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Shewanella oneidensis MR-1 Sensory Box Protein Involved in Aerobic and Anoxic Growth

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Although little is known of potential function for conserved signaling proteins, it is hypothesized that such proteins play important roles to coordinate cellular responses to environmental stimuli. In order to elucidate the function of a putative sensory box protein (PAS domains) in *Shewanella oneidensis* MR-1, the physiological role of SO3389 was characterized. The predicted open reading frame (ORF) encodes a putative sensory box protein that has PAS, GGDEF, and EAL domains, and an in-frame deletion mutant was constructed (Δ SO3389) with approximately 95% of the ORF deleted. Under aerated conditions, wild-type and mutant cultures had similar growth rates, but the mutant culture had a lower growth rate under static, aerobic conditions. Oxygen consumption rates were lower for mutant cultures (1.5-fold), and wild-type cultures also maintained lower dissolved oxygen concentrations under aerated growth conditions. When transferred to anoxic conditions, the mutant did not grow with fumarate, iron(III), or dimethyl sulfoxide (DMSO) as electron acceptors. Biochemical assays demonstrated the expression of different *c*-type cytochromes as well as decreased fumarate reductase activity in the mutant transferred to anoxic growth conditions. Transcriptomic studies showed the inability of the mutant to up-express and down-express genes, including *c*-type cytochromes (e.g., SO4047/SO4048, SO3285/SO3286), reductases (e.g., SO0768, SO1427), and potential regulators (e.g., SO1329). The complemented strain was able to grow when transferred from aerobic to anoxic growth conditions with the tested electron acceptors. The modeled structure for the SO3389 PAS domains was highly similar to the crystal structures of FAD-binding PAS domains that are known O₂/redox sensors. Based on physiological, genomic, and bioinformatic results, we suggest that the sensory box protein, SO3389, is an O₂/redox sensor that is involved in optimization of aerobic growth and transitions to anoxia in *S. oneidensis* MR-1.

In the postgenomic era, uncharacterized genes pose a major challenge in biology (16), and many sequenced genomes still contain a large fraction (up to 30 to 40%) of genes without defined physiological roles (13). In fact, many aspects of signaling and stress response most likely reside in this fraction of genes for a given organism and underlie the lack of understanding in physiological responses and control of metabolism. Many uncharacterized genes/proteins are classified as sensory box proteins based upon conserved domains, but signals and cellular responses for most presumptive sensory box proteins are not known.

Shewanella oneidensis MR-1, a facultative anaerobe classified as a gammaproteobacterium, can utilize numerous inorganic compounds as electron acceptors (e.g., oxygen, nitrate, and metals). The *S. oneidensis* MR-1 genome sequence was determined (20), and the most recent annotation of the 5.1-Mb genome estimated 4,467 genes, of which 1,623 had hypothetical functions (28). Many bacteria considered to be “environmen-

tal” organisms typically have diverse metabolic capacity, particularly facultative organisms such as *Shewanella* spp., and aside from well-studied systems (e.g., *Escherichia* and *Bacillus* pathogens) little is known about how these organisms sense and respond to changing environments.

An earlier study reported a correlation between the total number of PAS protein domains (*Drosophila* period clock, aryl hydrocarbon receptor, single-minded protein) (41) and the numbers of respiratory and photosynthetic electron transport-associated proteins in completely sequenced microbial genomes (59). The MR-1 genome has approximately 31 open reading frames (ORFs) that are predicted to encode PAS domains, and this compares to approximately 15 PAS domains in *Escherichia coli* K-12 MG1655 (www.microbesonline.org). PAS domains are usually part of larger polypeptides involved in signal transduction (one- and two-component systems), and previous studies have shown that PAS domains can sense changes in environmental stimuli, such as light, redox potential, oxygen, and energy state of the cell (53). A high number of PAS proteins likely coincide with the fact that MR-1 can utilize a wide variety of electron acceptors and must be able to coordinate a diverse metabolic capacity with various environmental conditions.

