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Genomic and physiological characterization of the chromate-reducing, aquifer-derived firmicute *Pelosinus* sp. strain HCF1

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Running title: Genome and physiology of *Pelosinus* sp. strain HCF1

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Abstract

*Pelosinus* spp. are fermentative firmicutes that were recently reported to be prominent members of microbial communities at contaminated subsurface sites in multiple locations. Here we report metabolic characteristics and their putative genetic basis in *Pelosinus* sp. strain HCF1, an isolate that predominated anaerobic, Cr(VI)-reducing columns constructed with aquifer sediment. Strain HCF1 ferments lactate to propionate and acetate (the methylmalonyl-CoA pathway was identified in the genome) and its genome encodes two [NiFe]- and four [FeFe]-hydrogenases for H₂ cycling. Reduction of Cr(VI) and Fe(III) may be catalyzed by a flavoprotein with 42-51% sequence identity to both ChrR and FerB. This bacterium has unexpected capabilities and gene content associated with reduction of nitrogen oxides, including dissimilatory reduction of nitrate to ammonium (two copies of NrfH and NrfA were identified along with NarGHI) and a nitric oxide reductase (NorCB). In this strain, either H₂ or lactate can act as a sole electron donor for nitrate, Cr(VI), and Fe(III) reduction. Transcriptional studies demonstrated differential expression of hydrogenases and nitrate and nitrite reductases. Overall, the unexpected metabolic capabilities and gene content reported here broaden our perspective on what biogeochemical and ecological roles this species might play as a prominent member of microbial communities in subsurface environments.

Introduction

The throughput, depth, and reduced cost of second-generation DNA sequencing facilitates our ability to gain insight into a broad range of microbiological processes in the environment, and genome sequencing of prominent members of environmental microbial communities should contribute to the understanding of complex biogeochemical systems. Members of the *Veillonellaceae*, and particularly *Pelosinus* spp., have recently been reported to
be among the more abundant bacterial taxa in chromate-reducing systems inoculated with material from the chromium-contaminated aquifer at the U.S. Department of Energy (DOE) Hanford 100H site in Washington state (1, 2) and in other contaminated aquifers (3-5).

Chromate-reducing bacteria are of interest because in situ reductive immobilization is favored as one of the more cost-effective approaches to remediation of aquifers contaminated with Cr(VI), a potent toxicant, mutagen, and carcinogen (6, 7). Fermentative bacteria such as Pelosinus spp. may be of particular relevance to a common bioremediation scenario in which metabolism of organic electron donors (e.g., lactate-based polymers) injected into the subsurface readily consume available electron acceptors (e.g., oxygen, nitrate, sulfate, ferric iron) and drive the treated zone towards fermentative/methanogenic conditions.

In this article, we report on a variety of metabolic capabilities and their possible underlying genetic basis in a Pelosinus isolate that dominated a chromate-reducing community derived from aquifer sediment from the Hanford 100H site. The metabolic capabilities explored include lactate fermentation to propionate and acetate (related to the methylmalonyl-CoA pathway identified in the genome), Cr(VI) and Fe(III) reduction (both potentially related to identified flavoproteins), nitrate and nitrite reduction (potentially related to NrfH and NrfA, as well as a membrane-bound, respiratory nitrate reductase), and H₂ metabolism (two [Ni-Fe]-hydrogenases and four [FeFe]-hydrogenases were identified). We also report on focused transcriptional studies designed to more clearly associate certain genes with specific metabolic activities (namely, H₂ cycling and nitrate or nitrite reduction). Some metabolic activities and gene content reported here are unexpected for Pelosinus species and broaden our perspective on what metabolic and ecological roles this species might play in microbial communities in contaminated (and uncontaminated) environments.
MATERIALS AND METHODS

Isolation and cultivation of *Pelosinus* sp. strain HCF1. *Pelosinus* sp. strain HCF1 was isolated from the effluent of an anaerobic, chromate-reducing, flow-through column containing aquifer sediment from the DOE Hanford 100H site and eluted with sterile, synthetic groundwater. The synthetic groundwater contained the following constituents: 1 mM phosphate buffer (0.58 mM K$_2$HPO$_4$ and 0.42 mM KH$_2$PO$_4$; pH 7), 1.25 mM NaHCO$_3$, 0.25 mM CaCl$_2$, 0.1 mM NH$_4$Cl, 5 mM sodium lactate (the carbon source and electron donor), 5 μM KCrO$_4$, and 7.5 mM Na$_2$SO$_4$. Highly purified water (18 MΩ resistance) obtained from a Milli-Q Biocel system (Millipore, Bedford, MA) was used to prepare the synthetic groundwater and all other aqueous solutions described in this article. Although sulfate was available as an electron acceptor, ion chromatographic monitoring of the effluent confirmed that lactate was fermented to acetate and propionate and that sulfate reduction was not required for this process.

Isolation of strain HCF1 from column effluent was accomplished using a modifiedHungate roll-tube technique (8) with serial dilutions to 10$^{-5}$. Direct microscopic cell counts showed that there were approximately 1x10$^7$ cells/mL in the effluent samples. Basal medium (0.58 mM K$_2$HPO$_4$, 0.42 mM KH$_2$PO$_4$, 0.25 mM CaCl$_2$, 0.1 mM NH$_4$Cl) containing 2% Noble agar and amended with 20 mM sodium lactate, 10 mM NaHCO$_3$, and vitamin, trace element, and selenite-tungstate solutions described elsewhere (9) was used for isolation. The inoculated roll tubes were incubated at 30°C for 7-10 days until bacterial colonies appeared. Second and third rounds of purification were conducted until the cells and colonies were uniform based on microscopic examination. Although yeast extract is not essential for the growth of strain HCF1, 0.5 g/L yeast extract was routinely used in the medium to facilitate growth and colony formation.
After isolation, routine cultivation of strain HCF1 was carried out under strictly anaerobic conditions in an anaerobic glove box (Coy Laboratory Products, Inc., Grass Lake, Mich.) with a nominal gas composition of 86% N₂ – 10% CO₂ – 4% H₂. The growth medium was the same as the medium used for isolation with the omission of Noble agar. Anaerobic techniques used in the preparation of growth medium and stock solutions are described elsewhere (10).

