Adaptation to Disruption of the Electron Transfer Pathway for Fe(III) Reduction in *Geobacter sulfurreducens*

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Previous studies demonstrated that an outer membrane c-type cytochrome, OmcB, was involved in Fe(III) reduction in Geobacter sulfurreducens. An OmcB-deficient mutant was greatly impaired in its ability to reduce both soluble and insoluble Fe(III). Reintroducing omcB restored the capacity for Fe(III) reduction at a level proportional to the level of OmcB production. Here, we report that the OmcB-deficient mutant gradually adapted to grow on soluble Fe(III) but not insoluble Fe(III). The adapted OmcB-deficient mutant reduced soluble Fe(III) at a rate comparable to that of the wild type, but the cell yield of the mutant was only ca. 60% of that of the wild type under steady-state culturing conditions. Analysis of proteins and transcript levels demonstrated that expression of several membrane-associated cytochromes was higher in the adapted mutant than in the wild type. Further comparison of transcript levels during steady-state growth on Fe(III) citrate with a whole-genome DNA microarray revealed a significant shift in gene expression in an apparent attempt to adapt metabolism to the impaired electron transport to Fe(III). These results demonstrate that, although there are many other membrane-bound c-type cytochromes in G. sulfurreducens, increased expression of these cytochromes cannot completely compensate for the loss of OmcB. The concept that outer membrane cytochromes are promiscuous reductases that are interchangeable in function appears to be incorrect. Furthermore, the results indicate that there may be different mechanisms for electron transfer to soluble Fe(III) and insoluble Fe(III) oxides in G. sulfurreducens, which emphasizes the importance of studying electron transport to the environmentally relevant Fe(III) oxides.

Microorganisms in the family of Geobacteraceae are the predominant Fe(III) reducers in many subsurface environments where dissimilatory Fe(III) reduction plays an important role in the degradation of organic matter or the bioremediation of organic or metal contaminants (12, 13). The mechanisms by which Fe(III)-reducing microorganisms reduce Fe(III) are poorly understood. It is known that phylogenetically distinct Fe(III) reducers reduce Fe(III) oxide with significantly different mechanisms. For example, Geothrix and Shewanella species produce soluble electron shuttles and/or Fe(III)-chelating compounds to alleviate the need for direct electron transfer from the cell surface to the surface of Fe(III) oxide (20–22, 26). However, current evidence suggests that Geobacter species require direct physical contact between cells and extracellular electron acceptors (19), which may be achieved with appendages such as flagella and pili (3).

Earlier biochemical studies suggested that *c*-type cytochromes are likely to be involved in Fe(III) reduction in *Geobacter* species (8, 9, 16). Further genetic studies, concentrating on *Geobacter sulfurreducens*, due to the availability of a genetic system (5) and the genome sequence (17), indicated that a periplasmic cytochrome, PpcA (11); an inner membrane-associated cytochrome, MacA (2); and an outer membrane cytochrome, OmcB (10), are important for Fe(III) reduction. Based on location, PpcA and MacA were proposed to be intermediary electron transfer components whereas the outer membrane localization of OmcB and the severe impact on Fe(III) reduction when *omcB* was deleted suggested that OmcB might be the terminal Fe(III) reductase (10).

However, sequencing of the *G. sulfureducens* genome has demonstrated that this organism contains genes for over 100 putative *c*-type cytochromes (17). A potential reason for this unprecedented number of cytochromes is that this might provide the opportunity for *G. sulfureducens* to form multiple routes for electron transfer to Fe(III) and thus better maximize rates of Fe(III) reduction and/or provide a high degree of flexibility to adapt to disruptions in electron transfer pathways. In order to investigate this potential for adaptation further, the long-term adaptation of the OmcB-deficient mutant was studied.

MATERIALS AND METHODS

Bacterial strains and culturing conditions. Geobacter sulfurreducens strain DL1 (5) and DL6 (omcB::cam) (10) were routinely cultured anaerobically in NBAF (acetate/fumarate) or FWFC [acetate/Fe(III) citrate] medium at 30°C as previously described (5). Steady-state growth was investigated in chemostats under strict anaerobic conditions (N₂-CO₂ [80:20, vol/vol]) in a freshwater medium containing Fe(III) citrate (56 mM) as the electron acceptor and limited acetate (5.5 mM) as the electron donor as previously described (4, 6). The dilution rate was 0.05 h⁻¹.

Detection of cytochromes in the membrane fraction. The membrane fractions of *G. sulfurreducens* were isolated as described earlier (10). Proteins (50 μ g) of membrane fractions were analyzed with Tris-Tricine denaturing polyacrylamide gel electrophoresis (1). *c*-type cytochromes were detected by staining with *N*,*N*,*N'*,*N'*-tetramethylbenzidine as previously described (7, 25). Molecular standard markers were purchased from Bio-Rad Laboratories (Hercules, CA) and were stained separately with Coomassie blue R-250.

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Target gene Primer		Sequences (5'-3')	Length (bp)	Reference(s)	
GSU0594	994F	CCACAGCAGTAAGGATATCG	375	This work	
	994R	GCATCGATATGCAGATCCG			
GSU2494	4120F	GCACTCAAGTTCAAGCTCAAG	469	This work	
	4120R	GGATGATATCGCTCTTCAGC			
GSU2495	4123F	GCTCATTCTTCACCTGTCGG	574	This work	
	4123R	GATAGTGCAGCGCTTGCTTTC			
GSU2503 (omcT)	4140-1F	CCAACCAGTTCAGCTGCATC	382	This work	
	4140-1R	GAAGGGGCCAAGGTTCTGATC			
GSU2504 (omcS)	4142F	ACGTTCGTGGTCTCAACAC	775	This work	
~ /	4142R	GATGGTCGTGAACTCGTATG			
GSU2731 (omcC)	8914	GCCAGAGTGAGGCCCAGA	543	This work; 4	
	8915	GGGTGTTGTGGTAGAAGGG			
GSU2737 (omcB)	8916	GGACTGCGCACCATCAAGG	435	This work; 10	
	8908-2	GGTCAGCAGGCCACCGG			
GSU2808	4655F	GGTGACAGTAGGAGTACCTG	354	This work	
	4655R	GGTGTGACAGAGGTAGCAGAC			
GSU2811	4659F	GTGTGACCAGCTATGCTCC	395	This work	
	4659R	GTCAAAGTCCAGGTCGAACC			
GSU2813	4662F	CCATCCTCTGTGCAGTCGC	599	This work	
	4662R	GGTCTGTTTTCCGTCAAGG			
GSU2887	4789-1F	CCATACGAGCACCAATGAGC	445	This work	
	4789-1R	GCTGTTGCTGTCAGAGGAGG			
GSU3259	5426F	CGTCTATCCTTGGCTTCGAG	426	This work	
	5426R	CGATATGCCAGTGGATACC			