The Pfam database (version 24.0) reports 33,313 sequences

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classified in the PAS protein clan, which contains 7 families, and the PAS and PAC motifs adopt a single globular fold of approximately 100 residues now known as the PAS domain (34). PAS domains are observed in all three domains of life, including metazoans, in which the domains can be part of hypoxia-inducible factors that play important roles in anaerobic metabolism, oxygen delivery, and angiogenesis (46). These observations indicate the importance of PAS proteins in biology; however, physiological roles for the vast majority of putative PAS proteins are unknown.

The fact that most sequenced genomes contain a high percentage of conserved hypothetical proteins with unknown physiological roles represents a knowledge gap across biology, and the current work set out to characterize the physiological role of a conserved sensory box protein. The mutant was affected in low-oxygen growth responses and transitions to anoxia, and these phenotypes could be at least partially explained by altered *c*-type cytochrome content, oxygen consumption, and reductase activity. In addition, the modeled PAS domain structure suggested high structural similarity to known flavin adenine dinucleotide (FAD)-based O₂/redox sensors.

MATERIALS AND METHODS

Sequence comparisons. The Smart (Simple Modular Architecture Research Tool) database (version 6.0) and the MicrobesOnline website were used for domain predictions and comparisons as previously described (11, 31). Phylogenetic and molecular evolutionary analyses were conducted with MUSCLE (version 3.7) (www.ebi.ac.uk/Tools/muscle/index.html) and within the MicrobesOnline workbench (www.microbesonline.org). Maximum likelihood trees were constructed with a Jones, Taylor, and Thornton (JTT) model with no correction for across-site rate variation with the MicrobesOnline workbench. The protein model for SO3389 was predicted with CPHmodels (version 3.0) and I-Tasser (37, 43). The CPHmodels quality was checked with QMEAN (3), ProSA (58), and MolProbity (10). I-Tasser has an internal method to check model quality (43). Protein Structural Alignment (version 3.0.4) with GANGSTA⁺, DalLite (version 3.1), Phyre (version 0.2), and I-Tasser were used for structural alignments with crystal structures of known PAS domains (19, 23, 25, 43).

Mutant construction. An approximately 2.4-kbp fragment was deleted from SO3389 (2,639 bp) with PCR-based crossover amplification as described previously (1, 17, 56). The primer pairs used to generate two DNA fragments with 3'-staggered ends were as follows: No-5'-CGCGAGCTCGTTCCGACCTTCGATAAATC-3' and Ni-5'-TGTTAACTAGTGGATGGGGCCCAATCTGTAACTGCTT-3' and Co-5'-CGCGAGCTCGACCTTGACGCAAATGATC-3' and Ci-5'-CCATCCACTAAGTTAAACAGCGGATGTTGAGGCCTTTT-3' (noncomplementary tag sequences are underlined), and the outside primers were used to amplify a fusion molecule. The crossover PCR product was purified and digested by *Sac*I, cloned into treated suicide vector pDS3.0, and electroporated into *E. coli* S17-1/λ_{pir}. The suicide plasmid construct was moved into MR-1 as previously described (56), and the mutated SO3389 gene was verified by PCR (5'-GAGTGGCATTTCAGCACTAGA-3' and 5'-CATACGGTCGGTTCATCAA-3') and sequence determination. The resulting mutant did not possess any of the predicted domains after completion of the in-frame deletion. The mutation was an in-frame deletion that removed approximately 2,400 bp of the gene.

