

A Previously Uncharacterized, Nonphotosynthetic Member of the *Chromatiaceae* Is the Primary CO₂-Fixing Constituent in a Self-Regenerating Biocathode

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Biocathode extracellular electron transfer (EET) may be exploited for biotechnology applications, including microbially mediated O₂ reduction in microbial fuel cells and microbial electrosynthesis. However, biocathode mechanistic studies needed to improve or engineer functionality have been limited to a few select species that form sparse, homogeneous biofilms characterized by little or no growth. Attempts to cultivate isolates from biocathode environmental enrichments often fail due to a lack of some advantage provided by life in a consortium, highlighting the need to study and understand biocathode consortia in situ. Here, we present metagenomic and metaproteomic characterization of a previously described biocathode biofilm (+310 mV versus a standard hydrogen electrode [SHE]) enriched from seawater, reducing O₂, and presumably fixing CO₂ for biomass generation. Metagenomics identified 16 distinct cluster genomes, 15 of which could be assigned at the family or genus level and whose abundance was roughly divided between Alpha- and Gammaproteobacteria. A total of 644 proteins were identified from shotgun metaproteomics and have been deposited in the the ProteomeXchange with identifier PXD001045. Cluster genomes were used to assign the taxonomic identities of 599 proteins, with Marinobacter, Chromatiaceae, and Labrenzia the most represented. RubisCO and phosphoribulokinase, along with 9 other Calvin-Benson-Bassham cycle proteins, were identified from Chromatiaceae. In addition, proteins similar to those predicted for iron oxidation pathways of known iron-oxidizing bacteria were observed for Chromatiaceae. These findings represent the first description of putative EET and CO₂ fixation mechanisms for a selfregenerating, self-sustaining multispecies biocathode, providing potential targets for functional engineering, as well as new insights into biocathode EET pathways using proteomics.

ioelectrochemical systems (BES) use microorganisms as cata-Blysts to drive complex electrochemical reactions, such as electricity generation by microbial fuel cells (MFCs) (1), wastewater treatment (2), and microbial electrosynthesis (3-6), that would not be possible without living cells. The term "biocathode" refers to a biofilm, constituted of a single organism or microbial consortium, that has formed on the cathode of a BES and consumes electrons (e⁻). Cathodes hold great potential as a stable electron source to drive microbial metabolism (7); however, little is known about the underlying extracellular electron transfer (EET) pathways that could be exploited for biocathode functional engineering. Although biocathode EET has been demonstrated for a variety of microorganisms, including acetogens (5) and a methanogenic archaeon (6), studies aimed at identifying EET conduits from the electrode to cells have mostly been confined to the model organisms Geobacter (8) and Shewanella (9), due to the massive effort put forth to understand how these iron-reducing bacteria are able to catalyze EET at bioanodes (10-12). The ability of iron-reducing bacteria to reduce anodes led to the hypothesis that iron-oxidizing bacteria (FeOB) would be able to perform EET with cathodes, which has been demonstrated for at least two FeOB, Mariprofundus ferrooxydans and Rhodopseudomonas palustris (13, 14). While many bacteria have been shown to attach to electrodes and "consume" electrons, electrode-dependent growth has thus far been demonstrated only for *M. ferrooxydans* (13). Furthermore, most cathode EET processes studied to date rely on a fairly negative electrode potential (between 0 and -0.400 V standard hydrogen

electrode [SHE]) to catalyze CO_2 or O_2 reduction. Biocathodes developed at higher potentials, such as those used in this study, need to rely on EET mediators with much higher potentials (>+0.300 V SHE). Identifying and understanding such mediators could provide flexibility to engineering functionality in BES applications where higher operating potentials are desired.

A challenge in developing the use of BES under environmentally relevant conditions (i.e., in seawater and under changing pH and changing temperature) is a lack of understanding of cathodic microbial communities. Little effort has been put into developing microbial consortia as biocathode catalysts, even though they have been shown to outperform homogeneous bacterial populations in

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terms of current density (15). Marshall et al. (3, 16) have demonstrated long-term biocommodity production using an acetogenic biocathode consortium but have not yet reported on the underlying EET pathways. Attempts to cultivate isolates from biocathode environmental enrichments typically result in loss of the electrochemical phenotype (4), as individual biofilm constituents may lack some essential cofactor provided by life in a consortium, such as vitamins or amino acids.

Advances in systems biology tools and large-scale, culture-independent, community level "omic" (e.g., metagenomic, metatranscriptomic, and metaproteomic) measurements now allow us to predict and potentially direct interactions in naturally occurring, stable microbial consortia (17). Metaomics data provide a perspective on the physiological state of organisms thriving from their associations with one another that may be different than when they are grown in homogeneous microbial populations (18). When applied to study biocathode microbial consortia, combined metagenomic and metaproteomic analyses may be particularly useful for generating information about the functions of biofilm constituents in relation to the biocathode lifestyle (19, 20). Biocathode metaproteomics can provide functional information from electrode-grown cells in order to predict biofilm EET pathways for targeted biofilm engineering, particularly when key EET biofilm constituents cannot be cultivated or when ex situ cultivation conditions are not relevant to electrode growth. In relation to BES, the use of metaproteomics to study bioanodes has been limited (21), and there are currently no published reports on the proteome of a biocathode.

We previously reported on the electrochemical features of an aerobically grown, nonphototrophic biocathode community enriched from seawater (22). This biocathode is a durable, multicelllayer-thick biofilm that is self-regenerating and self-sustaining. We hypothesized that the biocathode uses electrons supplied by a poised electrode (+310 mV versus SHE) to drive CO₂ fixation and O2 reduction, since no other electron donor or carbon source is provided. Portions of the biocathode biofilm can be removed and used to inoculate subsequent biocathode reactors that achieve reproducible electrochemical characteristics of the parent biofilm. 16S rRNA gene clone libraries initially showed the biocathode biofilm to be a low-complexity consortium that consisted primarily of Marinobacter, a ubiquitous biofilm-forming member of the Gammaproteobacteria known to oxidize iron under aerobic and circumneutral conditions (23), as well as other bacteria most closely related to marine Alpha- and Gammaproteobacteria.

The primary objective of the current study was to obtain an initial understanding of in situ biocathode EET and carbon fixation pathways at maximum current. We expand upon our previous work using metagenomics, reverse transcription (RT)-PCR, and shotgun metaproteomics to (i) confirm the identities of the primary biofilm constituents, (ii) provide an initial survey of the biocathode metaproteome, (iii) identify proteins of putative EET pathways, and (iv) identify carbon fixation pathways that may confer autotrophy on the biocathode community using electrons from the electrode as an energy source. We show that an unknown member of the family Chromatiaceae expresses proteins for both CO₂ fixation and EET. While Marinobacter may have some capacity for EET, no known EET pathways were identified. Roles for other abundant biofilm constituents, including Labrenzia and Kordiimonas, could not readily be predicted but are likely important for biofilm formation and carbon cycling.

MATERIALS AND METHODS

Biocathode biofilm cultivation. The bioelectrochemical reactors were 2-liter dual-chambered microbial fuel cell reactors (Adams and Chittenden Scientific Glass) without membrane separation. The working electrodes were either graphite coupons (length, 3.0 cm; height, 10.0 cm; width, 0.2 cm; total geometric surface area, 65.2 cm², or 0.00652 m²) or carbon cloth flags (length, 3.5 cm; height, 3.5 cm; total geometric surface area, 24.5 cm², or 0.00245 m²). Following initial proteomics analysis from graphite coupons, carbon cloth electrodes were used to grow biocathodes for protein analysis to improve protein recovery, since their electrochemical features were identical. Further descriptions of reactor and electrochemical measurements can be found in the supplemental material. The reactors were filled with artificial seawater medium (ASW) (see the supplemental material for the composition), and scrapings from previously described enriched biocathode biofilms were used as a source of inoculum (22). The reactors were maintained at 30°C with stirring (VWR standard multiposition stirrer; setting 2 [150 to 200 rpm]) under atmospheric conditions. The working electrodes were maintained at approximately +0.310 V versus SHE (+0.100 V versus Ag/AgCl) using a multichannel potentiostat (Solartron 1470E) under software control (Multistat; Scribner). All the potentials reported here are versus SHE unless specifically noted otherwise. Once maximum current was reached, biocathode electrochemical features were characterized and confirmed to be consistent with those previously reported (see Fig. S1 and S2 in the supplemental material) (22).

