inherent variability in relative abundance and activity between replicate reactors makes analysis of the transcriptome data challenging, the differences that we found are more likely to be robust because they are visible above the noise of such large biological variability found by using paired, replicated samples as described by Leary et al. (19).

The magnitude of changes in gene expression was relatively small compared to that seen in other transcriptomic studies (see, e.g., references 32 and 33). However, in one of those studies, replicate samples were acquired by repeatedly sampling the same biofilm, which may have inflated the reported statistical confidence, and in the other, no replicates were sampled. The 52-h adjustment period after changing the potential likely decreased the magnitude of the transcriptional response, but it was chosen to facilitate comparison to proteomics data (19). On a shorter time scale, transcription of genes for upregulated pathways might experience a burst, as the cells adjust to the new regulatory regimen, but would relax back to the new steady state once the proper balance of proteins has been reached (see, e.g., reference 34). This also allowed us to be certain that there were no lingering effects from running CV and that any changes that were detected in the transcriptome represented a steady-state adjustment and not a transient spike due to short-term stress. Also, the fundamental metabolism of ‘Ca.

**FIG 4** Change in expression of central carbon metabolism genes. Open boxes indicate genes that are more highly expressed at 470 mV, and gray boxes indicate genes that are more highly expressed at 310 mV. Error bars indicate standard deviations of the LogFC of pathways containing three or more genes. Details of the genes included in each category and of their expression can be found in Table S2. Alt. Comp., alternative complex; Cyt., cytochrome; Put., putative; TCA cycle, tricarboxylic acid cycle.
Tenderia electrophaga” does not change. Electrons are still being transferred from the cathode to the ETC, and CO2 fixation is still the sole carbon source.

Our results suggest that “Ca. Tenderia electrophaga” is primarily responsible for EET linked to CO2 fixation in biocathode-MCL biofilms. The dominance of “Ca. Tenderia electrophaga” transcriptions activity and its correlation to current density suggest that it can account for the majority of the current. Further substantiating this hypothesis are two recent reports of unclassified bacteria, thought to be within the Chromatiales, which were dominant members of freshwater biocathode communities (17, 18). While Marinobacter spp. are capable of cathode oxidation (21, 22, 35), isolates of Marinobacter sp. and Labrenzia sp. from the MCL community are capable of only weak current production at the biocathode or iron oxidation in monoculture (13).

On the basis of a prior analysis of biocathode-MCL performed using slow-scan-rate CV (19), we predicted the current to be approximately halved when the electrode potential was switched from 310 mV to 470 mV. However, as previously observed (19), for biocathode-MCL, switching to and maintaining a more positive potential (470 mV) over a much longer time period (>50 h) resulted in an increase in current attributed to O2 reduction after the initial adjustment period. This increase in the magnitude of the current is interpreted as the result of the need to make up for the decrease in energy available per electron at the higher potential. Employing a derivation of the Nernst equation (\( \Delta G = -nF\Delta E^0 \)) to estimate the theoretical yield of O2 reduction to H2O using an electron donor at a potential of 310 mV yields −47 kJ/mol e\(^{-}\), indicating the requirement of an additional ~61 kJ/mol e\(^{-}\) to reduce NAD(P)\(^{+}\). Assuming that the energy required to pump protons across the membrane is ~21 kJ/mol, “Ca. Tenderia electrophaga” could export two H\(^{+}\) per e\(^{-}\) at the optimal potential and would need to use three H\(^{+}\) to reduce NAD(P)\(^{+}\). This results in a theoretical balance for electron utilization by the forward versus reverse electron transport pathways of ~60%/40% to produce NAD(P)H. More electrons would need to go to the forward path to generate a proton gradient for ATP production. In comparison, using an electron donor at a potential of 470 mV reduces the \( \Delta G \) to −32 kJ/mol e\(^{-}\), lowering the theoretical yield to ~1.5 H\(^{+}\) exported per e\(^{-}\), and the larger \( \Delta E^0 \) between the electrode and NAD(P)H requires at least four H\(^{+}\) translocated per NAD(P)\(^{+}\) reduced. This results in an electron utilization balance of at least 80% and 20% for the forward and reverse pathways, respectively. Thus, theoretically, at least twice as many electrons are needed to generate the same amount of reducing equivalents for CO2 fixation at the more positive potential. During growth under standard culture conditions, the acidophilic iron oxidizer A. ferrooxidans is predicted to have a ratio closer to 90/10% for the forward versus reverse pathways, including the proton gradient necessary to generate ATP (36). Recent experimental evidence suggests that the ratio of electron utilization by the forward and reverse pathways during growth on an electrode is close to 15:1 in A. ferrooxidans (15).