**Cell suspension experiments testing Cr(VI) and Fe(III) reduction by strain HCF1.** All cell suspension experiments described in this article were performed under strictly anaerobic conditions in an anaerobic glove box. The glass and plastic materials used to contain or manipulate the cultures were allowed to degas in the glove box for at least one day before use. Cell suspension assays performed to assess chromate reduction by strain HCF1 under fermentative conditions were conducted in a similar manner to those described previously for *Pseudomonas* sp. strain RCH2 (9). Strain HCF1 was grown anaerobically at 30°C in a glove box, harvested in mid log phase (OD ≈ 0.2, after 48 hr of incubation) by anaerobic centrifugation (11,899 x g, 15 min, 15 °C), washed with 100 mL anaerobic basal medium, and resuspended anaerobically in the glove box. Assays were performed in butyl rubber-stoppered Balch tubes and initiated by adding 1 mL of concentrated cells to 9 mL of basal medium amended with the appropriate compounds, such as 50 μM chromate and 20 mM lactate (final concentration). Assays were run in duplicate at ~30°C. Samples (500 μL) were collected before cells were added, immediately after cell addition, and typically every 30 min thereafter. 200 μL of the suspension was used for OD₆₀₀ (optical density at 600 nm) measurement and the remainder was spun down at ~20,800 x g (4°C, 4 min). The freshly prepared supernatants and cell pellets were used for various analyses described below. Controls included assays that did not contain lactate (normally added at 20 mM) and in which cells were inactivated by heating (under anaerobic
conditions) in a boiling water bath for 15 min. The potential role of extracellular reductants in Cr(VI) reduction was investigated by conducting assays in 10 mL of filtered (0.2-μm pore size) spent medium after growth or 10 mL of filtered, spent medium that had been heat-denatured anaerobically in a boiling water bath for 15 min. These tests of extracellular reductants were initiated by adding 50 μM KCrO$_4$.

Anaerobic cell suspension experiments to assess reduction of Fe(III) (2 mM) under fermentative conditions were conducted as described for Cr(VI) reduction studies, except that un-centrifuged samples (not supernatants) were used for determination of the reduced metal. Fe(III)-NTA stock solutions (100 mM) were prepared by dissolving 0.82 g NaHCO$_3$, 1.28 g of trisodium nitrilotriacetic acid (Na$_3$NTA), and 1.35 g of FeCl$_3$•6H$_2$O in 50 mL anaerobic H$_2$O in an anaerobic glove box and filter-sterilizing into an anaerobic serum bottle. Amorphous Fe(OH)$_3$ was prepared as described previously (11) and stored in an anaerobic glove box for at least several weeks before use to de-oxygenate.

**Analytical methods.** Analytical methods for all analytes other than Fe(II) were described in detail previously (9) and are briefly summarized here. For routine determination of cell density, OD$_{600}$ of diluted samples was measured using a Model UV160U spectrophotometer (Shimadzu Corporation). It was determined that 1 OD$_{600}$ unit was equivalent to 1x10$^9$ cells/mL. Direct microscopic cell counts were also determined in cell suspension experiments. Lactate, acetate, propionate, nitrate, and nitrite concentrations were measured by ion chromatography (IC) using a Model ICS-2000 IC (Dionex Corporation) (9). Cr(VI) in cell suspension supernatants was determined with a spectrophotometric DPC (diphenylcarbazide) microplate assay (9). Total Cr in cell suspension supernatants was measured by inductively coupled plasma-mass spectrometry (ICP-MS) (9).
Fe(II) generated from Fe(III)-NTA or Fe(OH)_3 reduction by HCF1 cells was measured with a spectrophotometric Ferrozine assay conducted in an anaerobic glove box. The microplate assay used in this study was adapted from a method described by Stookey (12). Microplates (96-well) that had been stored in an anaerobic glove box for at least one day were amended with 90 μL of 1N HCl, and a 10 μL cell suspension sample was added to the HCl immediately after sampling. Then, 100 μL of Ferrozine solution (1 g/L Ferrozine, 500 g/L ammonium acetate) was added to the acidified sample. After a 10-min incubation, absorbance at 555 nm was measured using a Model 550 microplate reader (Bio-Rad). Fe(II) standards (0.2, 0.5, 1, and 2 mM ferrous ammonium sulfate hexahydrate in 1N HCl) were included on each microtiter plate.

**Phylogenetic analysis of microbial communities in Hanford flow-through columns.** DNA was extracted from effluent of the same fermenting columns from which strain HCF1 was originally isolated. Effluents were collected directly into 50-mL tubes containing 30 ml of an RNA preservation reagent (25 mM sodium citrate, 10 mM EDTA, 10 M ammonium sulfate, pH 5.2). After overnight incubation at 4 ºC, the solution was filtered (0.22-μm pore size) and DNA and RNA co-extracted from filter sections using a modification of the extraction procedure reported by Ivanov and co-workers (13). This modification involved the addition of 1 g of Chelex-100 to the extractions to chelate iron; all other procedures were as reported. Extracted DNA (10 ng) was used as a template for PCR amplifications using primers (515F and 907R) targeting the V4-V5 hypervariable regions of the SSU rRNA gene with error-correcting barcodes (14):

515F-454: 5' CCTATCCCCTGTGTGCCTTGGCAGTCTCAGAGGTGCCAGCMGCCGCGGTAA 3'