Metagenomic DNA sequencing and assembly. Metagenomic DNA was extracted from a graphite coupon electrode grown as described above using the MoBio PowerBiofilm DNA isolation kit. DNA integrity and concentration were verified using a Bioanalyzer 2100 (Agilent, Palo Alto, CA). Approximately 1.0 µg of high-quality DNA was processed using an Illumina TruSeq DNA sample preparation kit following the manufacturer's instructions (Illumina, San Diego, CA, USA). The library was validated and sequenced on an Illumina HiSeq 2000 sequencer using 100-bp paired-end reads. Approximately 31.3 million filtered raw read pairs were generated for this study. Raw sequence reads from the whole-genome shotgun (WGS) approach were first trimmed using SolexaQA (24), a Perlbased software package calculating quality statistics from FASTQ files generated by Illumina sequencers. The default setting (P = 0.05) was used. Additional quality control steps included the removal of low-quality reads based on error probabilities and trimming low-quality tails. After these steps, only paired-end reads of >90 bp were retained. These processed reads were then assembled using Velvet v1.2.10 (25) (kmer length = 55; insertion library length = 400). The expected coverage was set to auto.

ORF calling and annotation. Metagene (26) was used to predict 79,765 open reading frames (ORFs) from the biocathode metagenome, of which 63,097, 86,537, and 77,136 ORFs were assigned COG, pfam, and KEGG identifiers (IDs), respectively (see Table S3 in the supplemental material). Annotations were assigned using RPS-BLAST and the NCBI Conserved Domain Database (27). The predicted ORFs were submitted to WebMGA (28) to generate KEGG identifiers (29). Contigs from clusters containing fewer than 500 contigs (see below) were submitted as individual groups to the RAST pipeline for annotation (http://rast.nmpdr.org). Amino acid sequences for known proteins of functional interest were used to identify homologous proteins from the translated metagenome using the BLASTp algorithm and the NCBInr protein database with default settings (http://blast.ncbi.nlm.nih.gov).

Taxonomic/phylogenetic analysis. Predicted ORFs were processed using AMPHORA2 to identify and assign taxonomy to 31 different housekeeping genes (30). Single-copy housekeeping genes were used to infer the relative abundances of the biofilm constituents based on the Velvet assembly coverage of each unique gene. Since ORF calling on contigs could potentially identify paralogs or partial fragments, results returned from AMPHORA2 were further curated manually (see the supplemental material for details). During manual curation, ORFs identified by AMPHORA2 as containing a particular housekeeping gene from the same species, and

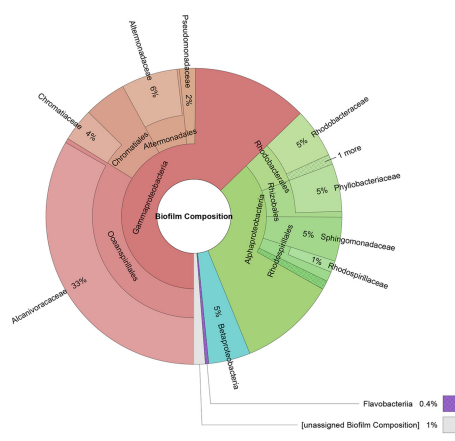


FIG 1 Relative abundances (weighted by sequence coverage) and taxonomic distribution of the biocathode community based on the most likely phylogenetic assignment of 31 distinct housekeeping genes identified from assembled metagenomic contigs. Class, order, and family level identifications with at least 1% relative abundance are shown (except for *Flavobacteriia*, which is shown at 0.6% due to a high-confidence identification).

from contigs with similar coverage, were grouped together. Relative abundance was estimated for each housekeeping gene by summing the Velvet assembly coverage of the contigs from which the genes were identified. These individual gene coverage results were averaged across a given taxonomic assignment and are reported in Fig. 1. Additionally, filtered raw reads were analyzed using MetaPhyler v1.25 to generate an abundance count at various taxonomic levels using taxonomic marker genes from complete genomes (31).

Clustering of assembled contigs. Clusters were defined for groups of AMPHORA2 genes that resolved to the same taxonomic level and for which similar coverage was found for the contigs from which the ORFs were generated. Contigs longer than 200 bp were processed using the Metawatt binner (v1.7), which uses multivariate statistics of tetranucleotide frequencies combined with use of the interpolated Markov model (IMM) (GLIMMER 3.02) to cluster contigs (32). Initial clusters based only on tetranucleotide frequencies were built of at least 0.1 Mbp using the medium confidence level. The resulting clusters were merged after inspection if contigs contained therein were previously identified as being in the same cluster according to AMPHORA2 analysis. The IMM modeling step was then run to obtain a refined group of clusters. The average and standard deviation of coverage of contigs longer than 2 kb in each cluster were computed. In clusters where the standard deviation was more than 10% of the average, the large contigs (>5,000 bp) were compared to this average, and if the coverage deviated significantly from the average, it was subjected to further checks. ORFs from contigs identified as being suspect were searched against the NCBInr database using BLASTp. Consensus identification was attempted for each contig by counting the top genus hit of each ORF from that contig. If the closest match for more than 37% of predicted proteins on a contig were assigned to the same genus, the contig

was designated as belonging to that genus. Contigs that had 25 to 37% proteins with their closest matches within a single genus were submitted for discontiguous megablast. If a large fraction of the contig was assigned to an organism in the same genus as those with >37% call identity, it was assigned that designation. For purposes of assigning functional proteins to a specific cluster genome, all unclassified clusters were treated as a single cluster genome.

Housekeeping genes. The KEGG orthology (KO) functional annotation has been determined for 107 marker genes that are typically found to be in single copies in more than 95% of bacterial genomes (33, 34). The number of these marker genes in each Metawatt cluster was determined from the KO assignments made in the KEGG annotation of the metagenome contigs.

Multiheme *c*-type cytochrome prediction. Multiheme *c*-type cytochromes (*c*-Cyts) were predicted using hmmsearch in the HMMER software package v3.1b1 (E value < 1E-5) (35). *In silico*-translated protein sequences derived from the metagenome were searched using a doubled CXXCH motif (pfam09699), which represents two copies of the hemebinding CXXCH motif. Predicted multiheme *c*-Cyts were screened for lipoprotein domains using the LipoP 1.0 server (36). The average molecular weight (MW) was predicted using the MoreFASTA utility of the DTASelect program (37).

Metaproteomics. Proteins were extracted from graphite coupon and cloth biofilms using several extraction methods to help alleviate some of the issues surrounding protein extraction bias (38). A low protein yield per electrode prevented simultaneous sampling for both DNA and protein; therefore, separate samples were grown for proteomics and evaluated for electrochemical traits identical to those of reactors used for DNA extraction (see Table S1 in the supplemental material). The methods of

protein extraction and analysis were adapted from previously developed methods (38, 39) and are briefly described below (a complete description can be found in the supplemental material). For graphite block biofilms, biofilms were scraped with razor blades from graphite blocks into 2% sodium dodecyl sulfate (SDS) in 50 mM ABC (ammonium bicarbonate) or B-Per Tris solution (Thermo Scientific, Rockford, IL). All samples were sonicated on ice, mixed with Tris-buffered phenol (1:1 ratio), incubated for 30 min at room temperature with mixing, and then subjected to centrifugation for phase separation. The phenol phase was collected from all samples and combined with ice-cold 100 mM ammonium acetate in 100% methanol (1:4 ratio) prior to incubation at -80°C for a minimum of 16 h. For graphite cloth biofilms, biofilm samples from graphite cloth were submerged in B-Per Tris solution (Life Technologies), sonicated on ice, and centrifuged (4°C; 5,000 \times g; 10 min) to sediment small graphite particles. The cleared supernatants were transferred into new tubes and precipitated using 100 mM ammonium acetate in 100% methanol as described above. The extracted and precipitated proteins were collected by centrifugation, dissolved in SDS-PAGE running buffer, separated by SDS-PAGE, and stained using Coomassie blue. Distinct protein bands were cut, washed, and destained. Each band was then in-gel digested overnight using modified porcine trypsin, and the tryptic digests were collected into new tubes, dried via speed-vac, and stored at -20°C. The dried digests were reconstituted in 5% acetonitrile, 0.1% formic acid in water immediately prior to analysis by reverse-phase liquid chromatography-tandem mass spectrometry (LC-MS-MS) using a TempoMDLC system coupled to a QStar Elite mass analyzer. Tandem mass spectra were extracted using the AB Sciex MS Data Converter (version 2.0) and searched by Mascot (Matrix Science, London, United Kingdom; version 2.4.1) and X! Tandem (The Global Proteome Machine [http://thegpm.org]; version CYCLONE [2010.12.01.1]). Scaffold (version Scaffold_4.2.1; Proteome Software Inc., Portland, OR) was used to validate MS-MS-based peptide and protein identifications.