TABLE 1. Primers used for Northern blot analyses for *c*-type cytochrome genes whose expression levels were significantly altered in the Fe(III)-adapted OmcB-deficient mutant

Analytical techniques. Protein concentration was determined with the bicinchoninic acid method with bovine serum albumin as a standard (24). Cells were counted using epifluorescence microscopy with acridine orange staining (15). Fe(II) concentrations were determined with the ferrozine assay as previously described (14). Acetate concentrations were measured with high-pressure liquid chromatography on a Hewlett-Packard series 1100 (Agilent Technologies, Inc., Albany, NY) with a Bio-Rad Aminex HPX-87H column (300 × 7.8 mm) and a mobile phase of 8 mM H₂SO₄.

DNA microarray hybridization experiments and data analysis. Total RNA was isolated from two sets of identically treated chemostat cultures of both the wild type and the adapted OmcB-deficient mutant. A total of 11 replicate hybridizations were carried out: five replicate hybridizations were from one set of the biological samples, and six were from the other set. DNA microarray hybridization and data analyses were described previously (18). Briefly, total RNA (5 μ g) that was isolated from steady-state continuous cultures of the wild type and the adapted OmcB-deficient mutant was used to synthesize cDNA labeled with Cy3-Cy5 fluorescent dyes, which was hybridized to gene arrays. The signal from each spot in the arrays served as a measure of the expression level of each gene and was used to calculate the expression ratio between the wild type and the mutant.

Quantitative reverse transcription-PCR (RT-PCR). For cDNA synthesis, SuperScript III RNase H⁻ reverse transcriptase (Invitrogen Co., Carlsbad, CA) was used according to the manufacturer's instructions with gene-specific antisense primers 8908-2 (10) and 8915 (4) for *omcB* and *omcC*, respectively.

The quantitative real-time PCR was carried out as previously described (4). Two specific primer sets, 8912/8908-2 and 8917/8915 (4), were used to determine levels of mRNA for *omcB* and *omcC*, respectively. The temperature profile was composed of an initial incubation step for 2 min at 50°C (activation of the polymerase) followed by a 10-min denaturation step at 95°C, 40 cycles of denaturation for 45 s at 95°C, annealing for 1 min at 58°C, elongation for 1 min at 72°C, and a final elongation step for 6 min at 72°C.

DNA and RNA manipulations. PCR product purification was carried out using QIAGEN PCR purification kits (QIAGEN Inc., Valencia, CA). Probes for Northern blot analysis were labeled with $[\alpha^{-32}P]dATP$ using a Strip-EZ DNA probe synthesis and removal kit (Ambion Inc., Austin, Texas). $[\alpha^{-32}P]dATP$ was purchased from PerkinElmer Life and Analytical Sciences, Inc., Boston, MA. All primers used to amplify *G. sulfurreducens* sequences were designed using the *G. sulfureducens* genome sequence (17). QIAGEN *Taq* DNA polymerase (QIA-GEN Inc., Valencia, CA) was used for all PCR amplifications.

Total RNA was isolated from mid-log cultures using RNeasy Midi kits (QIA-GEN Inc., Valencia, CA) followed by treatment with RNase-free DNase (Ambion Inc., Austin, Texas). Northern blot analyses were carried out with the NorthernMax-Gly system (Ambion Inc., Austin, Texas) according to the manufacturer's instructions. All probes for Northern blot analyses were purified PCR product. Primers for amplifying probes are listed in Table 1. The PCR amplification program was as follows: 96°C for 40 s followed by 25 cycles of 96°C for 40 s, 58°C for 1 min, and 72°C for 1 min and a final extension at 72°C for 10 min.

RESULTS

Adaptation of OmcB-deficient mutant (DL6) to grow on soluble Fe(III). As previously reported (10), the OmcB-deficient mutant did not reduce Fe(III) citrate for periods of time well beyond those in which the wild-type cells had reduced all the Fe(III) (Fig. 1). However, after incubation for more than a



FIG. 1. Adaptation of OmcB-deficient mutant (DL6) to grow with Fe(III) citrate as the sole electron acceptor. Mid-log (optical density at 600 nm, \sim 0.3) fumarate-grown cells served as the inoculum. Symbols: filled or empty squares, wild-type Fe(II) or cell numbers, respectively; shaded or empty circles, OmcB-deficient mutant Fe(II) or cell numbers, respectively. Data are means ± standard deviations of triplicates.



FIG. 2. Growth of the OmcB-deficient mutant after adaptation and two additional transfers on Fe(III) citrate medium. Symbols: filled or empty squares, wild-type Fe(II) or cell numbers, respectively; shaded or empty circles, OmcB-deficient mutant Fe(II) or cell numbers, respectively. Data are means \pm standard deviations of triplicates.

week the OmcB-deficient mutant began to reduce Fe(III) (Fig. 1). The growth rate (doubling time, 35 h) was lower than that of the wild type (6 h), and the number of cells produced was only 38% of that of the wild type. The OmcB-deficient mutant never adapted to grow with Fe(III) oxide as the terminal electron acceptor (data not shown).