907R-454: 5' GGACTACHVGGGTATCTAATCC 3'
169 5′CCATCTCATCCCTGCGTGTCTCCGACTCAGGTGAGGTCGCTAACCCGTCAATTCC
170 TTRAGTTT ′3
171
172 where underlined text represents a spacer, italics represents a barcode, and bold text represents a
173 priming region.
174
175 PCR reactions were performed using Takara Ex Taq polymerase, 0.625 units (Takara,
176 Madison, WI), 1X reaction buffer, 200 µM dNTPs, 0.56 mg/mL BSA, and 1 µM each primer
177 with the following thermocycling parameters: initial denaturation at 95 °C for 1 min followed by
178 25 cycles of 95 °C for 20 sec, 30 sec of annealing at 66 °C and extension at 72 °C for 1 min.
179 Final product extension was at 72 °C for 10 min. Reaction primer dimers were removed from the
180 PCR products via SPRI bead purification according to the manufacturer’s protocol (AMPure XP,
181 Beckman Coulter Genomics, Danvers, MA) before being checked for quality and quantity on a
182 Bioanalyzer 2100 using a DNA 7500 chip (Agilent Technologies, Santa Clara, CA). Each PCR
183 sample was normalized to 30 ng and samples were combined for multiplex sequencing.
184 Sequencing libraries were created using the LV emu-PCR kit (Lib-L, Roche, Indianapolis, IN)
185 and sequencing conducted on a GS-FLX sequencer (Roche, Indianapolis, IN) at the Veterans
186 Medical Research Foundation, La Jolla, CA.
187
188 Analysis of pyrotag sequences was performed using the QIIME suite of tools (15),
189 including quality filtering to Q20, clustering at 97% sequence homology and classification using
190 the RDP classifier (16).
191
192 **Phylogenetic analysis of strain HCF1.** The 16S rRNA gene of strain HCF1 was amplified by
193 the PCR using a universal bacterial 16S forward primer 27F (5′-
194 AGAGTTTGATCCTGGCTCAG-3′) and reverse primer 1492R (5′-TACCTTGTTACGACTT-
195 3′) (17). PCR products were cloned using One Shot TOP10 Chemically Competent *E. coli* and a
TOPO TA Cloning Kit (Invitrogen) and were sequenced at the UC Berkeley DNA Sequencing Facility (Berkeley, CA). The 16S rRNA gene sequence of strain HCF1 was BLASTed (18) against the GenBank nr and RDPII databases. A 16S rRNA phylogenetic tree was generated using MEGA version 4 (19).

**Genome sequencing, assembly, and annotation.** Genomic DNA library construction for Illumina sequencing of strain HCF1 was performed using the Nextera DNA Sample Prep Kit (Illumina Compatible) according to the manufacturer’s instructions using the LMW buffer (Epicenter Biotechnologies, WI). Fragment size distribution estimated using an Agilent Bioanalyzer (Agilent, CA) was centered at 400 bp with a distribution between 200-700 bp. The genome sequencing was performed on the Illumina platform (GAI1x) using a paired-end read library of read length 100 bp with an insert size of 450 bp. We obtained a total of 4,183,216 pairs of reads, corresponding to an estimated 40- to 80-fold coverage of the genome assuming a genome size of 5 to 10 Mb. All raw reads were quality-trimmed to an error probability of 1 in 1000 (Q30). After trimming, reads shorter than 20 nucleotides were removed. 3,957,260 reads were assembled *de novo* using Velvet (20) (k = 57, coverage cutoff = 24, expected coverage = 48, insert size = 450), generating 214 contigs with a N50 length of 79.5 kb.

Genome annotation was first performed automatically using the JGI IMG portal (21) and then manually curated (as described in the Results and Discussion). A search for the CXXCH heme-binding motif that characterizes c-type cytochromes was carried out using JGI IMG and ScanProSite (http://prosite.expasy.org/scanprosite/) search tools.

**Transcriptional (RT-qPCR) analysis of nitrate and nitrite reduction by strain HCF1**

**DNA/RNA extraction.** Duplicate 50-mL cultures of HCF1 were grown anaerobically using routine medium supplemented with 20 mM lactate with or without 5 mM NaNO3. Five-mL
samples of HCF1 cultures were collected in triplicate after 36 hours and 72 hours and immediately mixed with two volumes of the RNA preservation reagent described previously. Cells were then harvested anaerobically by centrifugation (10,300 x g, 20 min, 4°C) and DNA and RNA were extracted using an AllPrep DNA/RNA mini Kit (QIAGEN Sciences, Inc.). Cell pellets were resuspended in 600 µL RLT Plus buffer containing 1% β-mercaptoethanol and homogenized using a FastPrep-24 homogenization system (MP Biomedicals). DNA/RNA samples were then extracted according to the manufacturer’s instructions except that the residual DNA in RNA samples was digested on-column using RNase-Free DNase (QIAGEN) for 15 min. DNA/RNA samples were finally eluted in 30 µl RNase-free water and quantified using a NanoDrop 1000 Spectrophotometer (Thermo Fisher Scientific Inc.). DNA/RNA samples were stored at -80°C.

**PCR amplification.** For calibration standards, gene-specific primers were designed to amplify the *rpoB*, *narG*, *nrfH1*, and *nrfH2* genes from strain HCF1 (Table S1). PCR primers were designed with Primer 3 software (22). Fifty-µL PCR mixtures contained: 5 µL of 5X Ex Taq Buffer; 4 µL of dNTP Mix (2.5 mM of each dNTP); 2.5 µM forward and reverse primer (2 µL each); 0.3 µL TaKaRa Ex Taq DNA polymerase (Takara Bio Inc.); 100 ng DNA template, and nuclease-free water. PCR amplification was performed using a BioRad iCycler (Bio-Rad Laboratories) and the following thermocycling program: 2 min at 98°C; 30 cycles of 10 sec at 98°C, 30 sec at 55°C, 2 min at 72°C; 7 min at 72°C; maintained at 4°C. PCR products were purified using a QIAquick PCR purification Kit (QIAGEN). Amplicons of the *rpoB*, *narG*, *nrfH1*, and *nrfH2* genes were purified, quantified (NanoDrop 1000 spectrophotometer), and serially diluted as DNA templates for calibration (10 to $10^5$ copies/reaction).
**RT-qPCR analysis.** qPCR primers were designed with Primer 3 software. For each RNA sample, cDNA products were synthesized by SuperScript III reverse transcriptase and random hexamers (Invitrogen Corporation) according to the manufacturer’s instructions except that 2 µL 10 mM dNTP and 1 µL SuperaseIn (Ambion) were used in each RT reaction to inhibit RNA degradation. Eleven µL RNA was used as template in each 20-µL RT reaction. After RT, 20 µL final cDNA was ten-fold diluted by adding 180 µL nuclease-free water. For qPCR, 20-µL qPCR reaction mixtures contained 10 µL 2X SYBR Green Supermix (Bio-Rad Laboratories), 2.5 µM forward and reverse primers (Table S1) (2 µL each), diluted cDNA template (5 µL), and 1 µL nuclease-free water. qPCR analysis was performed with a MyiQ Single-Color Real-Time PCR Detection System (Bio-Rad Laboratories) using the following temperature cycling program: 2 min at 50°C; 8.5 min at 95°C; 40 cycles of 15 sec at 95°C, 30 sec at 50°C, 30 sec at 70°C, and 15 sec at 81°C. The real-time data were acquired at the 81°C-stage to exclude noise from primer dimers. For each gene, 5-point standards containing 10, 10², 10³, 10⁴, and 10⁵ copies/reaction were included in the same qPCR run to generate the calibration curve for absolute quantification.