All identified proteins were assigned to the cluster genomes described above. In order to predict protein functions and metabolic pathways, amino acid sequences were annotated using the BLASTp algorithm and the NCBInr protein database with default settings (http://blast.ncbi.nlm .nih.gov). An annotation of the first highest-scoring protein was accepted if the E value was <5. For hypothetical proteins, conserved domains were used to predict function if they were present. Protein localization was predicted by PSORTb (v3.0.2 [http://www.psort.org/psortb/]). All annotated proteins were manually assigned to three functional categories: (i) enzymes (all annotated proteins for which an EC number could be obtained or a domain known to be in enzymes was present; cytochromes, peroxiredoxins, thioredoxin, molybdopterin, and ferredoxins were included in this category, and the enzyme database BRENDA [http://www .brenda-enzymes.org/index.php] was used to obtain EC numbers and KEGG pathway numbers for all identified enzymes); (ii) receptors, transporters, and membrane proteins (which were grouped according to their substrates/cargo); and (iii) structural proteins and proteins with unknown function. All information regarding protein annotation, predicted function, and predicted localization can be found in Data Set S2 in the supplemental material.

RT-PCR. Biofilm RNA was extracted using a previously described method (40) with the following modification: 250 μ l of Zirconia beads (Life Technologies) was used instead of 0.8 g of 0.5-mm glass beads. The extracted RNA samples were subjected to Turbo DNase (Life Technologies) treatment according to the manufacturer's recommended protocol and purified using the RNA Clean and Concentrator-5 kit (Zymo Research). The DNase treatment was repeated once for each sample to ensure the removal of potential contaminating metagenomic DNA. The Complete Whole Transcriptome Amplification kit (Sigma-Aldrich, St. Louis, MO) was used to amplify biofilm total RNA according to the manufacturer's recommended protocol. The amplified products were purified using the DNA Clean and Concentrator-5 kit (Zymo Research) and quantified using a NanoDrop 2000c UV-Vis Spectrophotometer (Thermo Sci-

entific, Waltham, MA). The quantitative PCRs (qPCRs) were carried out in 25-µl reaction volumes containing $1 \times iQ$ SYBR Green Supermix (Bio-Rad Laboratories, Inc., Hercules, CA), 200 nM (each) forward and reverse primers, and 25 ng of amplified products using a MyiQ single-color real-time PCR detection system (Bio-Rad Laboratories, Inc.). Reactions were performed with an initial denaturation at 95°C for 3 min, followed by 35 cycles of 95°C for 10 s, 56°C for 20 s, and 72°C for 20 s. The sequences of the primers used in this study are listed in Table S2 in the supplemental material.

Accession numbers. The mass spectrometry proteomics data (raw sequence reads) have been deposited in the ProteomeXchange Consortium (41) via the PRIDE partnership repository with the data set identifier PXD001045. The Sequence Read Archive (SRA) accession (SRX621521) of the raw Illumina reads can be found at http://www.ncbi.nlm.nih.gov/sra/SRX621521. Other proteins (see Tables 4 and 5) were also deposited in the PRIDE partnership repository under the same identifier.

RESULTS

Biocathode metagenome. To further characterize the biological composition and functional potential of the community while simultaneously providing a matched-sample database for subsequent metaproteomic studies, a representative biocathode was subjected to whole-metagenome sequencing. Biocathode metagenome sequencing resulted in approximately 31.3 million filtered raw read pairs that were assembled into 32,870 contigs using the Velvet assembler (mean contig length = 2,016 bp; maximum contig length = 627,596 bp; N50 = 55,338 bp) (see Table S3 in the supplemental material). N50 is defined as the length (N) for which 50% of all bases are represented in fragments of length L (<N). Multiple data analysis techniques focusing on single- to low-copynumber housekeeping genes were used to determine the taxonomic composition of the biofilm and to estimate the relative abundances of constituents. A summary of the results from all the techniques used can be found in Table S4 in the supplemental material. First, AMPHORA2 was used to determine the average biofilm composition by examining the taxonomic assignment of 31 housekeeping genes from the assembled contigs (Fig. 1). Fifteen distinct cluster genomes could be identified by AMPHORA2 from 316 contigs that contained at least one of the housekeeping genes. Taxonomic identifications were found to be reliable down to the family level, with 99% of the housekeeping genes identified to the class level. A direct taxonomic analysis performed on the filtered but unassembled reads using MetaPhyler was consistent with those derived from the AMPHORA2 analysis (see Fig. S3 in the supplemental material). In both instances, sequencing fragments that could not be sorted into distinct cluster genomes were taxonomically classified as Betaproteobacteria or Flavobacteriia. These fragments may represent biocathode constituents present at very low abundance that could not be further resolved with the analysis techniques used here. Second, an additional clustering approach based on tetranucleotide frequency was performed on all contigs that were >200 bp in size using Metawatt (Fig. 2). Metawatt analysis resulted in 23 refined cluster genomes, 16 of which contained at least 1 Mb of sequence. All except the Unclassified-1 cluster were considered to be representative of a single organism. The RAST pipeline provided further confirmation of taxonomic assignments of refined cluster genomes with fewer than 500 contigs. Overall, the combined RAST, AMPHORA2, and Metawatt analyses agreed on all but one identification and even enabled genus level identification of some cluster genomes. The sole discrepancy was the RAST-identified Kordiimonas cluster genome,

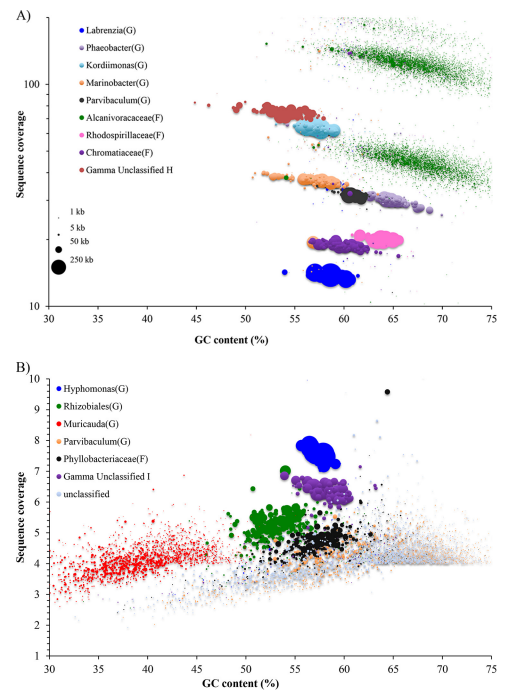


FIG 2 Contig sequence coverage is shown as points whose sizes are proportional to the contig length versus its GC content The color of each contig is based on the cluster to which it was assigned. (A) Sequence coverage in log scale depicting only higher-coverage organisms (>10). (B) Sequence coverage in linear scale depicting organisms with lesser coverage (<10).

which the AMPHORA2/Metawatt analyses identified as a member of the Sphingomonadaceae. A draft genome sequence of Kordiimonas was recently deposited in GenBank (NZ_AQXF00000000.1) and was not part of the database version used in the AMPHORA2 analysis. As such, the RAST identification of Kordiimonas was used for this cluster genome designation. Finally, 107 essential single-copy genes with conserved KEGG identifiers in 95% of all sequenced bacteria were enumerated from the predicted ORFs (see Data Set S1 in the supplemental material) (33, 42). Out of the 16 cluster genomes predicted by AMPHORA2 and Metawatt, all but *Parvibaculum*-2 contained at least 73 of the 107 housekeeping genes. The total length of all cluster genomes makes up 89% of the total assembled metagenome length. Cluster genomes were subsequently used to predict the origins of proteins identified by biocathode metaproteomics.