ΔSO3389 complementation. Multiple vectors (pACYC184, pJB3Cm6, pBBR1MCS-5, and pBAD; see Table 1) were attempted for complementation, but construction of the correct, stable *E. coli* strain was not successful or expression was not detected in the case of pBAD. Therefore, pBBR1MCS-2 and pBBAD were used, and the following complementation primers were used to amplify genomic DNA: 3389FWDCOM (GAATTCTGTCTGCGGTAATATC TGCG) and 3389REVCOM (GGATCCACAAAAGCAATCATGTGCG) (Table 1). The amplicon was cloned into a TOPO vector and digested with *Eco*RI and *Bam*HI. After digestion, the insert was purified with a QIAquick gel extraction kit as per the instructions and subsequently cloned by ligation into vectors pBBR1MCS-2 and pBBAD at 1:3 and 1:10 molar ratios (vector to insert), also digested using *Eco*RI and *Bam*HI. To increase transformation efficiency, ligations were transformed into *E. coli* UQ950 and then *E. coli* WM3064, a

TABLE 1. Strains and vectors

Strain and vector	Description	Reference
Strains		
<i>S. oneidensis</i> MR-1		20
ΔSO3389 mutant	In-frame deletion of SO3389	This study
ΔSO3389-pBBR1MCS-SO3389 mutant	Complemented strain	This study
<i>Escherichia coli</i>		
S17-1/λ _{pir}	Mating strain	56
UQ950	MDH5α/λ _{pir}	45
WM3064	DAP autotroph	45
Vectors		
pJB3Cm6	Expression vector	4
pBBR1MCS-2	Expression vector	29
pBBR1MCS-5	Expression vector	Gift from Y. Yang
pACYC184	Expression vector	36
pBAD	Expression vector	50
pBBAD	Expression vector	52
pDS3.0	Pir-dependent suicide vector	56

diaminopimelic (DAP) auxotroph. Transformants thus obtained were verified for inserts by PCR. The correct transformant was then mated with ΔSO3389 onto an LB-DAP (0.3 mM) plate and incubated at 30°C for 24 h. Transconjugants were selected by streaking mating mixtures onto LB-kanamycin-50.

Growth. The mutant was grown in aerobic and anoxic defined minimal medium. The defined medium (HBa) contained the following ingredients: PIPES [piperazine-*N,N'*-bis(2-ethanesulfonic acid)], 3 mM; NaOH, 7.5 mM; NH₄Cl, 28 mM; KCl, 1.3 mM; NaH₂PO₄, 4.4 mM; Na₂SO₄, 30 mM; minerals, 10 ml l⁻¹; vitamins, 10 ml l⁻¹; and lactate, 90 mM. Amino acids, CaCl₂ (0.7 mM), and electron acceptors [fumarate, Fe(III)-citrate, dimethyl sulfoxide (DMSO)] were added after autoclaving. The vitamin mix contained biotin, 0.002 g liter⁻¹; folic acid, 0.002 g liter⁻¹; pyridoxine HCl, 0.01 g liter⁻¹; riboflavin, 0.005 g liter⁻¹; thiamine, 0.005 g liter⁻¹; nicotinic acid, 0.005 g liter⁻¹; pantothenic acid, 0.005 g liter⁻¹; B-12, 0.0001 g liter⁻¹; *p*-aminobenzoic acid, 0.005 g liter⁻¹; and thioctic acid, 0.005 g liter⁻¹. The amino acid mix contained glutamic acid, arginine, and serine (2.0 g liter⁻¹ each). The mineral mix contained nitrilotriacetic acid, 1.5 g liter⁻¹; MgSO₄, 3.0 g liter⁻¹; MnSO₄ · H₂O, 0.5 g liter⁻¹; NaCl, 1.0 g liter⁻¹; FeSO₄ · 7H₂O, 0.1 g liter⁻¹; CaCl₂ · 2H₂O, 0.1 g liter⁻¹; CoCl₂ · 6H₂O, 0.1 g liter⁻¹; ZnCl₂, 0.1 g liter⁻¹; CuSO₄ · 5H₂O, 0.01 g liter⁻¹; AlK(SO₄)₂ · 12H₂O, 0.01 g liter⁻¹; H₃BO₃, 0.01 g liter⁻¹; Na₂MoO₄, 0.03 g liter⁻¹; NiCl₂ · 6H₂O, 0.03 g liter⁻¹; and Na₂WO₄ · 2H₂O, 0.03 g liter⁻¹. For aerobic growth, cultures (50 ml) were grown in 300-ml shake flasks (various rpm at 30°C). Static cultures were grown as 50-ml cultures in 300-ml side-arm flasks but were not shaken. Anoxic medium was prepared by boiling under O₂-free N₂ gas, dispensed into sparged tubes (N₂) or bottles, and sealed with butyl stoppers and aluminum crimp seals. Anoxic growth was done at 30°C, and growth was monitored via a spectrophotometer (600 nm).