Redundancy of electron transport chain components suggests metabolic flexibility in “Ca. Tenderia electrophaga” dependent upon the potentials of electron donors and the local redox environment (Fig. 5). Thermodynamics dictates that the balance of electrons passing through the two branches of the ETC depends upon the electrode potential. One possible method of controlling this ratio would be modulating the abundance of the periplasmic links between EET and the ETC. Several proteins could potentially make up this link, but their roles are presently unclear. One highly differentially expressed gene (Tel_12755) encodes a soluble monoheme cyt-c that could fulfill this role. Hierarchical clustering of potential electron transfer components by expression led to identification of several distinct clusters (Fig. 3). One cluster contains the ACIII and NUOR genes, which suggests that these complexes may be functionally linked. The canonical cytochrome cbb3 oxidase is part of a cluster with succinate dehydrogenase (complex II) and the cytochrome bc1 complex. This may indicate that they form part of a forward ETC, which would enable the use of stored reserves in the form of glycogen.

Among the proteins possibly involved in ET at the outer membrane, the Cyc2 homolog encoded by Tel_03480 was more highly expressed at 470 mV, consistent with...
The lower energetic yield per electron. This protein has previously been implicated in EET in the acidophilic iron-oxidizing bacterium *A. ferrooxidans* ATCC 19859 (31, 37, 38). The $E_m$ of Cyc2 in *A. ferrooxidans* is reported to be 560 mV at pH 4.8 (27). This is within the range that might be expected for a protein accepting electrons from a cathode at the potentials tested, although the midpoint potential of Cyc2 in *Ca. Tenderia electrophaga* is likely different. This strongly suggests that Cyc2 is involved in EET with the electrode in *Ca. Tenderia electrophaga,* although its exact role given the single heme and lack of transmembrane helices remains unknown.

The majority of CBB cycle genes and pentose phosphate cycle genes appear to be more highly expressed at 310 mV. This likely reflects a higher rate of CO$_2$ fixation at 310 mV than at 470 mV. Furthermore, the higher relative expression of the form IA$\nu$ RuBisCO genes and their associated carboxysomes at 470 mV may reflect the need to increase the efficiency of CO$_2$ fixation due to the lower energy availability. However, even at 470 mV, the overall transcript abundance of form IA$\alpha$ is still twice as high as that of form IA$\nu$. Metatranscriptomics of biocathode MCL supports previous work by our group that identified "Ca. Tenderia electrophaga" as the primary electroautotroph. It appears to be the key member of the community, and the notion of its predicted lifestyle as an electroautotroph is supported by several key findings. "Ca. Tenderia electrophaga" was more active than any other constituent, and its activity was positively correlated with current density. Genes for proteins previously predicted to be involved in EET in "Ca. Tenderia electrophaga," including Cyc2, were more highly expressed at 470 mV, when more electrons are needed to generate the same amount of PMF. The membrane-bound ETC that is necessary to generate the ATP and reducing power for CO$_2$ fixation was active at both potentials, but a potential complex IV analog and several that may be involved in the "downhill" branch of the ETC showed increased expression in response to changes in electrode potential. These changes were consistent with the need to use more electrons to obtain the same amount of energy. Most pathways of central carbon metabolism were more highly expressed at 310 mV, though not at the individual gene level, suggesting a higher metabolic rate. The evidence for
involvement of specific proteins or complexes in EET and ETC is correlative in nature but reduces the list of potential targets for the future confirmatory studies that are in progress. Further investigations with better temporal resolution may yield additional insights into the dynamics of the response to the changes in potential. Heterologous expression and biochemical measurements of predicted components of EET and the ETC may help identify functional roles. This work provides insight into the mechanisms for electroautotrophic growth in this biocathode community that will assist in engineering biocathode communities.

MATERIALS AND METHODS

Biofilm growth and sampling. Samples for RNA-seq were grown in artificial seawater medium (ASW) as previously described (13). Inocula consisted of cells detached by vortex mixing and sonication from a 3-cm-by-3-cm portion of a carbon cloth electrode and were counted using an Accuri C6 flow cytometer (BD Bioscience, San Jose, CA) as described previously (19). Sterile 2-liter dual-chambered microbial fuel cell reactors (Adams and Chittenden Scientific Glass) containing a graphite coupon (3 cm by 9 cm by 0.2 cm) were inoculated with standardized inocula estimated to contain 2 × 10⁸ cells.

Four reactors were inoculated with the same inoculum, and cells were grown under standard conditions (310 mV SHE) until the maximum current (12 to 65 mA·m⁻²) was reached. At this point, CV was recorded from 610 mV to 260 mV and back to 610 mV at a scan rate of 0.2 mV/s. After CV, two reactors remained poised at 310 mV, and two were shifted to a suboptimal potential (470 mV SHE) where CV indicates that the rate of electron uptake is approximately half of that at the “optimal” potential. After 52 h, the biofilm was scraped from both sides of the cathode with a sterile razor blade, immediately immersed in RNA-later (Life Technologies, Grand Island, NY), and stored at 4°C overnight prior to freezing at −20°C until total RNA extraction was performed. This experiment was repeated 7 days later with a new inoculum. This gave two pairs of biological replicates for each treatment and for each inoculum, which helps to control for variability caused by small differences in the community present in each inoculum.