With two successive transfers of the adapted OmcB-deficient cells in Fe(III) citrate medium, the rate of Fe(III) reduction approached that of the wild type (Fig. 2). However, the final cell yield of OmcB-deficient mutant remained less than half of that of the wild type. The same pattern of adaptation was observed repeatedly when fumarate-grown cultures of the OmcB-deficient mutant were inoculated into Fe(III) citrate medium, suggesting that the adaptation was not due to spontaneous mutations; instead the growth may have resulted from a subpopulation of phase variants or cells with epigenetic modifications. Furthermore, when membrane fractions were isolated from fumarate or Fe(III) citrate-adapted cultures OmcB was not detected (Fig. 3, lanes 3 and 4). However, at least six other cytochromes, including OmcC, which is a close relative



FIG. 3. Tricine-polyacrylamide gel electrophoresis and heme staining of membrane fractions prepared from wild type (WT) and the OmcB-deficient mutant. Membrane fractions were prepared from cultures grown with fumarate (lanes 1 and 3) or Fe(III) citrate (lanes 2 and 4). OmcB and OmcC are indicated with arrows on the left side of the gel. Other cytochromes which were highly expressed in the adapted Fe(III)-grown OmcB-deficient mutant are indicated with arrows on the right side of the gel.

of OmcB but is not required for Fe(III) reduction (10), were present in higher concentrations in the OmcB-deficient mutant than in wild-type cells when membrane fractions were isolated from Fe(III) citrate cultures (Fig. 3, lane 4). The higher concentration of OmcC in the adapted OmcB-deficient mutant was consistent with the finding that levels of *omcC* transcripts as determined with quantitative RT-PCR were ca. 10^3 times higher in adapted, Fe(III)-grown cells than in the wild type.

Transcriptional profile comparison of the OmcB-deficient mutant and the wild type. To further investigate potential differences in gene expression in the adapted OmcB-deficient mutant, transcript levels in continuous cultures of the adapted OmcB-deficient mutant grown with Fe(III) citrate as the electron acceptor were compared to those in wild-type cells with a whole-genome microarray (18). Although the OmcB-deficient mutant reduced Fe(III) at a rate comparable to that of the wild type versus mutant, 36.5 ± 1.2 mM versus 35.6 ± 2.9 mM, respectively; mean \pm standard deviation; n = 3], the steady-state cell yield of the mutant was only 61% of that of the wild type (total protein of wild type versus mutant, 0.036 ± 0.003 versus 0.022 ± 0.007 mg/ml, respectively).

The adaptation of the OmcB-deficient mutant to grow on Fe(III) citrate was associated with changes in expression of a variety of genes involved in electron transfer and metabolism (Tables 2 and 3). A total of 83 genes had higher transcript levels in the OmcB-deficient mutant than in the wild type (Table 2), and 88 genes appeared to have lower transcript levels in the mutant (Table 3). Eighty-six genes had transcript levels that changed more than twofold in the OmcB-deficient mutant, with 47 genes up-regulated and 39 genes down-regulated.

The greatest increase in transcript levels was for GSU2504 and GSU2503 (Tables 2 and 4), which are cotranscribed and are predicted to encode two outer membrane c-type cytochromes, OmcS and OmcT, with high similarity to each other (D. Holmes et al., submitted for publication; T. Mehta et al., submitted for publication). OmcS, the cytochrome encoded by GSU2504, is readily sheared off from the outer surface of G. sulfurreducens and is involved in insoluble Fe(III) reduction and electricity production (T. Mehta, submitted for publication). The protein encoded by GSU2503, OmcT, has yet to be detected in any proteomic analysis of G. sulfurreducens (R. Ding, unpublished data). Transcripts of omcS and omcT were detected in wild-type cells during growth on insoluble Fe(III) oxide but not on soluble Fe(III) citrate (T. Mehta et al., submitted for publication). However, transcripts of both omcS and omcT were detected in the adapted OmcB-deficient mutant during growth on Fe(III) citrate (Table 2; see also Fig. 5). Thus, it is likely that the heme-staining protein recovered in higher levels from the adapted OmcB-deficient mutant (Fig. 3) with a molecular mass of ca. 45 kDa was OmcS.

Genes with significant changes in their transcript levels could be categorized into 16 different groups according to their functions (Fig. 4). The largest functional groups that had higher levels of transcripts in the OmcB-deficient mutant than in the wild type were those related to energy metabolism and genes for hypothetical proteins. Other than *omcS* and *omcT*, the other genes related to energy metabolism which had transcript levels over twofold higher in the adapted mutant were