Growth for transcriptomic analysis. Wild-type and mutant cells were grown as described above or anoxically in 2-liter medium bottles, and then the cells were harvested, centrifuged at 4°C, and snap-frozen in triplicate for RNA extraction. For wild-type growth, samples were collected during aerobic exponential-phase growth, 10 h postinoculation in anoxic medium (anoxic, exponential-phase growth), and approximately 70 h into anoxic growth (anoxic, stationary-phase growth). Samples for mutant cell growth were harvested at similar time points, and 10 h postinoculation into anoxic medium represented the lag phase for the mutant when transferred to anoxia. All samples were centrifuged and snap-frozen in triplicate, and RNA was extracted and used subsequently for microarray analysis.

Transcriptomics. Microarray fabrication, hybridization, probe labeling, image acquisition, and processing were carried out as described previously (1, 17). Gene expression analysis was performed using three biological replicates for each microarray experiment, and each slide contained two replicates of two independent probes for each considered gene in the *S. oneidensis* MR-1 transcriptome (32). Microarray data analyses were performed using gene models from NCBI.

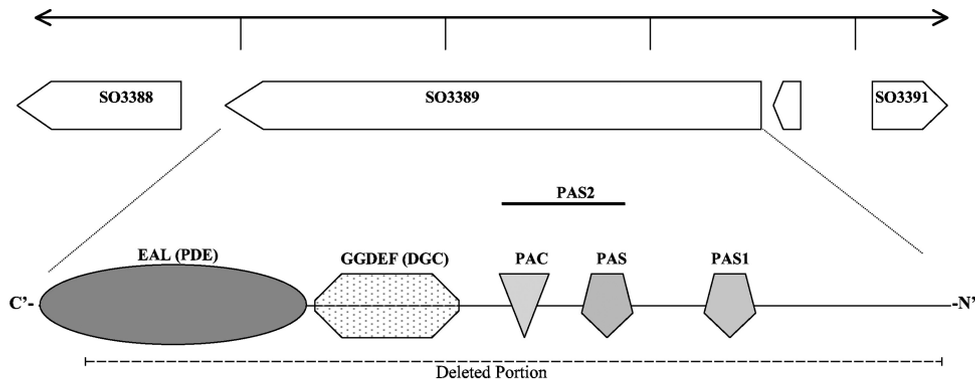


FIG. 1. Genome region view and domain architecture of SO3389. SO3388 is annotated as an RNA helicase, SO3390 is annotated as a small (36-amino-acid) hypothetical protein, and SO3391 is annotated as an ATP-dependent protease. The domains were predicted with SMART version 6.0 (<http://smart.embl-heidelberg.de/>).

All mRNA changes were assessed with total genomic DNA as a control. Log_2 ratios and z scores were computed as previously described (32).

DO and ORP measurements. Dissolved oxygen (DO) was measured with an SG6 DO meter (Mettler Toledo) at 5-s intervals for 4 min. Oxidation-reduction potential (ORP) was measured via an HQ20 probe (Hach) at 5-s intervals for 4 min. Measurements were made for wild-type and mutant cultures at similar cell densities (optical density [OD] at 600 nm of approximately 0.3) during static incubation of an aerobic culture or during growth at a respective shake speed. It should be noted that cultures were pregrown at the respective condition and the inoculums were in mid-exponential phase.