**Transcriptional (RT-qPCR) analysis of hydrogenases in strain HCF1**

**DNA/RNA extraction.** One hundred-mL cultures of HCF1 were grown anaerobically using routine medium supplemented with 20 mM lactate or 20 mM fructose. Anaerobically, cells were then spun down, washed, resuspended in 10 mL basal medium supplemented with the same substrates (20 mM lactate or 20 mM fructose), 10 mM NaHCO₃, and fixed with a headspace of 80% N₂ / 20% CO₂. In addition, 100 mL of lactate-grown HCF1 cells were resuspended in 10 mL basal medium with 10 mM NaHCO₃, and fixed with a headspace of 90% H₂ / 10% CO₂. After 1 hr of incubation, triplicate 3-mL samples (~3 x 10⁹ cells) were mixed with 6 mL of the RNA preservation reagent described previously prior to RNA isolation.
**PCR amplification and RT-qPCR analysis.** PCR amplification to make calibration standards and RT-qPCR conditions were the same as described previously, except that PCR and qPCR primers for the six target hydrogenase genes from strain HCF1 are given in Table S2.

**Nucleotide sequence accession numbers.** Draft genome sequence data for *Pelosinus* sp. strain HCF1 have been deposited in GenBank under accession number [XX-will be added in proofs-XX]. 16S rRNA gene pyrotag sequence data for the flow-through column from which strain HCF1 was isolated have been deposited in the MG-RAST database under accession number 4502869.3.

**RESULTS AND DISCUSSION**

**Lactate fermentation to propionate and acetate.** *Pelosinus* sp. strain HCF1 ferments a range of sugars and short-chain mono- and dicarboxylic acids to propionate and acetate. For example, lactate, which has been used in polymeric form (glycerol polylactate) to stimulate *in situ* reductive immobilization of chromium in groundwater (23), is fermented to propionate and acetate in a 2:1 ratio (Figure S1). In addition to lactate, carboxylic acids and sugars that serve as growth substrates and are fermented to propionate and acetate include the following (propionate:acetate ratios are in parenthesis): fumarate (1.5:1), fructose (2:1), glucose (2:1), cellobiose (2:1), and mannitol (4:1). These ratios were determined during growth in the absence of yeast extract or H₂ in the initial headspace.

**Chromate reduction by strain HCF1 under fermentative conditions.** Anaerobic suspensions of lactate-grown cells of strain HCF1 reduced chromate in a minimal, anaerobic buffer containing 20 mM lactate as the electron donor and no electron acceptor (other than ~45 μM chromate)(Figure 1A). The initial, specific rate of Cr(VI) reduction was ~1.5 x 10⁻¹¹ μmol hr⁻¹ cell⁻¹. Chromate reduction was much slower in the absence of lactate than in its presence, and
was negligible in controls with killed cells and lactate (Figure 1A). Whereas Figure 1A depicts dissolved Cr(VI) concentrations based on the DPC assay, which is specific to Cr(VI), Figure 1B depicts total dissolved Cr concentrations as determined by ICP-MS. Comparison of the results shown in Figures 1A and 1B, which are based on aliquots from the same samples, makes it clear that strain HCF1 reduced Cr(VI) but that the reduced Cr did not precipitate, even though Cr(III) would be expected to form poorly soluble oxides or hydroxides at pH values > 5 (24). The bacterial production of reduced, soluble Cr has been observed previously, for example in _Pseudomonas stutzeri_ strain RCH2 (9), and may be attributable to formation of Cr(III)-organic complexes [e.g., Cr(III)-NAD⁺ complexes formed during Cr(VI) reduction by flavin reductases; (25, 26)] or to kinetic limitations on Cr(III) precipitation. Notably, the Cr(VI) and total Cr concentration profiles for the no-lactate control were similar to each other (Figure 1), suggesting that the small amount of Cr(VI) reduced under these conditions precipitated. It is not clear why reduced Cr precipitated more readily in the cell suspensions without lactate, but it is possible that a soluble Cr(III)-propionate complex formed in lactate-containing cultures. In the absence of lactate, the electron donor may have been hydrogen, which is typically present in the glove box atmosphere at 1 to 2%. Additional experiments suggested that H₂ could serve as an electron donor for Cr(VI) reduction, albeit not a very favorable one (Figure S2). Experiments were also conducted to test the possibility that soluble chemical reductants or redox-active proteins were released by strain HCF1 during metabolism of lactate that might cause extracellular Cr(VI) reduction. These cell-free experiments used filtrates of spent medium after growth of strain HCF1 with lactate; the filtrates were tested with and without heat denaturation under anaerobic conditions. In all cases, no Cr(VI) reduction was detected (data not shown), ruling out the role of extracellular chromate reduction by biogenic chemical reductants. These experiments did not
address the possibility of chromate-inducible extracellular enzymes, because the filtrates derived
from cells grown in the absence of chromate.