Locus ID ^a	Common name	Role category	Log_2 ratio \pm SD
GSU0944 GSU0945 GSU1906	Cystathionine beta-lyase Cystathionine beta-lyase 2-Isopropylmalate synthase	Amino acid biosynthesis	$\begin{array}{c} 0.884 \pm 0.059 \\ 0.577 \pm 0.095 \\ 0.504 \pm 0.125 \end{array}$
GSU0265 GSU2078 GSU2497 GSU2498 GSU2526	Putative membrane protein Rod shape-determining protein Putative lipoprotein Putative lipoprotein Putative membrane protein	Cell envelope	$\begin{array}{c} 0.855 \pm 0.114 \\ 1.185 \pm 0.264 \\ 1.618 \pm 0.170 \\ 0.964 \pm 0.095 \\ 0.622 \pm 0.121 \end{array}$
GSU1140 GSU1899 GSU2236	Methyl-accepting chemotaxis protein Virulence factor Mce family protein GTP pyrophosphokinase	Cellular processes	$\begin{array}{c} 0.873 \pm 0.193 \\ 0.625 \pm 0.145 \\ 0.947 \pm 0.195 \end{array}$
GSU2502 GSU2537	Spermine/spermidine synthase family protein Biosynthetic arginine decarboxylase	Central intermediary metabolism	$\begin{array}{c} 0.723 \pm 0.121 \\ 0.638 \pm 0.079 \end{array}$
GSU3245	Putative DNA polymerase II	DNA metabolism	2.133 ± 0.199
GSU0193 GSU0594 GSU0782 GSU0783	L-Sorbosone dehydrogenase Cytochrome <i>c</i> Nickel-dependent hydrogenase, small subunit Nickel-dependent hydrogenase, iron-sulfur cluster-binding	Energy metabolism	$\begin{array}{c} 0.716 \pm 0.111 \\ 1.133 \pm 0.271 \\ 2.165 \pm 0.250 \\ 2.547 \pm 0.370 \end{array}$
GSU0784 GSU0785 GSU1640 GSU1641 GSU1707 GSU1722 GSU1761 GSU2018 GSU2098 GSU2494 GSU2494 GSU2501 GSU2501 GSU2504 GSU2811 GSU2812 GSU2813 GSU2246	protein Nickel-dependent hydrogenase, membrane protein Nickel-dependent hydrogenase, large subunit Cytochrome <i>d</i> ubiquinol oxidase, subunit I Cytochrome <i>d</i> ubiquinol oxidase, subunit II Group II decarboxylase Creatinine amidohydrolase Cytochrome <i>c</i> Glycine cleavage system H protein Carbon monoxide dehydrogenase subunit Cytochrome <i>c</i> Cytochrome <i>c</i> Thioredoxin family protein Cytochrome <i>c</i> 551 peroxidase Thioredoxin peroxidase		$\begin{array}{c} 2.605 \pm 0.193 \\ 2.416 \pm 0.306 \\ 1.779 \pm 0.776 \\ 2.508 \pm 0.427 \\ 0.664 \pm 0.052 \\ 0.570 \pm 0.103 \\ 0.652 \pm 0.164 \\ 0.575 \pm 0.112 \\ 1.782 \pm 0.286 \\ 1.863 \pm 0.388 \\ 1.810 \pm 0.294 \\ 0.870 \pm 0.204 \\ 3.340 \pm 0.154 \\ 3.676 \pm 0.626 \\ 0.745 \pm 0.112 \\ 0.815 \pm 0.124 \\ 0.839 \pm 0.186 \\ 1.049 \pm 0.229 \end{array}$
GSU1916	Phosphatidate cytidylyltransferase	Fatty acid and phospholipid metabolism	0.556 ± 0.146
GSU0192 GSU0208 GSU0593 GSU0647 GSU0788 GSU0919 GSU1160 GSU1309 GSU1770 GSU1947 GSU1947 GSU1948 GSU2496 GSU2499 GSU2500 GSU3310 GSU3310 GSU3403 GSU3400 GSU3410	Hypothetical protein Hypothetical protein	Hypothetical proteins	$\begin{array}{c} 1.433 \pm 0.203 \\ 1.029 \pm 0.354 \\ 1.083 \pm 0.217 \\ 1.002 \pm 0.139 \\ 1.453 \pm 0.329 \\ 1.587 \pm 0.326 \\ 1.220 \pm 0.134 \\ 0.701 \pm 0.216 \\ 0.762 \pm 0.117 \\ 1.464 \pm 0.098 \\ 1.168 \pm 0.162 \\ 1.418 \pm 0.374 \\ 1.998 \pm 0.238 \\ 1.792 \pm 0.532 \\ 0.664 \pm 0.091 \\ 0.670 \pm 0.160 \\ 2.586 \pm 0.168 \\ 1.248 \pm 0.215 \\ 1.588 \pm 0.170 \end{array}$

TABLE 2. Genes that were significantly up-regulated in the adapted OmcB-deficient mutant compared to the wild type grown in acetatelimiting Fe(III) citrate continuous culture

Continued on following page

Locus ID ^a	Common name	Role category	Log_2 ratio \pm SD
GSU0786	Hydrogenase maturation protease	Protein fate	1.894 ± 0.178
GSU0787	TatA/E family twin-arginine translocation protein		0.820 ± 0.155
GSU1159	Ptp1 family intracellular protease		1.185 ± 0.245
GSU0646	tRNA (guanine-N ₁)-methyltransferase	Protein synthesis	0.607 ± 0.129
GSU0648	Ribosomal protein L19		0.788 ± 0.185
GSU0537	Sensory box/GGDEF family protein	Regulatory functions	1.138 ± 0.081
GSU2506	Sigma-54-dependent DNA-binding response regulator		2.264 ± 0.345
GSU2507	Sensor histidine kinase		2.797 ± 0.432
GSU3387	AraC/XylS family transcriptional regulator		0.561 ± 0.146
GSU0189	ATP-dependent RNA helicase	Transcription	1.097 ± 0.139
GSU1161	RND family efflux transporter	Transport and binding proteins	0.949 ± 0.198
GSU1346	Sulfate ABC transporter, periplasmic sulfate-binding protein		1.047 ± 0.224
GSU1723	Mechanosensitive ion channel family protein		0.812 ± 0.177
GSU3401	Branched-chain amino acid ABC transporter, periplasmic amino acid-binding protein		2.590 ± 0.255
GSU3404	Amino acid ABC transporter, ATP-binding protein		2.452 ± 0.341
GSU3405	Amino acid ABC transporter, permease protein		0.973 ± 0.187
GSU0500	GTP-binding protein TypA	Unknown functions	0.610 ± 0.086
GSU0534	RrF2 family protein		0.814 ± 0.248
GSU0664	GTP binding protein YchF		0.546 ± 0.091
GSU0769	RarD protein		1.262 ± 0.177
GSU1643	GGDEF/response regulator receiver domain protein		2.053 ± 0.162
GSU1708	Atz/Trz family chlorohydrolase		0.599 ± 0.098
GSU1877	2-Nitropropane dioxygenase family oxidoreductase		1.059 ± 0.287
GSU1945	Fibronectin type III domain protein		1.553 ± 0.177
GSU2493	NHL repeat domain protein		1.656 ± 0.166
GSU2505	NHL repeat domain protein		2.102 ± 0.573
GSU2527	Nitrate/sulfite reductase domain protein		1.200 ± 0.296

TABLE 2—Continued

^a ID, identification.

those associated with a nickel-dependent hydrogenase, a cytochrome d ubiquinol oxidase, and a carbon monoxide dehydrogenase (Table 2). The transcript of a relA homolog (GSU2236), whose protein product is predicted to be involved in a stringent response during stress or nutrient depletion, was ca. twofold up-regulated in the mutant. The largest functional groups that had lower levels of transcripts in the OmcB-deficient mutant than in the wild type were those coding for hypothetical proteins, mobile and extrachromosomal element functions, and transport and binding proteins (Fig. 4). Four c-type cytochrome genes including the omcB gene were among the genes with lower transcript levels than those of the wild type (Tables 3 and 4). Genes coding for acetate metabolism (acetate kinase, GSU2707, and phosphate acetyltransferase, GSU2706) were among the groups of genes with over twofold-lower transcript levels.