SDS-PAGE and cytochrome stain. In order to determine the *c*-type cytochrome content of wild-type and mutant cells, spheroplast and periplasmic fractions were obtained as previously described (42). Prior to electrophoresis, samples were incubated with urea and SDS at room temperature (1 h) as previously described (42) and separated on a gradient polyacrylamide gel (4 to 20%). The *c*-type cytochromes were observed via a heme-linked peroxidase stain with TMBZ (3,3',5,5'-tetramethylbenzidine) as previously described (42). Protein was quantified via the Lowry method, with serum albumin as the standard.

Fumarate reductase. Cells were harvested and sonicated at a constant rate (1-s bursts) for 12 s on ice 4 times. After sonication, cells were centrifuged at 4°C for 15 min at $9,000 \times g$. Protein concentrations were obtained via the Lowry method, and equal amounts of protein were loaded. Fumarate reductase activity in native gels was detected as a clear band as a result of oxidized methyl viologen after the addition of fumarate (250 mM). Gels were incubated in a round-bottom flask with 10 mM methyl viologen in 0.05 M potassium phosphate (10 ml; pH 7.2; 1 mM sodium dithionite) that was gassed continuously with oxygen-free N_2 (33).

Fumarate reductase activity was quantified by the benzyl viologen-linked reductase assay as described previously (49, 57). Reagents (75 mM sodium phosphate buffer [pH 6.8], 0.2 mM benzyl viologen, and cell extract) were added to a 1-ml cuvette through the stopper while being gassed continuously. Sodium dithionite (20 mM) was added to the cuvette until absorbance at 585 nm reached 0.8 to 0.9 (half-reduced benzyl viologen). Sodium fumarate (5 mM, final concentration) was then added to the mixture, and oxidation was immediately recorded at 585 nm. Activity was expressed in millimoles of benzyl viologen oxidized minute^{-1} microgram $^{-1}$.

Quantitative PCR. Quantitative PCR (qPCR) was performed with RNA extracts from wild-type and mutant cells under aerobic and anoxic conditions. cDNA was prepared by reverse transcription with 5 μg RNA. A mixture was prepared containing RNA and random primers and was incubated at 70°C for 10 min and then cooled on ice. Reverse transcription was performed by adding 4 μl dithiothreitol (DTT), 1 μl deoxynucleoside triphosphates (dNTPs) at 10 mM, 1 μl RNase inhibitor, and 1 μl R reverse transcriptase (Invitrogen). cDNA was generated with incubation at 42°C for 2 h and 70°C for 15 min. After being cooled, the concentration of cDNA was measured at 260 nm. In order to perform quantitative PCR, cDNA was first diluted to 20 ng μl^{-1} , and 2 μl of diluted sample was used per reaction. Primers were added at a 10 mM concentration along with SYBR green Master Mix (Applied Biosystems) and nuclease-free water. Standards were prepared by carrying out serial dilution of genomic DNA, starting with 10^8 copies μl^{-1} to 10^1 copies μl^{-1} . Negative control without cDNA was included along with all runs. qPCR was performed using Smart Cycler II (Cepheid) by using the following conditions: initial denaturation, 95°C for 9.5

min; amplification, 95°C for 15 s, 55°C for 30 s, and 60°C for 30 s. Stage 2 was repeated to complete 45 cycles. Final extension was at 60°C for 95 s.

Total heme quantification. Wild-type and mutant cells were harvested and sonicated at a constant rate for 15 s (4 times) on ice. Cells were then centrifuged at 4°C for 15 min at $9,000 \times g$. Protein was quantified by the Lowry assay of the fractions obtained after sonication. Total heme was measured per μg of protein using the QuantiChrom heme assay kit according to the manufacturer's instructions (BioAssay Systems).