Biocathode metaproteome. A total of 644 proteins were iden-

TABLE 1 Summary of	f proteins identified	from the biocathode	metaproteome
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Identification	No. of proteins	No. of enzymes ^{<i>a</i>}	No. of receptors, transporters, and membrane proteins	No. of structural and hypothetical proteins
Marinobacter	177	69	50	58
Chromatiaceae	137	63	33	41
Labrenzia	59	20	32	7
Unclassified	45	5	22	18
Kordiimonas	44	8	20	16
Gammaproteobacteria-2	30	14	9	7
Rhodospirillaceae	29	12	10	7
Phaeobacter	26	6	15	5
Alcanivoracaceae	23	10	6	7
Rhizobiales	19	2	12	5
Gammaproteobacteria-1	12	7	1	4
Phyllobacteriaceae	11	4	2	5
Parvibaculum-1	10	3	4	3
Muricauda	8	2	6	0
Hyphomonas	7	3	3	1
Parvibaculum-2	7	4	2	1
Total	644	232	227	185

^{*a*} Enzymes include all proteins for which an EC number could be obtained or for which an enzyme domain was found to be present from the NCBI annotation. Cytochromes, peroxiredoxins, thioredoxin, molybdopterin, and ferredoxins were included in the enzyme category.

tified from the biocathode biofilm (see Data Set S2 in the supplemental material), 599 (93%) of which could be mapped to cluster genomes (Table 1). The *Marinobacter*, *Chromatiaceae*, and *Labrenzia* cluster genomes were the most represented taxa in the metaproteome, with 177, 137, and 59 proteins identified, respectively. Annotation, classification, and predicted localization information for each of the identified proteins can be found in Data Set S2 in the supplemental material; it was used to search for proteins associated with biocathode EET and carbon fixation, as described below.

Evidence for biocathode EET by an unknown member of the *Chromatiaceae.* A suite of consensus genetic markers is not known for cathode EET. Based on the identification and relative abundance of *Marinobacter* bacteria (a known FeOB) from our previous work (22), as well as recent demonstrations of cathode EET by other FeOB (13, 14), we hypothesized that our biocathode consortium utilizes iron oxidation pathways for biocathode EET. We explored the biocathode metagenome for genetic evidence of known or predicted iron oxidation pathways from all characterized FeOB (43). Table 2 summarizes the results of this search, including instances in which either the protein product was identified by proteomics or gene expression was confirmed by RT-PCR. The most interesting finding was the discovery that the Chromatiaceae cluster genome contains homologs of genes for proteins from two different putative FeOB EET pathways. The family Chromatiaceae is most known for the purple sulfur bacteria, which use light energy to oxidize sulfide under anoxygenic conditions. The *pufLM* (photosynthetic unit forming) operon represents a conservative marker for photosynthesis within the Chromatiaceae (44); however, we were not able to find pufLM operon homologs within the Chromatiaceae cluster genome or within the entire metagenome, suggesting that the Chromatiaceae

TABLE 2 Distribution of proteins among cluster genomes reported to be important for iron oxidation in FeOB

Protein ID ^{<i>a</i>} (iron oxidation)	Contig identifier	Cluster genome	RT-PCR ^b	Protein observed	Reference
Cytochrome <i>c</i> family protein	NODE_83_length_128150_cov_18.722910_73	Chromatiaceae			46
MopB, molybdopterin	NODE_83_length_128150_cov_18.722910_74	Chromatiaceae	+	Yes	46
oxidoreductase Fe ₄ S ₄ region					
4Fe-4S ferredoxin, iron-sulfur	NODE_83_length_128150_cov_18.722910_75	Chromatiaceae		Yes	46
binding protein					
Cytochrome <i>c</i> family protein	NODE_83_length_128150_cov_18.722910_76	Chromatiaceae			46
Quinol-cytochrome <i>c</i> oxidoreductase	NODE_83_length_128150_cov_18.722910_77	Chromatiaceae		Yes	46
Cytochrome <i>c</i> family protein	NODE_83_length_128150_cov_18.722910_78	Chromatiaceae		Yes	46
Quinol-cytochrome <i>c</i> oxidoreductase	NODE_83_length_128150_cov_18.722910_79	Chromatiaceae			46
Cyc2, cytochrome <i>c</i> family protein	NODE_9998_length_33057_cov_18.081980_35	Chromatiaceae		Yes	47, 48
Outer membrane protein (MtrB)	NODE_12611_length_8169_cov_61.299057_2	Kordiimonas	-		56
Cytochrome <i>c</i> family protein (MtrA)	NODE_12611_length_8169_cov_61.299057_3	Kordiimonas	_		56
Cytochrome <i>c</i> family protein (MtrA)	NODE_40928_length_81659_cov_6.155341_59	Gammaproteobacteria-1	_		56
Outer membrane protein (MtrB)	NODE_40928_length_81659_cov_6.155341_60	Gammaproteobacteria-1	-		56

^{*a*} Shaded and unshaded proteins were identified as belonging to distinct cluster genomes.

^b +, RT-PCR was positive for gene expression for the protein; -, no gene expression was detected.

TABLE 3 Distribution of	predicted multiheme	c-type cytochromes a	among cluster genomes

	No. of CXXCH	Predicted molecular			
Protein ID ^a	domains	mass (kDa)	Contig identifier	Cluster genome	$RT-PCR^b$
Hypothetical protein	5	47	NODE_127_length_181001_cov_19.410473_22	Marinobacter	+
Cytochrome c family protein	4	29	NODE_83_length_128150_cov_18.722910_103	Chromatiaceae	+
Cytochrome c family protein	11	98	NODE_83_length_128150_cov_18.722910_112	Chromatiaceae	+
Cytochrome <i>c</i> family protein (lp)	6	62	NODE_304_length_51348_cov_19.644348_22	Chromatiaceae	
Cytochrome c family protein	5	27	NODE_1003_length_21147_cov_19.239466_11	Chromatiaceae	+
Cytochrome <i>c</i> (554)	5	84	NODE_414_length_627542_cov_13.795955_434	Labrenzia	
Cytochrome <i>c</i> (554)	4	37	NODE_414_length_627542_cov_13.795955_437	Labrenzia	+
Cytochrome <i>c</i> family protein	10	94	NODE_414_length_627542_cov_13.795955_440	Labrenzia	+
Cytochrome <i>c</i> family protein	4	25	NODE_341_length_165593_cov_13.624054_22	Labrenzia	+
Cytochrome <i>c</i> family protein	8	71	NODE_750_length_164268_cov_13.758200_104	Labrenzia	
Cytochrome <i>c</i> family protein (MtrA)	9	35	NODE_12611_length_8169_cov_61.299057_3	Kordiimonas	_
Cytochrome <i>c</i> family protein	6	35	NODE_203_length_183696_cov_75.607124_159	Gammaproteobacteria-2	+
Cytochrome <i>c</i> -type protein NapC	3	22	NODE_1790_length_115957_cov_20.493477_12	Rhodospirillaceae	+
Cytochrome <i>c</i> family protein	4	25	NODE_21_length_106609_cov_30.063549_87	Phaeobacter	-
Cytochrome <i>c</i> family protein	8	70	NODE_355_length_35054_cov_33.539139_11	Phaeobacter	
Cytochrome c family protein	5	53	NODE_32589_length_12848_cov_5.104218_7	Rhizobiales	_
Cytochrome c family protein	6	31	NODE_32589_length_12848_cov_5.104218_8	Rhizobiales	
Cytochrome <i>c</i> family protein (MtrA)	9	38	NODE_40928_length_81659_cov_6.155341_59	Gammaproteobacteria-1	-
Periplasmic nitrate (or nitrite) reductase c-type cytochrome NapC/NirT family	4	26	NODE_367_length_177668_cov_31.499155_15	Parvibaculum-1	+
Cytochrome <i>c</i> family protein	2	12	NODE_3395_length_306_cov_130.271240_1	Alcanivoraceae	
Cytochrome <i>c</i> family protein ^{<i>a</i>} Shaded and unshaded proteins were identified	2	22	NODE_10082_length_765_cov_4.294117_2	no cluster assignment	_

^a Shaded and unshaded proteins were identified as belonging to distinct cluster genomes. lp, predicted lipoprotein.