Validation by Northern blotting of expression ratios from transcriptional profiling. Transcript levels for cytochrome genes which were significantly different (ca. twofold) between the adapted mutant and the wild type from the whole-genome microarray analysis were evaluated with Northern blot analyses (Fig. 5). The results from the Northern analysis compared well with those from the microarray analysis. The only exception was that *omcC* was not detected as up-regulated in the microarray, but transcript levels of *omcC* were significantly higher in the Northern analysis, consistent with quantitative real-time RT-PCR results mentioned above. The predicted molecular weights of those cytochromes whose transcript levels were significantly higher (ca. twofold) than those of the wild type from both microarray and Northern analyses (Table 4) are in agreement with those protein bands that were identified from the membrane fractions of the OmcB-deficient mutant (Fig. 3, lane 4).

DISCUSSION

The results demonstrate that even though *G. sulfurreducens* has genes for ca. 30 outer membrane *c*-type cytochromes, including the closely related OmcC, it cannot fully adapt to the loss of OmcB, which is essential for the most effective growth on soluble Fe(III) as well as for any growth on Fe(III) oxide, the most abundant form of Fe(III) in most environments. As detailed below, these results suggest that outer membrane cytochromes, which are frequently viewed as rather promiscuous agents for electron transfer to a variety of extracellular electron acceptors, may in fact have quite specific functions that cannot be replicated by other cytochromes.

Adaptation of OmcB-deficient mutant for growth on soluble Fe(III). Previous short-term studies identified OmcB as an important component in electron transfer based on the finding that deleting *omcB* inhibited Fe(III) reduction and reintroducing *omcB* restored the capacity for Fe(III) reduction at a level

TABLE 3.	Genes that were significantly	down-regulated in the adapted	l OmcB-deficient	mutant compared to	o the wild type in acetate-limitin	ıg
		Fe(III) citrate-grown	continuous cultu	ire		

Locus ID ^a	Common name	Role category	Log_2 ratio \pm SD
GSU0656	Branched-chain amino acid aminotransferase	Amino acid biosynthesis	-0.765 ± 0.179
GSU0862	FolD bifunctional protein	Biosynthesis of cofactors, prosthetic groups and	-0.881 ± 0.176
GSU1577	Cob(I) alamin adenosyltransferase	carriers	-0.635 ± 0.087
GSU0967	Putative membrane protein	Cell envelope	-1.843 ± 0.242
GSU1010	Slt family transglycosylase	een envelope	-0.809 ± 0.186
GSU1493	Type IV pilus biogenesis protein PilC		-0.692 ± 0.128
GSU1013	Putative chemotaxis MotB protein	Cellular processes	-0.673 ± 0.191
GSU2707	Acetate kinase	Central intermediary metabolism	-1.150 ± 0.162
GSU0096	Recombination protein RecR	DNA metabolism	-0.896 ± 0.098
GSU0338	NADH dehydrogenase I; A subunit	Energy metabolism	-0.980 ± 0.156
GSU1538	Putative methylamine utilization protein MauG		-1.708 ± 0.244
GSU2612	Putative rubrerythrin/rubredoxin protein		-0.765 ± 0.098
GSU2706	Phosphate acetyltransferase		-1.189 ± 0.271
GSU2737	Cytochrome c (OmcB)		-0.940 ± 0.152
GSU2808	Cytochrome c		-0.730 ± 0.132
GSU2887	Cytochrome c		-1515 ± 0.208
GSU3250	Cytochrome c		-1.069 ± 0.150
GSU3444	NADH dehydrogenase I B/C/D subunits		-0.669 ± 0.136
GSU3061	Squalene-hopene cyclase	Fatty acids and phospholipid metabolism	-1.180 ± 0.349
GSU0384	Hypothetical protein	Hypothetical proteins	-0.785 ± 0.190
GSU0712	Hypothetical protein		-0.867 ± 0.115
GSU0875	Hypothetical protein		-1.037 ± 0.259
GSU0964	Hypothetical protein		-1.460 ± 0.196
GSU0965	Hypothetical protein		-1.557 ± 0.543
GSU0966	Hypothetical protein		-1.731 ± 0.304
GSU0976	Hypothetical protein		-0.858 ± 0.254
GSU0977	Hypothetical protein		-0.912 ± 0.254
GSU0980	Hypothetical protein		-0.592 ± 0.176
GSU0981	Hypothetical protein		-0.847 ± 0.231
GSU0982	Hypothetical protein		-0.824 ± 0.231
GSU0088	Hypothetical protein		-0.652 ± 0.201
GSU0900	Hypothetical protein		-1.066 ± 0.291
GSU0990	Hypothetical protein		-1.000 ± 0.290 -0.507 ± 0.116
CSU1220	Insponetical protein		-0.397 ± 0.110 1 274 ± 0.412
GSU1559	Hypothetical protein		-1.574 ± 0.412 1 212 ± 0.194
GSU1945	Hypothetical protein		-1.212 ± 0.184
GSU1994	Hypothetical protein		$-1.6/5 \pm 0.211$
GSU2353	Hypothetical protein		-1.264 ± 0.296
GSU2/80	Hypothetical protein		-0.902 ± 0.223
GSU3079	Hypothetical protein		-0.709 ± 0.171
GSU3080	Hypothetical protein		-1.280 ± 0.150
GSU3081	Hypothetical protein		-1.051 ± 0.198
GSU3084 GSU3085	Hypothetical protein Hypothetical protein		-1.344 ± 0.144 -1.227 ± 0.246
	JI TOTAL		
GSU0555	ISGsu7, transposase OrfA	Mobile and extrachromosomal element functions	-0.785 ± 0.188
GSU0556	ISGsu7, transposase OrfB		-1.069 ± 0.343
GSU0761	ISGsu7, transposase OrfB		-0.844 ± 0.289
GSU0762	ISGsu7, transposase OrfA		-0.742 ± 0.173
GSU0975	Phage tail sheath protein		-0.866 ± 0.283
GSU0986	Tail lysozyme		-0.767 ± 0.097
GSU1355	ISGsu7, transposase OrfA		-0.882 ± 0.222
GSU1356	ISGsu7, transposase OrfB		-1.029 ± 0.366
GSU1847	ISGsu7, transposase OrfA		-0.856 ± 0.212
GSU1848	ISGsu7, transposase OrfB		-1.057 ± 0.327
GSU2127	ISGsu7, transposase OrfA		-0.784 ± 0.204
GSU2128	ISGsu7, transposase OrfB		-0.969 ± 0.204
GSU2139	ISGsu7, transposase OrfA		-0.734 ± 0.198