RESULTS AND DISCUSSION

SO3389 in *Shewanella*. The predicted monocistronic ORF (SO3389) encodes a putative 879-amino-acid polypeptide, and the predicted protein was assigned to COG5001, which contained EAL (E value of $4.0e-111$; named for conserved amino acids), GGDEF (E value of $3.6e-62$; named for conserved amino acids), and PAS ($5.5e+0.0$; E value of $2.4e-08$) domains (Fig. 1; E values are from the SMART database). The respective Interpro families were IPR003018, IPR000014, IPR013656, and IPR013767 (<http://www.ebi.ac.uk/interpro/>). A signal sequence was not predicted via SignalP (version 3.0) (14) or TatP (version 1.0) (2), although one hydrophobic α -helix was predicted from residue 137 to 162 with TMphred (22) and DAS (9, 9a) and weakly predicted with TMHMM (version 2.0) (35). Further work is needed to confirm the localization of SO3389 in the cytoplasm or cytoplasmic membrane.

Phylogenetic relationships. The proteins assigned to COG5001 are identified in 708 genomes from very diverse bacteria, and over 20 genomes contain 20 or more COG5001 proteins (data not shown). To date, 22 *Shewanella* genome sequences are available in public databases (<http://img.jgi.doe.gov/>), and 15 contained an ortholog to SO3389 with peptide sequence identity between 52 and 85% (<http://microbesonline.org/>). *Shewanella putrefaciens* CN-32, *S. putrefaciens* 200, *Shewanella woodyi*, *Shewanella pealeana*, *Shewanella violacea*, and *Shewanella* W3-18-1 did not contain a predicted ortholog to SO3389. The closely related sequences of SO3389 were detected in *Shewanella* ANA-3, *Shewanella* MR-4, and *Shewanella* MR-7 (84 to 85% identity) as well as in *Vibrio* species (Fig. 2). When the entire protein sequence of SO3389 was compared to sequenced genomes, the closest non-*Shewanella* relatives were putative sensory box proteins in *Vibrio* species, and sequences were also detected in *Cyanothecce*, *Neptuniibac-*

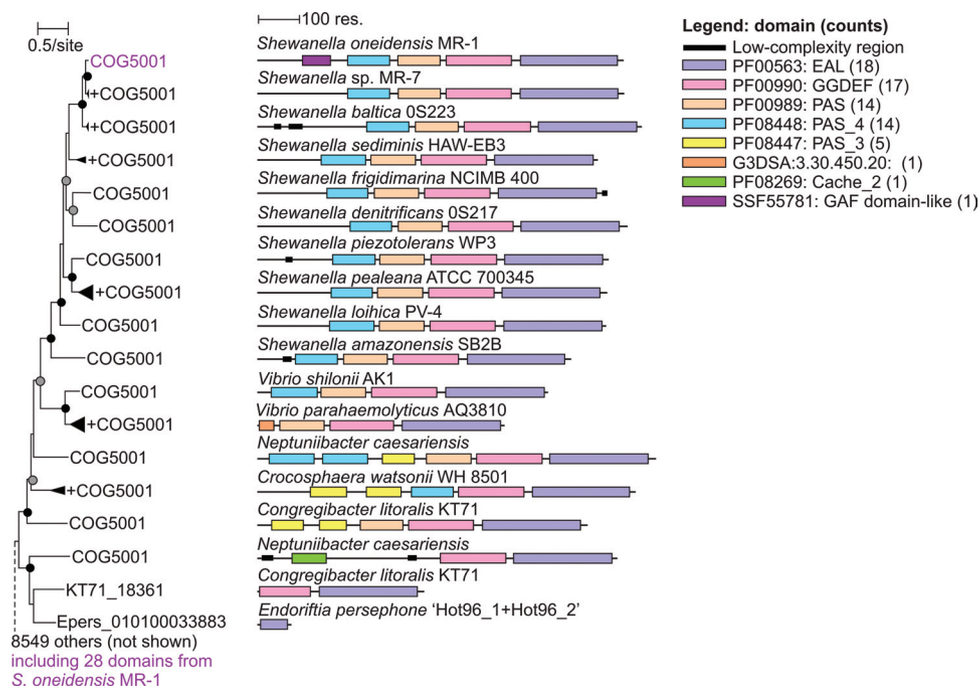


FIG. 2. Phylogram based upon PAS domains related to SO3389 with a neighbor-joining method within the MicrobesOnline workbench (<http://microbesonline.org>).