**Anaerobic Fe(III) reduction by strain HCF1.** Although strain HCF1 is primarily
characterized by its fermentative metabolism, it is capable of anaerobic Fe(III) reduction under
the tested growth conditions (data not shown) and in cell suspensions. Anaerobic suspensions of
lactate-grown cells of strain HCF1 reduced soluble Fe(III) in a minimal, anaerobic buffer
containing 20 mM lactate as the electron donor and Fe(III)-NTA as the electron acceptor (Figure
2). In the absence of cells or when cells were anaerobically autoclaved before resuspension,
Fe(III) reduction was negligible (Figure 2). Analogous to observations for Cr(VI) reduction
studies, a small amount of Fe(III) was reduced when cells were suspended in the absence of
lactate. Presumably, reducing equivalents were supplied by H₂ in the glove box atmosphere, as
suggested by cell suspension studies in which H₂ was supplied as the sole electron donor at a
range of concentrations (Figure S3). Clearly, H₂ is a more effective electron donor for Fe(III)
reduction than Cr(VI) reduction (Figures S2 and S3). The specific rate of Fe(III) reduction in the
presence of lactate was ~1.5 x 10⁻⁹ μmol hr⁻¹ cell⁻¹ [approximately 100-fold faster than the rate of
Cr(VI) reduction]. When solid-phase Fe(III), such as synthetic, amorphous Fe(OH)₃, was used
rather than Fe(III)-NTA, the Fe(III) reduction rate was orders of magnitude slower (data not
shown).

**Anaerobic nitrate reduction by strain HCF1.** Strain HCF1 is capable of reducing millimolar
quantities of nitrate when growing with lactate (Figure 3A). However, under these conditions,
nitrate reduction did not closely mirror lactate consumption; the most rapid lactate consumption
occurred between 24 and 36 hr whereas the most rapid nitrate reduction occurred between 36 and
48 hr (Figure 3A). Beginning at 36 hr, nitrite began to accumulate and propionate and acetate
production began to deviate from that observed during growth without nitrate (specifically, acetate production was greater and propionate production lower in the presence of nitrate, despite similar lactate consumption rates in the presence and absence of nitrate). Since most nitrite accumulation occurred after lactate was depleted, it is not clear what electron donor was driving nitrate reduction after 48 hr. Possibilities include lactate-derived formate and H₂, although these two compounds should also have been present during lactate fermentation. Indeed, cell suspension studies in which H₂ was the only electron donor supplied (90% H₂/10% CO₂ in the headspace) resulted in nitrate reduction to nitrite, whereas controls with no electron donor supplied (80% N₂/20% CO₂ in the headspace) did not generate nitrite (Figure S4). For the study represented in Figure 3, it is likely that nitrite was reduced further to ammonium, based upon two observations: (a) the amount of accumulated nitrite accounted for only approximately half of the nitrate reduced and (b) an ammonium-forming nitrite reductase (nrfA; two copies) is encoded in the genome (see below). More insight on nitrate/nitrite reduction was provided by targeted transcriptional analysis (reverse transcription quantitative Polymerase Chain Reaction, or RT-qPCR) of three putative nitrate or nitrite reductase genes conducted for samples collected at 36 and 72 hr in the presence and absence of nitrate (Figure 3B); these data are addressed in the section on c-type cytochromes below.

**Phylogeny of Pelosinus sp. strain HCF1.** Based on its 16S rRNA gene sequence, strain HCF1 belongs to the *Pelosinus-Sporotalea* group in the Firmicutes (Figure 4). It is most closely related to *Pelosinus fermentans* R7 (27) and *Sporotalea propionicum* TmPN3 (28), whose 16S rRNA gene sequences share 98.0% and 95.0% identity, respectively, with that of strain HCF1. Notably, two distinct 16S rRNA gene copies of differing lengths were amplified from strain HCF1 using universal forward (27F) and reverse (1492R) primers. The longer 16S rRNA gene
of strain HCF1 contains an additional ~100-bp insertion region at the 5’ end, whereas the rest of
the two sequences are identical. Similarly, it has been reported that two length-variable copies of
the 16S rRNA gene exist in Pelosinus sp. UFO1 and Sporotalea sp. TM1 (29), suggesting that
this is not an unusual phenomenon in the Pelosinus-Sporotalea group (Figure 4). Whether or not
the longer copy in the strain HCF1 genome is a pseudogene remains to be determined.

Overview of the Pelosinus sp. strain HCF1 genome. A draft sequence of the genome of
Pelosinus sp. strain HCF1 was obtained by paired-end sequencing on an Illumina GAIIx
platform. After quality control, the 100-bp paired-end reads were assembled into 214 contigs
using Velvet (20) and coverage of 48x with an N50 of 79.5 Kb and maximum contig length of
288 Kb. The genome size is estimated at 4.98 Mb with a GC content of 39.8 % (Table 1).