^b +, RT-PCR was positive for gene expression for the protein; –, no gene expression was detected.

cluster genome probably represents a nonphotosynthetic member of the family. The entire genomic locus for a MopB-containing alternative complex III (ACIII), recently described as part of a putative iron oxidation pathway in Zetaproteobacteria (45, 46), was observed on a single contig within the Chromatiaceae cluster genome. The cluster includes genes for two multiheme c-Cyts, a 4Fe-4S ferredoxin, iron-sulfur binding protein, an integral transmembrane polysulfide reductase (NrfD), and two quinol-cytochrome c oxidoreductases and showed synteny with M. ferrooxydans and Sideroxydans lithotrophicus, as well as Geobacter uraniireducens (see Table S5 in the supplemental material). Proteins for four out of seven components of the putative ACIII cluster were observed by proteomics (Table 2). No specific iron oxidase has been assigned linking the ACIII to EET in Zetaproteobacteria, but the current model suggests that an outer membrane c-Cyt may be involved, as predicted for other FeOB. Two potential electrode oxidase proteins were identified within the Chromatiaceae cluster genome. Expression of a monoheme c-Cyt with sequence similarity to the monoheme Cyc2 from Acidithiobacillus ferrooxidans (also known as Cyt572) was identified by proteomics (Table 2). Cyc2 has been demonstrated to be important for iron oxidation (47, 48) and is suspected of directly oxidizing Fe(II) at low pH. In spite of significant divergence in the overall protein sequences (sequence alignments are shown in Fig. S4A and B in the supplemental material), the positions of the heme-binding site and some surrounding residues in the N termini are conserved among the three proteins. The predicted average molecular mass of the putative monoheme Cyc2-like protein from Chromatiaceae is 56 kDa, compared to 46 kDa for A. ferrooxidans Cyc2 and 61 kDa for Cyt572.

Another *c*-Cyt of note identified from the *Chromatiaceae* cluster genome is a predicted undecaheme *c*-Cyt residing on the

same contig (Node_83) as genes for the ACIII (Table 3). If such a protein were associated with the outer membrane, it could potentially participate in EET, as is proposed for membrane-associated multiheme *c*-Cyts of *Shewanella* and *Geobacter* (49, 50). Other predicted proteins of interest within the same contig are genes for the *c*-Cyt biogenesis system, an outer membrane porin (whose expression was confirmed by proteomics [see Data Set S2 in the supplemental material), and three periplasmic triheme *c*-Cyt family proteins (see Data Set S3 in the supplemental material).

We did not identify homologs of the well-known A. ferrooxidans rusticyanin, of Cyc1, or of the aa3-type oxidase pathway linking Cyc2 to oxygen reduction from the biocathode metagenome, leading us to investigate other oxygen reduction pathways associated with FeOB. With some known exceptions, cytochrome *cbb*₃ oxidases are unique to Proteobacteria and are found in FeOB, such as M. ferrooxydans (45). Cytochrome cbb3 oxidases play a significant role in microaerobic respiration, with a high affinity for O₂ (51). Genes encoding CcoN (conserved subunit I of cytochrome *cbb*₃) were found in 12 different cluster genomes as part of the full ccoNOQP operon, and most were found by RT-PCR to be expressed (see Table S6 in the supplemental material). The only CcoN protein identified by proteomics was from the Chromatiaceae but was not part of the full ccoNOPQ operon (only ccoNO were present). A similar observation has been made for M. ferrooxydans, where only ccoNOP were identified in the genome (45). Partial reduction of O₂ during aerobic iron oxidation can lead to production of potentially toxic reactive oxygen species (ROS), such as hydrogen peroxide. Bacterial cytochrome *c* peroxidases (CCP) are responsible for catalyzing the two-electron reduction of hydrogen peroxide to water and are critical for detoxification of ROS (52). CCP have also been suggested to be important for iron oxidation by Marinobacter aquaeolei (53, 54). Six full-length genes encoding CCP were identified from the biocathode metagenome (see Table S7 in the supplemental material). The expression of four of these genes, of which three belong to the *Chromatiaceae* cluster genome and one belongs to the *Gammaproteobacteria*-1 cluster genome, was confirmed by RT-PCR. Expression of one of the three *Chromatiaceae* CCP was also observed by proteomics. The role of CCP in biocathode EET is not clear, but given a potential role in iron oxidation in other organisms, it warrants further investigation.

Aside from the FeOB pathways described above, homologs for *mtrAB* genes from both the *Kordiimonas* and *Gammaproteobacteria*-1 cluster genomes were identified in the metagenome (Table 2). The decaheme *c*-Cyt MtrA and the outer membrane protein MtrB are involved in *Shewanella oneidensis* EET (43, 55), and the homologs PioA and PioB from *R. palustris* are known to be involved in iron oxidation (43). Additionally, the MtrA/MtrB homologs MtoA and MtoB from *S. lithotrophicus* have been demonstrated to oxidize iron *in vitro* (56). Interestingly, gene expression was not detected by either RT-PCR or proteomics.

Genes for known iron oxidation proteins were not identified in any other cluster genome, including Marinobacter. We therefore surveyed the metagenome for additional putative EET pathways based on the assumption that electron transfer at the biocathode may be mediated by a membrane-associated multiheme, c-Cyt, as is thought to be the case for anode-respiring bacteria. Approximately 187 genes encoding putative c-Cyts were identified in the biofilm metagenome by searching for the conserved CXXCH heme-binding motif. Twenty-one of them contained multiple heme-binding sites and are listed in Table 3, along with their predicted molecular masses. Gene expression for most predicted c-Cyts was observed by RT-PCR for Labrenzia, Marinobacter, Rhodospirillaceae, Chromatiaceae, Parvibaculum-1, and Gammaproteobacteria-2; however, no proteins were observed. This could, in part, be due to known difficulties associated with detecting c-Cyts by mass spectrometry (57). Although at this time it is not possible to assign a role in EET to the identified c-Cyts, it is interesting that they were present in a number of biofilm constituents.

Eighteen representative isolates from six different cluster genomes could be cultivated and were qualitatively evaluated for the ability to oxidize iron as an initial screening for EET (see Table S8 in the supplemental material). A representative member of the Chromatiaceae could not be cultivated using the methods reported here, and efforts are ongoing. Since Marinobacter and Labrenzia are predicted to be relatively abundant in the biofilm and showed some indication of iron oxidation, isolates were also evaluated for EET with a poised electrode (+0.310 V SHE) with and without supplementation with acetate (2 mM) (see Table S8 and Fig. S5A and B in the supplemental material). Supplementation with acetate resulted in Marinobacter EET, with a maximum current 2 orders of magnitude lower than that typically measured for the biocathode community. No electrode EET was observed for Labrenzia. When acetate was omitted, an initial spike in current was observed for Marinobacter, possibly associated with cells initially attaching to the electrode surface; however, no further increase in current was observed. This sharp spike in current by the Marinobacter-inoculated reactor contrasts with an increase in current over time for biocathode community reactors and suggests that the biocathode consortium is needed to develop and sustain Marinobacter when no organic carbon source is provided.

Potential for biocathode-linked carbon fixation by an unknown member of the Chromatiaceae. The biocathodes described here are grown at equilibrium with atmospheric concentrations of O_2 (i.e., no additional aeration is provided), while CO_2 is the only carbon source provided and the electrode is assumed to be the primary electron donor. Purging the reactor of O₂ was previously shown to eliminate the current (22); therefore, we hypothesized that the biocathode biofilm was fixing CO₂ to support growth through an aerobic pathway. Table 4 summarizes 32 key Calvin-Benson-Bassham (CBB) cycle (58) genes and accessory genes that were identified in the biocathode metagenome almost exclusively from the Chromatiaceae cluster genome. Three genes encoding RubisCO were identified: two RubisCO form I (IAq and IAc) genes sharing 79% identity at the amino acid level in the Chromatiaceae cluster and one gene encoding a form IV RubisCO-like protein (RLP) in the Labrenzia cluster. The IAq locus included the RubisCO structural genes rbcL, rbcS, and cbbQO, encoding proteins important in RubisCO assembly, and the *cbbR* gene, encoding a Lys-type regulator. The IAc operon includes *rbcL* and *rbcS*, followed by genes encoding carboxysome shell peptides that are assumed to enhance the effectiveness of CO₂ capture (see Fig. S6 in the supplemental material). Of the 32 genes identified in the metagenome, 11 were identified in the metaproteome analysis, including IAq RbcS, phosphoribulokinase, and two carboxysome shell proteins. RT-PCR analysis confirmed that both form I rbcL genes were expressed.