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FABLE	3—	Continued
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GSU2140 ISGsu' GSU2171 ISGsu' GSU2279 ISGsu' GSU2391 ISGsu' GSU2392 ISGsu' GSU3082 ISGsu' GSU3083 ISGsu' GSU1079 Cro/Cl GSU1072 IcIR fa GSU1626 GntR GSU2779 MerR GSU1068 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331	7, transposase OrfB 7, transposase OrfB 7, transposase OrfB 7, transposase OrfB 7, transposase OrfA 7, transposase OrfA 7, transposase OrfB 6 family transcriptional regulator family transcriptional regulator family transcriptional regulator family transcriptional regulator family transcriptional regulator family transcriptional regulator	Regulatory functions	$\begin{array}{c} -1.009 \pm 0.300 \\ -0.993 \pm 0.263 \\ -0.827 \pm 0.309 \\ -1.136 \pm 0.397 \\ -0.833 \pm 0.253 \\ -0.946 \pm 0.237 \\ -1.249 \pm 0.374 \\ -0.542 \pm 0.075 \\ -1.200 \pm 0.414 \\ -0.626 \pm 0.109 \\ -0.728 \pm 0.091 \end{array}$
GSU2171 ISGsu' GSU2279 ISGsu' GSU2391 ISGsu' GSU2392 ISGsu' GSU3082 ISGsu' GSU3083 ISGsu' GSU0079 Cro/Cl GSU1072 IclR fa GSU2046 DNA-I GSU2079 MerR GSU1070 Sodiur GSU1070 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331	7, transposase OrfB 7, transposase OrfB 7, transposase OrfB 7, transposase OrfA 7, transposase OrfA 7, transposase OrfA 7, transposase OrfB family transcriptional regulator mily transcriptional regulator family transcriptional regulator family transcriptional regulator family transcriptional regulator family transcriptional regulator	Regulatory functions	$\begin{array}{c} -0.993 \pm 0.263 \\ -0.827 \pm 0.309 \\ -1.136 \pm 0.397 \\ -0.833 \pm 0.253 \\ -0.946 \pm 0.237 \\ -1.249 \pm 0.374 \\ \end{array}$
GSU2279 ISGsu' GSU2391 ISGsu' GSU2392 ISGsu' GSU3082 ISGsu' GSU3083 ISGsu' GSU3083 ISGsu' GSU1079 Cro/Cl GSU1072 IcIR fa GSU1626 GntR GSU2046 DNA-I GSU2079 MerR GSU1070 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331	7, transposase OrfB 7, transposase OrfB 7, transposase OrfA 7, transposase OrfA 7, transposase OrfB family transcriptional regulator mily transcriptional regulator family transcriptional regulator poinding response regulator family transcriptional regulator family transcriptional regulator	Regulatory functions	$\begin{array}{c} -0.827 \pm 0.309 \\ -1.136 \pm 0.397 \\ -0.833 \pm 0.253 \\ -0.946 \pm 0.237 \\ -1.249 \pm 0.374 \\ \end{array}$ $\begin{array}{c} -0.542 \pm 0.075 \\ -1.200 \pm 0.414 \\ -0.626 \pm 0.109 \\ -0.728 \pm 0.091 \end{array}$
GSU2391 ISGsu GSU2392 ISGsu GSU3082 ISGsu GSU3083 ISGsu GSU3083 ISGsu GSU1072 IclR fa GSU1626 GntR GSU2046 DNA-I GSU2046 DNA-I GSU2779 MerR GSU1068 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331 RND	, transposase OrfB 7, transposase OrfA 7, transposase OrfA 7, transposase OrfB 6 family transcriptional regulator mily transcriptional regulator family transcriptional regulator poinding response regulator family transcriptional regulator family transcriptional regulator	Regulatory functions	$\begin{array}{c} -1.136 \pm 0.397 \\ -0.833 \pm 0.253 \\ -0.946 \pm 0.237 \\ -1.249 \pm 0.374 \\ \end{array}$ $\begin{array}{c} -0.542 \pm 0.075 \\ -1.200 \pm 0.414 \\ -0.626 \pm 0.109 \\ -0.728 \pm 0.091 \end{array}$
GSU2392 ISGsu GSU3082 ISGsu GSU3083 ISGsu GSU3083 ISGsu GSU1072 IclR fa GSU1626 GntR GSU246 DNA-I GSU2779 MerR GSU1068 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331 RND	, transposaseOrfA 7, transposaseOrfA 7, transposaseOrfB family transcriptional regulator mily transcriptional regulator family transcriptional regulator binding response regulator family transcriptional regulator hysolute symporter family protein	Regulatory functions	$\begin{array}{c} -0.833 \pm 0.253 \\ -0.946 \pm 0.237 \\ -1.249 \pm 0.374 \\ \end{array}$ $\begin{array}{c} -0.542 \pm 0.075 \\ -1.200 \pm 0.414 \\ -0.626 \pm 0.109 \\ -0.728 \pm 0.091 \end{array}$