Lactate fermentation by strain HCF1 proceeds via the methylmalonyl-CoA pathway.
Analysis of the genome sequence of strain HCF1 indicates that it ferments lactate to propionate
via the methylmalonyl-CoA pathway. The proposed enzymatic reactions involved in lactate
fermentation by strain HCF1 are shown in Figure 5A. As proposed, lactate is converted to
pyruvate by lactate dehydrogenase (multiple copies are present in the genome). Pyruvate can be
converted to acetyl-CoA by pyruvate formate-lyase (PFL; two copies are present in the genome,
Hcf1DRAFT_00614 and Hcf1DRAFT_04611, with adjacent genes encoding a PFL activase).
In addition, pyruvate can be converted to acetyl-CoA by pyruvate:ferredoxin/flavodoxin
oxidoreductase (PFOR; Hcf1DRAFT_02505), which can generate reduced ferredoxin as an
electron donor for hydrogenases. Acetyl-CoA can be converted to acetate (with ATP generation
from substrate-level phosphorylation) via phosphotransacetylase (Pta; Hcf1DRAFT_00266) and
acetate kinase (Ack; Hcf1DRAFT_03144).
The pathway from pyruvate to propionate through methylmalonyl-CoA appears similar to that used by the close relative *Veillonella parvula* strain DSM 2008 (GenBank NC_013520); to illustrate, similar organization for selected genes in the methylmalonyl-CoA pathway in both organisms is shown in Figure 5B. However, strain HCF1 and *V. parvula* may differ in the enzymes catalyzing pyruvate carboxylation to oxaloacetate. For *V. parvula*, this is likely catalyzed by pyruvate carboxylase (Vpar_0752), but BLASTP searching (18) for this gene product did not result in any hits in the HCF1 genome. Instead, it is possible that pyruvate is converted to oxaloacetate by oxaloacetate decarboxylase [OAD; Hcf1DRAFT_02350 shares 50% sequence identity to the alpha subunit of OAD in *Vibrio cholera* (PDB accession number 2NX9)]. However, this same gene (Hcf1DRAFT_02350) was also tentatively annotated as pyruvate carboxylase subunit B and does not occur in a cluster with the beta and gamma subunits of OAD as does the alpha subunit of OAD in *Vibrio cholera*. Subsequent steps (Figure 5A) could be catalyzed by malate dehydrogenase (Hcf1DRAFT_00468), fumarase (e.g., Hcf1DRAFT_00270), and succinate dehydrogenase (Hcf1DRAFT_02750-2). Propionate formation is proposed to occur via propionyl-CoA:succinate CoA transferase [Hcf1DRAFT_02208; this shares 41% sequence identity with *E. coli* YgfH, which was demonstrated to have this activity (30)]. Other enzymes in the proposed pathway (Figure 5A) include methylmalonyl-CoA mutase (Hcf1DRAFT_00246-7 and Hcf1DRAFT_02209-10), methylmalonyl-CoA epimerase (Hcf1DRAFT_03696 and Hcf1DRAFT_03033, which share 97% sequence identity), and the Na⁺ ion-translocating methylmalonyl-CoA decarboxylase (the alpha subunit is likely Hcf1DRAFT_03697, which shares 79% sequence identity with Vpar_1244 in *V. parvula*; the beta subunit is likely encoded by Hcf1DRAFT_03584, Hcf1DRAFT_04139, and Hcf1DRAFT_01795, which collectively share from 55 to 79%
sequence identity with Vpar_1240). Note that the last enzyme, methylmalonyl-CoA decarboxylase, is not used by a better-studied lactate-fermenting bacterium, Propionibacterium freudenreichii, which instead uses methylmalonyl-CoA carboxytransferase to simultaneously decarboxylate (S)-methylmalonyl-CoA to propionyl-CoA and carboxylate pyruvate to oxaloacetate (31, 32). Many of the genes cited in this section appear likely to have been mis-annotated by automated annotation pipelines. Our re-annotation of some of these genes was based in part on gene context. As shown in Figure 5B, a number of the genes putatively encoding the methylmalonyl-CoA pathway in V. parvula and strain HCF1 are organized in similar gene clusters. Experimental validation of the functions of some of these genes is required for more definitive annotation.

[FeFe]- and [NiFe]-hydrogenases in strain HCF1. The genome of fermentative strain HCF1 encodes an assortment of [NiFe]- and [FeFe]-hydrogenases, which are proposed to catalyze H₂ oxidation and H₂ production, respectively. The two [NiFe]-hydrogenases are likely membrane-associated respiratory uptake (H₂-oxidizing) hydrogenases [classified as Group 1 hydrogenases by Vignais and co-workers (33)]. The structural genes of the two [NiFe]-hydrogenases are localized in clusters that include genes for the three subunits (i.e., large, small, and b-type cytochrome). One [NiFe]-hydrogenase includes the genes Hcf1DRAFT_03506-03510 and the other includes Hcf1DRAFT_02596-02598 (with some "adjacent" genes on opposite DNA strands). Genes encoding proteins required for [NiFe]-hydrogenase assembly and maturation were identified, including hypFCDE (Hcf1DRAFT_00423-00420), hypAB (Hcf1DRAFT_00123-00122), and Hcf1DRAFT_03507-03506.

There is evidence for at least four [FeFe]-hydrogenases encoded in the genome of strain HCF1. Hcf1DRAFT_00661 was annotated as a ferredoxin hydrogenase large subunit, and
appears to belong to the Group B1 monomeric [FeFe]-hydrogenases [as described by Calusinska and co-workers (34)]. Hcf1DRAFT_02066 was also annotated as a ferredoxin hydrogenase large subunit but shares only 21% amino acid sequence identity with Hcf1DRAFT_00661. Hcf1DRAFT_02066 is part of a gene cluster that includes an alpha subunit of formate dehydrogenase (Hcf1DRAFT_02068), as has been observed for [FeFe]-hydrogenases identified in *Eubacterium acidaminophilum* and a variety of *Clostridia* spp. (34, 35). Considering the co-location of [Fe-Fe]-hydrogenase and formate dehydrogenase genes, it is possible that the formate dehydrogenase interacts with the hydrogenase in a complex linking formate oxidation to H⁺ reduction.

Another putative [Fe-Fe]-hydrogenase is Hcf1DRAFT_01773, which occurs in a 3-gene cluster (Hcf1DRAFT_01771-01773). Considerable sequence similarity exists between these three genes and NuoEFG, which are subunits of Complex I (NADH:ubiquinone oxidoreductase); such similarity has been observed previously for certain [FeFe]-hydrogenases (33).

A fourth putative [Fe-Fe]-hydrogenase, Hcf1DRAFT_02349, appears to belong to the Group B2 monomeric [FeFe]-hydrogenases [as described by Calusinska and co-workers (34)]. Notably, an adjacent gene, Hcf1DRAFT_02350, is putatively pyruvate carboxylase or oxaloacetate decarboxylase, which is involved in lactate fermentation to propionate (as discussed earlier).

Transcriptional studies of strain HCF1 were undertaken to explore differential expression of structural genes representing the two [NiFe]-hydrogenases and four [FeFe]-hydrogenases under several anaerobic conditions: (a) H₂ (a 90% H₂/10 CO₂ mixture) as the sole electron donor, (b) lactate with no H₂ (80% N₂/20% CO₂ headspace), and (c) fructose with no H₂ (80% N₂/20% CO₂ headspace). Although definitive trends were not apparent to distinguish expression among
three conditions tested, one clear trend was that one [NiFe]-hydrogenase (represented by HCF1DRAFT_03510) was expressed at significantly higher levels than the other five hydrogenases tested (on average, ~8-fold higher, normalized to rpoB and averaged across all conditions examined) (Table S3).