The gene for form IV RLP was identified in the *Labrenzia* cluster genome. RLPs are structural homologs of RubisCO that are unable to catalyze CO_2 fixation (59). They form six deeply branching subclades, only two of which have defined biochemical functions, participating in methionine salvage pathways (60, 61). Another clade (form IV-Photo) has an unknown role during growth using thiosulfate as an electron donor (62). The RLP present in the *Labrenzia* bin is related to the form IV-NonPhoto clade, which has no known function. The gene encoding this protein was expressed; however, the protein was not detected (Table 4). Marker genes associated with alternative CO_2 fixation pathways were not observed, including genes for ATP citrate lyase, bifunctional carbon monoxide dehydrogenases (CODH)/acetyl-coenzyme A (CoA) synthase, malonyl-CoA reductase, propionyl-CoA synthase, and 4-hydroxybutyryl-CoA dehydratase.

Taken together, genomic, RT-PCR, and proteomic evidence of CBB cycle proteins and complex inner membrane respiratory proteins (NADH/quinone oxidoreductase, cytochrome *bc*₁ complex, cytochrome cbb₃ oxidase, and ACIII), as well as putative EET proteins, identified from the Chromatiaceae cluster genome could account for electrode-driven autotrophy under microaerobic conditions. Due to the positive potential of the electrode, CO₂ fixation through the CBB cycle in Chromatiaceae must to be linked to O₂ reduction in order to generate proton motive force to power reverse electron transport from the cytochrome bc_1 complex through the NADH-quinone oxidoreductase complex, as in other chemolithoautotrophs (reviewed in references 43 and 63). The ACIII could take the place of the cytochrome bc_1 complex, accepting electrons from a transperiplasm redox module and reducing the quinone pool. Energy conservation may occur via electron bifurcation or via proton motive force (64). The estimated $\Delta G'$ (' denotes standard conditions) ($-nF\Delta E^{\circ}$, where n = 1 mol, F = 96,485 J, and ΔE° is the formal potential) for reduction of oxygen (+0.8 V SHE) with an electrode poised at +0.310 V SHE is

TABLE 4 Key components of the CBB cycle from the biocathode

Protein ID ^{<i>a</i>} (CBB cycle)	Contig identifier	Cluster genome	RT-PCR ^b	Protein observed
	0	0	K1-I CK	observed
NAD-dependent glyceraldehyde-3-phosphate dehydrogenase	NODE_1248_length_53554_cov_18.300053_33	Chromatiaceae		3.7
Ribose 5-phosphate isomerase A	NODE_138_length_121478_cov_19.056250_101	Chromatiaceae		Yes
NAD-dependent glyceraldehyde-3-phosphate dehydrogenase	NODE_138_length_121478_cov_19.056250_112	Chromatiaceae		
Hypothetical transmembrane protein coupled to NADH- ubiquinone oxidoreductase chain 5 homolog	NODE_1400_length_162497_cov_18.433632_23	Chromatiaceae		
Fructose-bisphosphate aldolase class II	NODE_1400_length_162497_cov_18.433632_75	Chromatiaceae		
RubisCO activation protein CbbO	NODE_1400_length_162497_cov_18.433632_77	Chromatiaceae		
RubisCO activation protein CbbQ	NODE_1400_length_162497_cov_18.433632_78	Chromatiaceae		
Ribulose bisphosphate carboxylase small chain	NODE_1400_length_162497_cov_18.433632_79	Chromatiaceae		Yes
Ribulose bisphosphate carboxylase large chain	NODE_1400_length_162497_cov_18.433632_80	Chromatiaceae	+	
RubisCO operon transcriptional regulator	NODE_1400_length_162497_cov_18.433632_81	Chromatiaceae		
Phosphoribulokinase	NODE_1926_length_32342_cov_18.998732_5	Chromatiaceae		Yes
RubisCO operon transcriptional regulator CbbR	NODE_3212_length_30296_cov_18.496204_15	Chromatiaceae		
Fructose-bisphosphate aldolase class I	NODE_5577_length_143978_cov_18.963264_100	Chromatiaceae		
Fructose-bisphosphate aldolase class I	NODE_5577_length_143978_cov_18.963264_118	Chromatiaceae		
Fructose-bisphosphate aldolase class II	NODE_6036_length_19485_cov_18.536259_1	Chromatiaceae		Yes
Phosphoglycerate kinase	NODE_6036_length_19485_cov_18.536259_3	Chromatiaceae		Yes
NAD-dependent glyceraldehyde-3-phosphate dehydrogenase	NODE_6036_length_19485_cov_18.536259_4	Chromatiaceae		Yes
Transketolase	NODE_6036_length_19485_cov_18.536259_5	Chromatiaceae		Yes
Carboxysome shell protein CsoS1	NODE_7231_length_64919_cov_18.593540_1	Chromatiaceae		Yes
Putative carboxysome peptide B	NODE_7231_length_64919_cov_18.593540_2	Chromatiaceae		
Putative carboxysome peptide A	NODE_7231_length_64919_cov_18.593540_3	Chromatiaceae		
Carboxysome shell protein CsoS3	NODE_7231_length_64919_cov_18.593540_4	Chromatiaceae		
Carboxysome shell protein CsoS2	NODE_7231_length_64919_cov_18.593540_5	Chromatiaceae		
Ribulose bisphosphate carboxylase small chain	NODE_7231_length_64919_cov_18.593540_6	Chromatiaceae		
Ribulose bisphosphate carboxylase large chain	NODE_7231_length_64919_cov_18.593540_7	Chromatiaceae	+	
Ribulose-phosphate 3-epimerase	NODE_83_length_128150_cov_18.722910_81	Chromatiaceae		Yes
Fructose-1,6-bisphosphatase, type I	NODE_890_length_140831_cov_19.766834_44	Chromatiaceae		Yes
Carbon dioxide-concentrating mechanism/carboxysome shell protein	NODE_9998_length_33057_cov_18.081980_1	Chromatiaceae		Yes
Possible pterin-4 alpha-carbinolamine dehydratase-like protein	NODE_9998_length_33057_cov_18.081980_3	Chromatiaceae		
Putative sodium-dependent bicarbonate transporter	NODE_9998_length_33057_cov_18.081980_5	Chromatiaceae		
RubisCO operon transcriptional regulator	NODE_9998_length_33057_cov_18.081980_7	Chromatiaceae		
Form IV RLP	NODE_90_length_211422_cov_13.680662_51	Labrenzia	+	

^a Shaded and unshaded proteins were identified as belonging to distinct contigs.

^b +, RT-PCR was positive for gene expression for the protein.

-47 kJ/mol e⁻, which is within the range for other organisms powering cell biosynthesis by reverse electron transport using iron as an electron donor (65). A schematic summarizing the predicted EET pathway identified from the Chromatiaceae cluster genome, and supported by proteomics and RT-PCR, is presented in Fig. 3. This proposed scheme does not explain how fixed carbon from Chromatiaceae may be distributed to other members of the biocathode consortia for biomass formation or how much total biomass could be generated. The total extractable cell biomass measured from surrogate reactors increased by an order of magnitude or more before reaching maximum current and ranged from 1.41E6 to 6.03E6 cells (see Fig. S7 and Table S9 in the supplemental material). The accumulated biomass was independent of the number of cells in the inoculum (2.0E4 versus 2.0E5 cells), indicating that the initial number of cells available to attach to the electrode surface does limit current. If all attached cells are assumed to contribute to the current, either by direct electron transfer or by driving EET through syntrophy, the rate of cell-normalized EET ranges from 0.29 to 2.62 pmol electrons h⁻¹ cell⁻¹. This estimate is ca. 4 to 40 times greater than the 0.075 pmol electrons h^{-1} cell⁻¹ reported for *M. ferrooxydans* cathodes (13). This could

be due to a real increase in the rate of EET for our biocathode community or to an underestimation of the number of electrodeassociated cells by an order of magnitude or more due to challenges in disaggregating the biofilm for flow cytometry.