RNA extraction, library preparation, and sequencing. Biofilm RNA was extracted using an established protocol (39) with the following modification: instead of 0.8 g of 0.5-mm-diameter glass beads, 250 µl of zirconia beads (Life Technologies, Inc.) was used. The extracted RNA samples were subjected to DNase treatment using Turbo DNase (Life Technologies, Inc.) and were cleaned up using RNA Clean & Concentrator-5 (Zymo Research). The DNase treatment was repeated once for each sample to ensure the removal of contaminating DNA. Ribosomal RNA was depleted from RNA samples using a Ribo-Zero rRNA Removal kit for Bacteria (Illumina, San Diego, CA). RNA library preparation was carried out using a NEBNext Ultra RNA Library Prep kit for Illumina (New England Biolabs, Ipswich, MA) according to the manufacturer’s recommended protocol. The sequence reads of RNA libraries were acquired on a MiSeq instrument after automated software control (version 2.2.0; Illumina, San Diego, CA). Raw reads that passed the initial quality filter step were submitted to the NCBI Sequence Read Archive (http://www.ncbi.nlm.nih.gov/bioproject).

Metatranscriptome alignment and statistical analysis of gene expression. Metatranscriptome reads were trimmed for quality using Trimmomatic 0.30 with a sliding window set to 5 bases with a minimum score of 25, and the first 8 bases of the sequence and any leading bases with a quality score less than 20 were cropped (40). Individual reads with less than 50 bases remaining were discarded. A biocathode-MCL metagenomic assembly, available at http://www.ncbi.nlm.nih.gov (A. P. Malanoski, B. Lin, Z. Wang, B. J. Eddie, W. J. Hervey IV, and S. M. Strycharz-Glaven, unpublished data), was modified by removing contigs from the bins corresponding to PacBio sequenced genomes of “Ca. Tenderia electrophaga,” Lobrenzie sp. CP4, and Marinobacter sp. CP1 (CP013099.1, CP013100.1, CP011929.1, CP011927.1, and CP011928.1) and contigs that were >98% identical to these genomes over >90% of their length. The modified metagenome sequence is available at https://figshare.com/articles/Biocathode_MCL_Curated_Metagenome/4282631. Bowtie 2 (41) was used to align trimmed paired-end reads and remaining singleton reads to the modified metagenome using the following options: -D 30 -R 3 -N 1 -L 15-1,5,0.50—score-min L0,—0.15.

The resulting SAM files were converted to sorted BAM files with SAMtools version 1.1, and read counts per contig were calculated using the iStatsd function (42). The relative activity of each genome or metagenomic bin was calculated as the sum of RNA-seq reads mapping to the genome or bin divided by the length of the genome or bin in nucleotides as a percentage of activity in the library. Read counts per gene were calculated with the featureCount tool in the subread package (43). EdgeR (44, 45) was used to normalize read counts by the trimmed mean of M-values method and to determine the statistical significance of differences in read counts for each annotated gene and pseudogene in “Ca. T. electrophaga” using the likelihood-ratio test. Benjamini-Hochberg adjustment was used to reduce the rate of false discovery due to multiple-hypothesis testing (46). A multifactor experimental design was used to separate changes in expression likely to be due to the use of the different inocula from changes in expression resulting from the change in electrode potential (47). Genes with an FDR of <0.05 for differential expression were considered differentially expressed. Values reported for gene expression under each set of conditions correspond to the log₂ value of the number of reads per gene after normalization (48) (see Table S3 in the supplemental material). To facilitate analysis of differences between genes in relative expression levels, overall gene expression was calculated as log counts per minute (LogCPM) minus the log₂ of the length of the gene divided by 1,000, roughly equivalent to the commonly used parameter of reads per kilobase per million (49). For clustering genes by change in
expression, gene expression was centered at 0 for each inoculum. Euclidean distance matrices were calculated, and genes were clustered using the Ward method implemented in R package “pheatmap” version 0.2.7 (50).

Correlations between current density and the relative abundance of mRNA from each metagenomic genome or bin were calculated as Pearson’s product-moment correlation coefficients (51). The critical value for a significant correlation for a two-tailed test with a sample size of \( n = 4 \) is 0.9 for a \( P \) value of 0.1. A higher \( P \) value cutoff for the correlation is justified because this is a different kind of statistical test and it assumes a linear relationship between the data, which is likely not the case for most genes. Furthermore, there was a large variability between samples collected under the same conditions (current magnitude, increasing/decreasing current, and community composition) that would prevent the nearly perfect correlation that a \( P \) value of 0.05 would require.

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