**Putative chromate reductases and c-type cytochromes in strain HCF1.** Although many chromate-reducing bacterial species have been reported, only for a relatively small subset of these bacteria have the proteins catalyzing Cr(VI) reduction been identified and characterized biochemically, as reviewed elsewhere (36, 37). To date, the best-studied chromate reductase is ChrR, a soluble, dimeric, NADH-dependent flavoprotein reported in *Pseudomonas putida* (38, 39). BLASTP searches of the strain HCF1 genome were conducted for the following chromate reductases: ChrR from *P. putida* (GenBank AF375642); an Old Yellow Enzyme (OYE) homolog from *Thermus scotoductis* [(40); GenBank AM902709]; YieF, a ChrR homolog from *E. coli* [(38); GenBank AAK62985.2]; the Fre flavin reductase from *E. coli* [(25); EcoCyc EG10334]; FerB from *Paracoccus denitrificans* [(41, 42); putatively GenBank YP_917833.1]; and an azoreductase from *Bacillus* sp OY1-2 [(43); GenBank BAB13746.1]. The best match to any of these proteins was Hcf1DRAFT_04683, whose predicted protein sequence shared 51% identity with ChrR from *P. putida*. Alignment of these two proteins (Figure S5) revealed high conservation of the LFVTPEYNXXXXXXLKNAIDXXS motif in Hcf1DRAFT_04683 (with a leucine-to-isoleucine deviation at position 90), indicating that it is a member of the NADH_dh2 family of putative flavin-binding quinone reductases (39). Notably, Hcf1DRAFT_04683 also shared 42% sequence identity with the FerB flavoprotein, which has been shown to be effective at reducing Fe(III)-NTA as well as Cr(VI) (41, 42) and thus could be relevant to Fe(III) reduction in strain HCF1. The only other BLASTP match of note was Hcf1DRAFT_00464,
which shared 45% protein sequence identity with the NAD(P)H:flavin oxidoreductase (OYE homolog) from *T. scotoductis*.

Motivated in part by a desire to identify candidate c-type cytochromes that might be catalyzing Fe(III) or Cr(VI) reduction in strain HCF1, we searched all strain HCF1 ORFs for the CXXCH heme-binding motif that characterizes c-type cytochromes. Of the 37 ORFs found to contain one or more instances of the CXXCH motif, most were clearly not c-type cytochromes (including many Fe-S proteins, the molecular chaperone DnaJ, and ribosomal proteins L31 and L32, among others). A list of encoded proteins containing a CXXCH motif is shown in Table 2; all genes with annotations clearly indicating that they do not encode c-type cytochromes have been excluded from the table, leaving only 13 genes, most of which are annotated as "hypothetical proteins". There was no evidence of the multi-heme c-type cytochromes that have been reported to catalyze Fe(III) reduction in *Geobacter* and *Shewanella* species (44). Instead, the five encoded c-type cytochromes that could be confidently annotated are all related to the reduction of nitrogen oxides (nitrate, nitrite, or nitric oxide). Included in the HCF1 genome are two highly similar (sharing 58 to 68% amino acid sequence identity) pairs of c-type cytochrome genes encoding a nitrite reductase complex composed of NrfH and NrfA (formate-dependent nitrite reductase): Hcf1DRAFT_02324-02323 and Hcf1DRAFT_00451-00450. In both cases, these genes are clustered with upstream cytochrome c biosynthesis genes. The NrfHA nitrite reductase complex could play a key role in the dissimilatory reduction of nitrate to ammonium coupled to the oxidation of formate or H₂, as has been reported in certain proteobacteria, such as *Wolinella succinogenes* (45). Notably, the genome of strain HCF1 also encodes a membrane-bound, respiratory nitrate reductase (*narGHIJ*; Hcf1DRAFT_02301-02298), which could reduce nitrate to nitrite, thereby supplying nitrite to the NrfHA nitrite reductase complex. Thus, it is...
possible that formate- or H₂-dependent dissimilatory reduction of nitrate to ammonium catalyzed by NarG, NrfA, and NrfH (among other proteins) is responsible for the previously discussed observation of nitrate reduction occurring during growth of strain HCF1 with lactate (Figure 3A).

Transcriptional (RT-qPCR) analyses of narG and the two putative nrfH copies in cultures of strain HCF1 growing with lactate in the presence and absence of nitrate provide some insight on which nitrate and nitrite reductases may be involved. It is clear (Figure 3B) that both narG (Hcf1DRAFT_02301) and "nrfH2" (Hcf1DRAFT_02324) are nitrate-inducible, whereas "nrfH1" (Hcf1DRAFT_00451) is apparently not and is less likely to be catalyzing nitrite reduction under the conditions tested.

Finally, to complete the list of well-defined c-type cytochromes encoded in the HCF1 genome, we identified a membrane-anchored nitric oxide reductase, which includes the c-type cytochrome NorC (Hcf1DRAFT_04668) and the b-type cytochrome NorB (Hcf1DRAFT_04669). The presence of nitric oxide reductase in strain HCF1 is unexpected, as this enzyme is typically associated with denitrification, and the genome sequence does not indicate that strain HCF1 has this capability (i.e., we could not identify genes encoding nitric oxide-forming nitrite reductases, such as NirS or NirK, or the nitrous oxide reductase, NosZ).

BLASTp searches of the closely related fermenter Pelosinus sp. strain HCF1 in laboratory systems inoculated with aquifer material from the Hanford 100H site. Analysis by PCR amplification and pyrosequencing of SSU rRNA genes from fermenting column effluents demonstrated that a 16S rRNA gene taxon, identical to strain HCF1 over the region amplified (V4-V5 hypervariable region), was the dominant taxon, representing almost 80% of the pyrotag reads (Figure 6).
Environmental relevance of *Pelosinus* sp. strain HCF1. *Pelosinus* sp. strain HCF1 and close relatives have been reported as dominant microbial community members in experimental systems inoculated with Hanford 100H aquifer sediment (Figure 6) and groundwater (2) as well as laboratory (46) and field (5) aquifer systems associated with the Integrated Field Research Challenge site located at Oak Ridge National Laboratory (Tennessee). Considering its ability to become a prominent member of aquifer microbial communities and its diverse metabolic capabilities, *Pelosinus* sp. strain HCF1 (and close relatives) could play an important role in multiple biogeochemical cycles in aquifer environments. For example, in addition to catalyzing *in situ* Cr(VI) reduction under fermentative conditions in contaminated aquifers, it could potentially also reduce U(VI) (2, 47), nitrate (to nitrite and ammonium), and Fe(III). Although these reductive processes may not all be coupled to energy conservation in strain HCF1, they nonetheless could directly or indirectly mediate the mobility, toxicity, and bioavailability of several elements of environmental relevance. Strain HCF1 can also ferment a range of substrates relevant to both native and engineered conditions and could thereby release important metabolites for interspecies transfer, including acetate, propionate, H₂, and formate. Overall, the metabolic activities of strain HCF1 could mediate biogeochemical processes affecting the physicochemical and microbial components of its local aquifer environment.