GSU3082 ISGsu GSU3083 ISGsu GSU3083 ISGsu GSU1072 IcIR fa GSU1626 GntR GSU2046 DNA-I GSU2779 MerR GSU1068 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331 RND	, transposaseOrfA 7, transposaseOrfB family transcriptional regulator mily transcriptional regulator family transcriptional regulator binding response regulator family transcriptional regulator h/solute symporter family protein	Regulatory functions	$\begin{array}{c} -0.946 \pm 0.237 \\ -1.249 \pm 0.374 \\ \end{array}$ $\begin{array}{c} -0.542 \pm 0.075 \\ -1.200 \pm 0.414 \\ -0.626 \pm 0.109 \\ -0.728 \pm 0.091 \end{array}$
GSU3083 ISGsu' GSU0079 Cro/Cl GSU1072 IclR fc GSU1626 GntR GSU2046 DNA-I GSU2779 MerR GSU1068 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331 RND	f, transposase OrfB family transcriptional regulator mily transcriptional regulator family transcriptional regulator binding response regulator family transcriptional regulator h/solute symporter family protein	Regulatory functions	$\begin{array}{c} -1.249 \pm 0.374 \\ -0.542 \pm 0.075 \\ -1.200 \pm 0.414 \\ -0.626 \pm 0.109 \\ -0.728 \pm 0.091 \end{array}$
GSU0079 Cro/Cl GSU1072 IclR fa GSU1626 GntR GSU2046 DNA-I GSU2779 MerR GSU1068 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331 RND	family transcriptional regulator mily transcriptional regulator family transcriptional regulator binding response regulator family transcriptional regulator n/solute symporter family protein	Regulatory functions	$\begin{array}{c} -0.542 \pm 0.075 \\ -1.200 \pm 0.414 \\ -0.626 \pm 0.109 \\ -0.728 \pm 0.091 \end{array}$
GSU1072 IclR fa GSU1626 GntR GSU2046 DNA-I GSU2779 MerR GSU1068 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331 RND	mily transcriptional regulator family transcriptional regulator pinding response regulator family transcriptional regulator n/solute symporter family protein		$\begin{array}{c} -1.200 \pm 0.414 \\ -0.626 \pm 0.109 \\ -0.728 \pm 0.091 \end{array}$
GSU1626 GntR GSU2046 DNA-I GSU2779 MerR GSU1068 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331 RND	family transcriptional regulator binding response regulator family transcriptional regulator n/solute symporter family protein		$\begin{array}{c} -0.626 \pm 0.109 \\ -0.728 \pm 0.091 \end{array}$
GSU2046 DNA- GSU2779 MerR GSU1068 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331 RND	pinding response regulator family transcriptional regulator n/solute symporter family protein		-0.728 ± 0.091
GSU1068 Sodiur GSU1068 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331 RND	family transcriptional regulator		
GSU1068 Sodiur GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331 RND	n/solute symporter family protein		-0.966 ± 0.172
GSU1070 Sodiur GSU1261 ABC t GSU1330 Metal prot GSU1331 BND		Transport and binding proteins	-1.771 ± 0.167
GSU1261 ABC t GSU1330 Metal prot GSU1331 RND	n/solute symporter family protein		-1.670 ± 0.231
GSU1330 Metal prot GSU1331 RND	ransporter, ATP-binding protein		-0.534 ± 0.053
GSU1331 RND	ion efflux outer membrane protein family ein		-1.142 ± 0.400
0001001	amily efflux transporter		-0.774 ± 0.144
GSU2005 Branch	ed-chain amino acid ABC transporter, plasmic amino acid-binding protein		-1.938 ± 0.395
GSU2006 Branch	hed-chain amino acid ABC transporter,		-1.776 ± 0.312
GSU2007 Branch	hed-chain amino acid ABC transporter,		-1.540 ± 0.346
GSU2008 Branch	hed-chain amino acid ABC transporter,		-1.259 ± 0.133
GSU2352 Sodiur	1/solute symporter family protein		-1.508 ± 0.193
GSU2490 Oxalat	e/formate antiporter		-1.957 ± 0.510
GSU2886 TonB-	dependent receptor		-1.501 ± 0.239
GSU0711 Endon	uclease/exonuclease/phosphatase family ein	Unknown functions	-0.745 ± 0.136
GSU0930 Sulfur	transferase		-0.758 ± 0.159
GSU1338 Heavy	metal-associated domain protein		-1.225 ± 0.341
GSU1398 SCO1/	SenC family protein		-0.774 ± 0.144
GSU2010 CBS d	omain protein		-1142 ± 0.141
GSU3059 Radics	1 SAM domain protein		-0.623 ± 0.114
GSU3157 Alpha	heta fold family hydrolase		-0.725 ± 0.114
GSU3435 Ankvri	IN LA DINE FAILURY UVUITUANE		0.725 = 0.100