An alternative source of biofilm carbon and energy: CO oxidation. CODHs have recently been found to be abundant in diverse marine bacteria (66, 67), suggesting that aerobic CO oxidation is a widespread metabolic strategy in the marine environment. CO is present in the atmosphere at 0.35 to 0.5 ppm (68) and could conceivably be available to biofilm organisms as an inorganic carbon source or a supplemental energy source. Labrenzia spp. are among the bacteria known to oxidize CO via carbon monoxide dehydrogenase to CO_2 (69) and are estimated to make up ca. 5% of the biofilm composition, leading us to examine the genomic potential of the biocathode biofilm for CO oxidation. CODH genes were classified into two known forms based on sequence divergence of the large subunit CoxL: form I, known as the OMP group (from Oligotropha, Mycobacterium, and Pseudomonas), is well characterized in the classic carboxydotrophs and is able to oxidize CO; form II, known as the BMS group (from Bradyrhizobium, Mesorhizobium, and Sinorhizobium), is a putative CODH inferred from se-

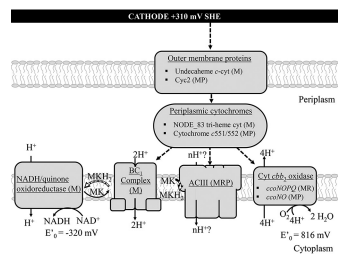


FIG 3 Schematic of the hypothetical electron transfer pathway between the electrode and *Chromatiaceae* respiratory proteins. The oxidative branch of the pathway generates energy and proton motive force through cytochrome cbb_3 oxidase. The proton motive force is used to generate ATP and to power the reductive branch, which produces NADH to provide reducing equivalents for CO₂ fixation. Both the cytochrome bc_1 complex and the putative ACIII are capable of reducing quinones, such as menaquinone (MK), accepting electrons from cytochromes at potentials within the range expected here. Support for the presence of these pathways in the uncultivated *Chromatiaceae* is indicated by the following: evidence from the metagenome only (M), an RT-PCR product found (for at least one gene in a complex) (P). E'₀ indicates the standard reduction potential.

quence homology to form I. The function of form II CoxL remains unknown, but it may preferentially oxidize alternate substrates (66). A total of nine CODH operons (coxLMS) were found in the biofilm metagenome and were distributed among the Labrenzia, Rhodospirillaceae, Rhizobiales, Parvibaculum-1, and Gammaproteobacteria-1 cluster genomes (Table 5). Of those nine, one of two form I and the form II CoxM protein from the Labrenzia cluster genome were observed by proteomics. RT-PCR of the large-subunit CoxL protein from the same operons in which CoxM was observed showed that it was also expressed. Expression of the second coxL form I operon of the Labrenzia cluster genome was not observed by either proteomics or RT-PCR. RT-PCR of genes encoding CoxL from the remaining six operons detected expression of a form II from the Parvibaculum-1 cluster genome and a form I from the Gammaproteobacteria-1 cluster genome. It is interesting that the Labrenzia, Rhodospirillaceae, and Parvibaculum-1 cluster genomes contain multiple copies or both forms of the cox operons. The presence of both forms may provide an ecological advantage under a range of CO conditions and substrates, as has been noted previously for a diversity of Labrenzia spp. (69). As noted above, genetic evidence for CO₂ fixation could not be found for CODH-containing cluster genomes, and a possible role for this metabolism is discussed further below.

DISCUSSION

In this study, we used metagenomics and metaproteomics to confirm the identities of major biofilm constituents, to identify the major CO_2 fixation pathway, and to provide an initial characterization of putative EET pathways of a previously described selfregenerating and self-sustaining biocathode community. The community is of low diversity, with *Gammaproteobacteria* estimated to make up ca. 60% of the constituents, while the remaining 35 to 40% belong mainly to the *Alphaproteobacteria*. This distribution is consistent with previous clone library analyses, where the majority of clones were *Gammaproteobacteria*, including *Marinobacter* and *Chromatiaceae* (22). *Alphaproteobacteria*, *Planctomycetes*, and *Actinobacteria* were also previously identified by clone library analysis; however, the last two phyla were not detected here by metagenomics, suggesting they were not essential to biocathode performance and may have been lost during subsequent transfers.

In general, the distribution of biocathode proteins among cluster genomes correlated with the predicted taxonomic distribution of the metagenome, with more abundant taxa representing a higher proportion of identified proteins. With the exception of Parvibaculum-1, cluster genomes with higher sequence coverage also had a greater number of proteins identified. For example, Labrenzia, Phaeobacter, and Kordiimonas had the highest sequence coverage among Alphaproteobacteria identified to the genus level and together represent 18% of all identified proteins. Likewise, Marinobacter and Chromatiaceae are two of the most abundant Gammaproteobacteria and made up 27% and 21% of all identified proteins, respectively. The family Alcanivoraceae was predicted to make up 33% of Gammaproteobacteria but was not previously identified by clone library analysis and accounted for only 4% of all identified proteins. This discrepancy may be due to short contig lengths, which can result in artificially high abundance counts and/or may result in undercounting Alcanivoraceae proteins, as short peptide sequences could be insufficient for MS identification.

Cluster genomes generated using metagenomics were used to address two fundamental questions currently facing biocathode technologies: can microorganisms gain energy for growth through electrode-driven autotrophy and, if so, what are the biocathode EET and CO₂ fixation pathways? The proteomic and genomic observations presented here confirm that an uncharacterized member of the family Chromatiaceae expresses proteins for autotrophic CO₂ fixation and could potentially use the electrode as an electron donor. Genes encoding proteins of proposed iron oxidation pathways, including Cyc2 (Cyt572) and the Zetaproteobacteria ACIII, were identified from the Chromatiaceae cluster genome, and expression was observed by proteomics or RT-PCR. Experimental evidence is still needed to confirm the involvement of ACIII in EET during iron oxidation and has thus far been inferred only from genomic and proteomic observations. The fact that a known metal reducer, G. uraniireducens, also contains a homologous ACIII but is not known to oxidize iron further confounds its role.

Key enzymes from the CBB cycle were identified only in the *Chromatiaceae* cluster genome. The presence of form I RubisCO suggests that these enzymes are directly involved in carbon metabolism associated with the CBB cycle (59, 70, 71). At this time, we have no direct evidence for electrode-driven growth of *Chromatiaceae*, as we have been unable to cultivate a representative isolate. Electrode-associated cell biomass increases over time as the magnitude of the current increases while no other electron donor is provided other than the poised electrode and no other source of carbon is intentionally provided other than CO₂. The rates of cell-normalized EET are at least as high as that previously noted for *M. ferrooxydans* growing on a cathode (13), suggesting that the num-