Genomic and focused transcriptional analyses of strain HCF1 reported here have provided some insights into the underlying genetic basis of some of this organism's metabolic capabilities, including the methylmalonyl-CoA pathway and associated enzymes that catalyze the fermentation of lactate (and other substrates) to propionate and acetate, a variety of [NiFe]- and [FeFe]-hydrogenases that likely catalyze both H₂ production and consumption, flavoproteins that may catalyze Cr(VI) and Fe(III) reduction, and nitrate and nitrite reductases (NrfH, NrfA,
and NarGHI) that likely play a role in dissimilatory reduction of nitrate to ammonium. These nitrite reductases \((\text{nrf}H\text{ and nrf}A)\), along with nitric oxide reductase \((\text{nor}CB)\), are among the unexpected genes identified in the genome of strain HCF1 (e.g., BLASTP searches revealed no homologs in the \(V.\text{parvula}\) genome). To date, identification of \(\text{nrf}A\) in any firmicute genome is rare (as indicated by the FunGene database; http://fungene.cme.msu.edu; 6/7/2012 release). The metabolic context relevant to some of the genes identified in strain HCF1, such as nitric oxide reductase, is not currently understood. Understanding should improve as a result of future studies that will focus on characterizing the gene expression of strain HCF1 in an environmentally relevant context, such as in flow-through columns constructed with Hanford 100H aquifer sediment and eluted with synthetic groundwater that simulates \textit{in situ} conditions.

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FIGURE 1. Results of cell suspension assays to determine reduction of Cr(VI) by strain HCF1 under fermentative conditions with lactate (20 mM) as the electron donor. (A) Dissolved Cr(VI) concentrations as determined by the DPC assay. (B) Dissolved total Cr concentrations as determined by ICP-MS in aliquots of the same samples represented in panel A. Cell densities were approximately 2.2 x 10⁹ cells/mL. Note that the slightly lower time-zero concentrations for lactate-containing samples in panel A compared to panel B are likely due to a minor effect of lactate on the DPC assay (9).

FIGURE 2. Results of cell suspension assays to determine reduction of Fe(III)-NTA by strain HCF1 under fermentative conditions with lactate (20 mM) as the electron donor. Data points represent averages of duplicates and error bars represent 1 standard deviation. Cell densities were approximately 1.3 x 10⁹ cells/mL.

FIGURE 3. Lactate fermentation under anaerobic growth conditions in the presence and absence of nitrate. (A) Concentrations of lactate, acetate, and propionate (left-hand axis) and nitrate and nitrite (right-hand axis); nitrate and nitrite are shown only for nitrate-amended replicates. Data points represent averages of duplicates and error bars represent 1 standard deviation. Black arrows indicate the sampling times for transcriptional analysis. (B) Transcript copy number (normalized to rpoB) determined by RT-qPCR for three genes: narG (Hcf1DRAFT_02301), nrfH-copy 1 (Hcf1DRAFT_00451), nrfH-copy 2 (Hcf1DRAFT_02324) in biological duplicates with and without nitrate (represented in panel A). Bar heights represent averages (n = 5 or 6, including technical replicates). Relative standard deviations averaged 24% and ranged from 13 to 46% (Table S4).

FIGURE 4. Maximum-likelihood phylogenetic tree of Pelosinus sp. strain HCF1 and representative Firmicutes. The bar represents distances calculated as described in Materials and Methods. Numbers at internal nodes are the percentage of 100 bootstrap samples in which the organisms to the right of the node were monophyletic.
FIGURE 5. (A) Proposed pathway for lactate fermentation to acetate and propionate in *Pelosinus* sp. strain HCF1 based on genomic analysis (see text). Abbreviations: Pta, phosphotransacetylase; Ack, acetate kinase; PFL, pyruvate formate-lyase; PFOR, pyruvate:ferredoxin/flavodoxin oxidoreductase; DH, dehydrogenase; CoA, coenzyme A. (B) Genomic organization of selected genes putatively involved in the methylmalonyl-CoA pathway in strain HCF1 and *Veillonella parvula*. Note that PCR finishing was used to establish that Hcf1DRAFT_2210 and Hcf1DRAFT_3695 are contiguous in the genome.

FIGURE 6. Phylogenetic composition of flow-through column microbial community from which strain HCF1 was isolated, based upon SSU rRNA gene pyrotag sequence analysis.
TABLE 1. *Pelosinus* sp. strain HCF1 sequencing and draft genome statistics

<table>
<thead>
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<th>Platform</th>
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<td>Estimated genome size</td>
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<tr>
<td>N50</td>
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<td>G+C percentage</td>
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<tr>
<th>Genes, total number</th>
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<th>Percentage</th>
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<tr>
<td>Protein coding genes</td>
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<tr>
<td>RNA genes</td>
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<td>2.46%</td>
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<tr>
<td>Protein coding genes with function prediction</td>
<td>3776</td>
<td>77.94%</td>
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| CRISPR Count | 2 |
TABLE 2. CDS potentially encoding c-type cytochromes in the *Pelosinus* strain HCF1 genome

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<td>Uncharacterized conserved protein</td>
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<td>Formate-dependent nitrite reductase, periplasmic cytochrome c_552 subunit (EC:1.7.2.2), NrfA</td>
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<td>5</td>
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<td>Putative nitric oxide reductase, NorC</td>
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740
741  a  As defined by the presence of at least one CXXCH heme-binding motif but excluding proteins that are clearly not c-type cytochromes (see text). For NrfA, one additional heme is represented by a CXXCK motif (45).
744
745  b  Best attempt at annotation based on examination of best BLASTP hits and genomic context.
747
748  c  Molecular mass predicted for the unprocessed gene product without cofactors.
Figure 1
Figure 2
Figure 3
Figure 4
Figure 5
Figure 6
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