^a ID, identification.

proportional to the level of OmcB production (10). Furthermore, there is a direct correlation between transcript levels of *omcB* and rates of Fe(III) reduction in continuous cultures (4). However, as shown here, after extended incubation the OmcBdeficient mutant is able to adapt and grow with soluble Fe(III) as the sole electron acceptor.

One of the more surprising aspects of this adaptation is that, during growth, adapted OmcB-deficient cultures reduced soluble Fe(III) at rates comparable to the wild type, but cell yields were substantially lower, indicating that there is less energy conservation from the pathway for Fe(III) reduction in the adapted mutant. It is unlikely that OmcB, an outer membrane protein, plays a direct role in coupling the flow of electron transport to the generation of a proton motive force, because it is expected that this is primarily established from proton pumping across the inner membrane. However, it is possible that the electron transport pathway from inner membrane electron transport components to OmcB is tightly coupled and that alternative electron transfer pathways to Fe(III) reduction, in the absence of OmcB, require electron transfer from alternative inner membrane electron transfer components that yield less proton pumping. Alternatively, the apparent need for increased biosynthesis of alternative *c*-type cytochromes, and possibly other proteins, may increase energy consumption and decrease growth yields.

The increased production of other outer membrane c-type cytochromes associated with adaptation in the OmcB-deficient mutant may account for the ability of the adapted mutant to reduce Fe(III) by providing an alternative route(s) for electron transfer to soluble Fe(III).

Five of the nine up-regulated *c*-type cytochrome genes with higher levels of transcripts in the OmcB-deficient mutant (GSU594, 2495, 2503 [OmcT], 2811, and 2813) were also found to have higher transcript levels when *G. sulfurreducens* was grown under electron-acceptor limiting conditions, rather than with the electron donor limiting growth (A. Esteve-Nunez, unpublished data). Thus, the physiological state associated with the disruption of the electron transport pathway in the OmcB-deficient mutant may mimic the physiological state under electron acceptor-limiting conditions, and this, as well as

TABLE 4. Genes coding for *c*-type cytochromes whose expression levels were altered significantly (ca. twofold) in the OmcB-deficient mutant versus the wild type via microarray assay^{*a*}

Locus ID	Predicted location	Microarray log ₂ ratio	Predicted mol mass (kDa)	Northern blot result
GSU0594	IM	1.133	38.2	Up
GSU2494	IM	1.863	47.7	Up
GSU2495	Р	1.810	69.7	Up
GSU2503	OM	3.340	45.5	Up
GSU2504	OM	3.676	45.4	Úp
GSU2811	OM	0.745	52.7	Úp
GSU2813	Р	0.839	36.9	Úp
GSU2731 (OmcC)	OM	NA	77.2	Úp
GSU2737 (OmcB)	OM	-1.189	77.1	Down
GSU2808	OM	-0.730	20.4	Down
GSU2887	OM	-1.515	89.7	Down
GSU3259	IM	-1.069	57.4	Down

^{*a*} Abbreviations: ID, identification; IM, inner membrane; P, periplasm; OM, outer membrane; NA, not applicable; mol, molecular.

possible other physiological signals, may account for the increased production of outer membrane cytochromes.

In addition to affecting cytochrome production, the disruption of electron flow in the OmcB-deficient mutant appeared to have an impact on other aspects of metabolism. This was evident from the microarray studies which suggested that there was a decrease in transcription of several genes involved in acetate metabolism and an increase in transcript levels for genes such as a nickel-dependent hydrogenase (GSU0782 to -0787), a cytochrome *d* ubiquinol oxidase (GSU1640-1641), and a carbon monoxide dehydrogenase (GSU2098), which presumably played a role in maintaining the appropriate balance of reducing equivalents in the OmcB-deficient mutant (23, 27). Evidence that the mutant was under stress includes the apparent up-regulation of a *relA* homolog (GSU2236) which has been shown to be important in regulating levels of ppGpp and stress responses in *G. sulfurreducens* (L. DiDonato et al., sub-



FIG. 5. Northern blot analyses of up- or down-regulated *c*-type cytochrome genes. Total RNA was isolated from mid-log Fe(III) citrate-grown cultures of wild type (lane 1) or OmcB-deficient mutant (lane 2). (A) *c*-type cytochrome genes that were identified to be up-regulated by a \geq 2-fold change in the adapted OmcB-deficient mutant. (B) *c*-type cytochrome genes that were identified to be down-regulated in the adapted OmcB-deficient mutant. (C) A replicate gel was run and stained with ethidium bromide, revealing the 16S and 23S rRNA, as a loading control.

mitted for publication), as well as higher transcript levels for other stress response proteins, such as thioredoxin peroxidase (GSU3246), glutaredoxin family protein (GSU2812), and proteins involved in cell envelope biosynthesis (GSU2078, 2497), etc.

Significance of failure to adapt for Fe(III) oxide reduction. The fact that the OmcB-deficient mutant was never able to adapt to grow on Fe(III) oxide emphasizes the central role that OmcB plays in the reduction of this most environmentally relevant form of Fe(III). Although it was initially considered that OmcB might function as the terminal Fe(III) reductase because of its location in the outer membrane (10), subsequent



FIG. 4. Functional groups of up- or down-regulated genes from the adapted OmcB-deficient mutant. The up- or down-regulated genes of the OmcB-deficient mutant were identified by comparing expression patterns with that of the wild type as described in Materials and Methods. Functional classes are determined using the *Geobacter sulfurreducens* genome page from TIGR (http://www.tigr.org/tigr-scripts/CMR2/ GenomePage3.spl?database=ggs). Functional groups of up-regulated genes from OmcB-deficient mutant are represented by black bars, and functional groups of down-regulated genes are shown by gray-striped bars. The number of genes represented in each bar is indicated in the *x* axis.

studies have identified other proteins, such as the outer membrane c-type cytochrome OmcS (T. Mehta et al., submitted for publication) and pili (23a), that are more exposed on the outside of the cell and are specifically required for the reduction of Fe(III) oxide, but not soluble Fe(III). Thus, the critical role of OmcB in Fe(III) oxide reduction may be as an intermediary electron transfer component that establishes electron transfer to the terminal Fe(III) oxide reductase (13). The finding that the mutant deficient in OmcB can reduce soluble Fe(III), but not Fe(III) oxides, demonstrates that G. sulfurreducens can reduce soluble Fe(III) via mechanisms different than those required for Fe(III) oxide reduction. This is an important consideration when extrapolating results from pure cultures to Fe(III) reduction in sedimentary environments where Fe(III) oxides are the predominant Fe(III) form readily available for microbial reduction and concentrations of soluble Fe(III) are likely to be low (13, 21).

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