TABLE 5 Distribution of the coxLMS operon among biocathode cluster genomes

Protein ID ^{<i>a</i>} (carbon monooxide oxidation)	Contig identifier	CODH form	Cluster genome	RT-PCR ^b	Protein observed
CoxL, carbon monoxide dehydrogenase, large	NODE_1964_length_374203_cov_14.158323_147	Ι	Labrenzia	+	
subunit CoxM, carbon monoxide dehydrogenase, medium subunit	NODE_1964_length_374203_cov_14.158323_149		Labrenzia		Yes
	NODE_1964_length_374203_cov_14.158323_148		Labrenzia		
CoxL, carbon monoxide dehydrogenase, large subunit	NODE_1964_length_374203_cov_14.158323_221	II	Labrenzia	+	
CoxM, carbon monoxide dehydrogenase, medium subunit	NODE_1964_length_374203_cov_14.158323_220		Labrenzia		Yes
CoxS, carbon monoxide dehydrogenase, small subunit	NODE_1964_length_374203_cov_14.158323_222		Labrenzia		
	NODE_440_length_159716_cov_13.704870_27	Ι	Labrenzia	-	
CoxM, carbon monoxide dehydrogenase, medium subunit	NODE_440_length_159716_cov_13.704870_29		Labrenzia		
CoxS, carbon monoxide dehydrogenase, small subunit	NODE_440_length_159716_cov_13.704870_28		Labrenzia		
CoxL, carbon monoxide dehydrogenase, large subunit	NODE_498_length_292238_cov_20.118721_27	Π	Rhodospirillaceae	-	
CoxM, carbon monoxide dehydrogenase, medium subunit	NODE_498_length_292238_cov_20.118721_26		Rhodospirillaceae		
CoxS, carbon monoxide dehydrogenase, small subunit	NODE_498_length_292238_cov_20.118721_25		Rhodospirillaceae		
CoxL, carbon monoxide dehydrogenase, large subunit	NODE_498_length_292238_cov_20.118721_82	Ι	Rhodospirillaceae	-	
CoxM, carbon monoxide dehydrogenase, medium subunit	NODE_498_length_292238_cov_20.118721_80		Rhodospirillaceae		
	NODE_498_length_292238_cov_20.118721_81		Rhodospirillaceae		
CoxL, carbon monoxide dehydrogenase, large subunit	NODE_303_length_99799_cov_30.081644_73	II	Rhizobiales	-	
CoxM, carbon monoxide dehydrogenase, medium subunit	NODE_303_length_99799_cov_30.081644_74		Rhizobiales		
	NODE_303_length_99799_cov_30.081644_72		Rhizobiales		
	NODE_610_length_56213_cov_74.588669_37	Ι	Gammaproteobacteria-1	+	
CoxM, carbon monoxide dehydrogenase, medium subunit	NODE_610_length_56213_cov_74.588669_35		Gammaproteobacteria-1		
	NODE_610_length_56213_cov_74.588669_36		Gammaproteobacteria-1		
	NODE_97_length_191039_cov_31.216541_178	II	Parvibaculum-1	+	
CoxM, carbon monoxide dehydrogenase, medium subunit	NODE_97_length_191039_cov_31.216541_177		Parvibaculum-1		
	NODE_97_length_191039_cov_31.216541_179		Parvibaculum-1		
CoxL, carbon monoxide dehydrogenase, large subunit	NODE_247_length_237777_cov_31.838627_173	II	Parvibaculum-1	-	
CoxM, carbon monoxide dehydrogenase, medium subunit	NODE_247_length_237777_cov_31.838627_174		Parvibaculum-1		
	NODE_247_length_237777_cov_31.838627_172		Parvibaculum-1		

^a Shaded and unshaded proteins were identified as belonging to distinct contigs.

^b +, RT-PCR was positive for gene expression for the protein; –, no gene expression was detected.

ber of electrons from the electrode is sufficient for autotrophic growth. Future studies using stable-isotope labeling need to address what proportion of accumulated biomass results from CO₂ fixation by *Chromatiaceae* and just how this fixed carbon is dis-

tributed to other biofilm constituents. Temporal analysis of biofilm development using fluorescence *in situ* hybridization (FISH) may also help to determine the dependence of the growth of one organism on another and may shed light on what limits biocathode current. Biocathodes are limited by oxygen diffusion to the electrode, as well as the buildup of ROS from oxygen reduction, both of which could limit biofilm growth. Due to the fact that the biocathode community is enriched from the environment and many constituents have not been characterized, we cannot rule out the possibility that degradation of reactor components (nylon screws, wire insulation, or chelating agents in mineral solution) could contribute to the biocathode carbon supply. Although degradation has not been specifically observed, at least two biocathode constituents, *Kordiimonas* and *Alcanivoraceae*, are related to known hydrocarbon-degrading bacteria with appetites for unconventional sources of carbon.

A significant role in CO₂ fixation or biocathode EET could not be predicted for biocathode constituents other than Chromatiaceae using the search parameters in this study. The Marinobacter, Labrenzia, Gammaproteobacteria-2, Rhodospirillaceae, and Parvibaculum-1 cluster genomes all contained genes for multiheme c-Cyt that may participate in EET, and expression was observed by RT-PCR. Several biocathode isolates exhibited the capacity to oxidize iron in gradient tubes when small amounts of organic carbon were supplied, and Marinobacter had some capacity for electrode EET, indicating possible mixotrophy or, as others have noted, "opportunitrophy" (54). Given its estimated abundance in the biofilm and observed interaction with the electrode, it was surprising that specific EET proteins could not be prescribed for Marinobacter. Other potential Marinobacter EET pathways need to be explored, aside from those currently known for FeOB, including those containing proteins with redox cofactors other than iron, such as molybdenum and copper.

This study focused on EET and CO₂ fixation; however, proteins that may be important for life at the biocathode were also identified. In general, enzymes from more abundant biofilm constituents were those involved in amino acid and nucleotide biosynthesis, carbohydrate metabolism, fatty acid metabolism, and mitigation of oxygen stress. With the exception of the Chromatiaceae, most cluster genomes contained a number of ABC or TRAP transporters and TonB receptors (see Data Set S2 in the supplemental material). This supports the idea that Chromatiaceae are the primary biofilm producers, while other constituents are most likely involved in carbon cycling. Proteins prevalent among all the cluster genomes included structural proteins for motility and biofilm formation, such as flagella, type IV pili, fimbria, and chemotaxis. Members of the order Rhodobacterales, which includes Phaeobacter, Labrenzia, and Hyphomonas spp., have been reported to be ubiquitous and dominant primary surface colonizers of biocorroding communities in temperate coastal waters (72) and deep-sea environments (23) and to be significantly abundant in acetogenic multispecies biocathodes (16). The Kordiimonas genome cluster was highly represented among the Alphaproteobac*teria*; however, the role of the organism within the biocathode community is also not clear. Many of the identified proteins from Kordiimonas were hypothetical due to the genome sequence only recently being made available. Few proteins were identified from cluster genomes with very low sequence coverage (i.e., Muricauda, Phyllobacteriaceae, and Hyphomonas), which would indicate a specific role in the biocathode biofilm.

An unexpected finding was that CO oxidation appeared to be active in the biocathode community and may represent a dynamic metabolic strategy for energy and carbon acquisition. We identified expression of the *coxL* gene from the *Labrenzia*, *Parvibacu*- *lum*-1, and *Gammaproteobacteria*-1 cluster genomes. While not much is known about CO oxidation by *Parvibaculum, L. aggregata* is categorized as a carboxydovore, where only low concentrations of CO are oxidized during mixotrophic metabolism in the presence of other organic substrates (69). While it is not immediately apparent how CO oxidation is incorporated into the carbon cycle of the biofilm, we can imagine a scenario where CO is oxidized by members of *Labrenzia, Parvibaculum*-1, or *Gammaproteobacteria*-1 to CO_2 for fixation by *Chromatiaceae*. Such a relationship may explain why these particular biocathode constituents remain part of the consortium and may also indicate a potential target for functional engineering to modify biocathode carbon acquisition.

Conclusions. In order to improve biocathode performance through functional engineering for MFCs, microbial electrosynthesis, or other biotechnology applications, there is a need to understand EET and energy conservation in such systems. Naturally enriched biocathode consortia operating at more positive electrode potentials and under aerobic conditions may have an advantage over homogeneous cell populations in terms of their robustness and stability under changing conditions, such as pH, temperature, and salinity, which are relevant to operating in the marine environment. In this study, metagenomics combined with metaproteomics provided a more comprehensive understanding, beyond previous 16S rRNA gene clone libraries, of the primary constituents of a self-regenerating and self-sustaining biocathode biofilm. We confirmed expression of a major autotrophic CO₂ fixation pathway (the CBB cycle) from a nonphotosynthetic, uncharacterized member of the family Chromatiaceae. We have also presented proteins for putative EET pathways from the electrode to this organism that could potentially drive CO₂ fixation. Thus far, efforts to cultivate Chromatiaceae off the electrode have been unsuccessful, highlighting the importance of using an omics approach to study microbial communities. Although the greatest number of proteins were identified from Marinobacter, which was also shown to engage in EET with iron and the electrode, no specific EET pathways could be identified. The metagenomic and metaproteomic analyses reported here will serve as the basis for future studies to determine the roles of other biocathode constituents and whether their relationships can be exploited to manipulate biocathode performance and metabolism.

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