ter, *Congregibacter*, *Crocospaera*, and *Endoriftia*. Based upon PAS domain sequence phylogeny (see Materials and Methods), the closest relatives to SO3389 were observed in other *Shewanella* and *Vibrio* species; however, physiological roles have not been assigned to these putative proteins (Fig. 2).

Growth phenotypes. The mutant was tested for growth defects in aerobic (static and aerated) and anoxic defined minimal medium. The cell cultures (wild-type and Δ SO3389 cultures) did not differ significantly in growth rates in shaken aerobic medium (50 to 200 rpm), but the mutant culture had a lower growth rate in aerobic, static cultures than the wild type (decreased 1.6-fold; $P = 0.05$) (Fig. 3a and b). Oxygen concentrations were depleted at a lower rate by mutant cells (first-order rate constants were approximately 1.5-fold lower) than those by wild-type cells when cultures grown at 150 rpm were then incubated statically (Fig. 4a). In addition, the oxidation-reduction potential (ORP) declined at a lower rate for the mutant culture than for the wild type (Fig. 4a). When the DO concentrations were measured in actively shaken cultures, the mutant also had lower rates of O_2 consumption (approximately 1.5-fold) but also had higher steady-state DO concentrations (0.35 mg l^{-1} [0.01 mM] versus 1.7 mg l^{-1} [0.05 mM] and 2.8 mg l^{-1} [0.09 mM] versus 3.7 mg l^{-1} [0.12 mM], respectively, for wild-type and mutant cultures) (Fig. 4b). These results indicated that wild-type cultures increased consumption rates in response to increasing O_2 concentrations. In addition, the mutant culture was deficient in O_2 regulation, which resulted in less-efficient O_2 consumption at low and high O_2 concentrations.

When cells were transferred from aerobic to anoxic medium with lactate and fumarate, the Δ SO3389 cells did not grow (Fig. 5). Similar results were observed when the anoxic me-

dium contained lactate as the electron donor and DMSO or Fe(III)-citrate as the electron acceptor (data not shown). These results indicated that the mutant was deficient in response to anoxia.

Complementation. Complementation was achieved with the pBBRMCS-2 vector and a full-length SO3389 gene. The complemented strain, Δ SO3389:pBBR1MCS-2-SO3389, was able to grow when transferred to anoxic medium without a lag (Fig. 6). In addition, the complemented strain grew more similarly to the wild type (empty vector strain) under aerobic conditions (Fig. 3b). These results confirmed that the growth defect during growth at low DO concentrations and transition from aerobic conditions to anoxia were due to the Δ SO3389 mutation.

qPCR was used to monitor the expression of SO3389 and SO3388 in wild-type and mutant cells under aerobic and anoxic conditions. Wild-type cells had higher expression levels of SO3389 under aerobic and anoxic conditions than the deletion mutant (2.5-fold), and the SO3388 and SO3390 genes had similar expression levels in both wild-type and mutant cells under aerobic and anoxic conditions (data not shown). These results indicated that the Δ SO3389 mutation had not altered SO3388 or SO3390 expression.

Cytochrome *c*-type content. Due to the response to anoxic growth, spheroplast and periplasmic fractions were obtained as previously described in order to determine *c*-type cytochrome content (42). Minor differences were observed in the spheroplast and periplasmic fractions for aerobically grown wild-type and mutant cells (i.e., intensity differences). When cells were allowed to transition from aerobic to anoxic conditions (5 to 10 h posttransfer), differences were observed in the spheroplast fraction, most noticeably for heme-containing polypeptides with estimated molecular weights of 57, 33, and 20 kDa

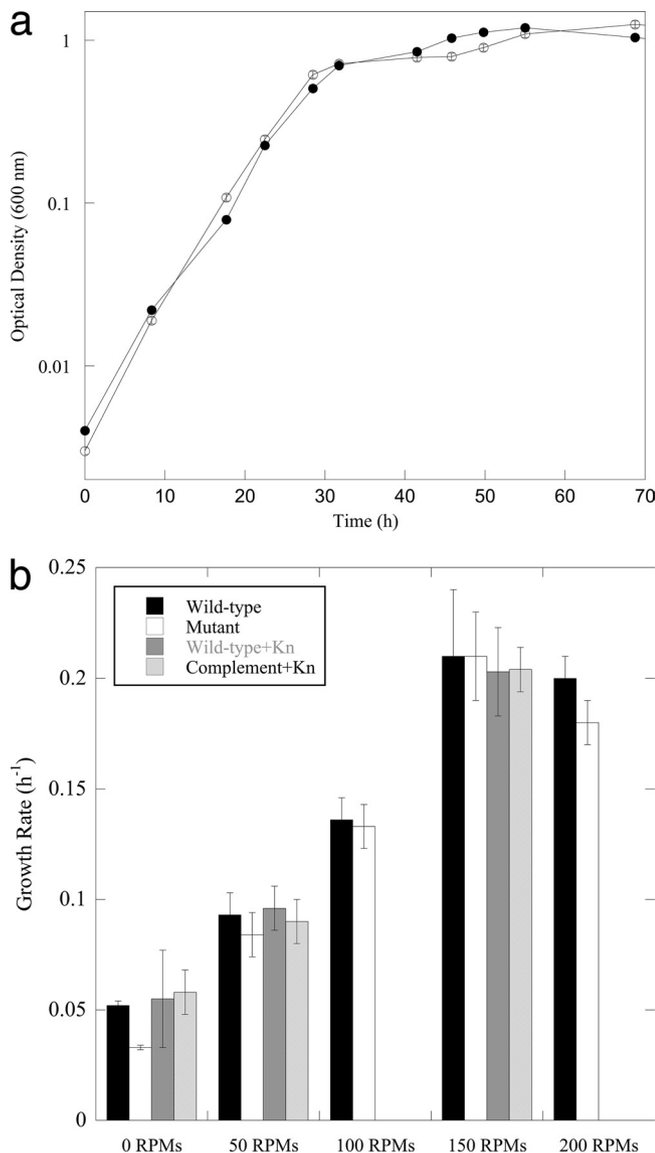


FIG. 3. (a) Growth curves of wild-type MR-1 (●) and Δ SO3389 (○) cells in aerobic, defined minimal medium with lactate, shaken at 150 rpm (error bars represent standard deviations). (b) Growth rates of wild-type (black) and mutant (white) planktonic cultures at different shake speeds (rpm). The static wild-type and mutant cultures were statistically different ($P = 0.05$). In addition, wild-type with empty vector (dark gray) and complement (light gray) strains were grown at the respective shake speeds in the presence of kanamycin.

(Fig. 7a and b). These results indicated that the Δ SO3389 mutant was deficient in at least three *c*-type cytochromes during the transition period from aerobic to anoxic conditions.

In addition to the qualitative analysis of cytochromes, total heme content (soluble and membrane) was quantified for wild-type and mutant cells grown under aerobic conditions and transition to anoxia. There was not a significant difference in the heme contents of wild-type and mutant cell fractions grown aerobically (1.4 ± 0.1 versus 1.5 ± 0.1 ng μ g⁻¹ protein), and these data coincided with the similar cytochrome profile of wild-type and mutant cells grown under aerobic conditions. During the transition to anoxia, the mutant cells exhibited

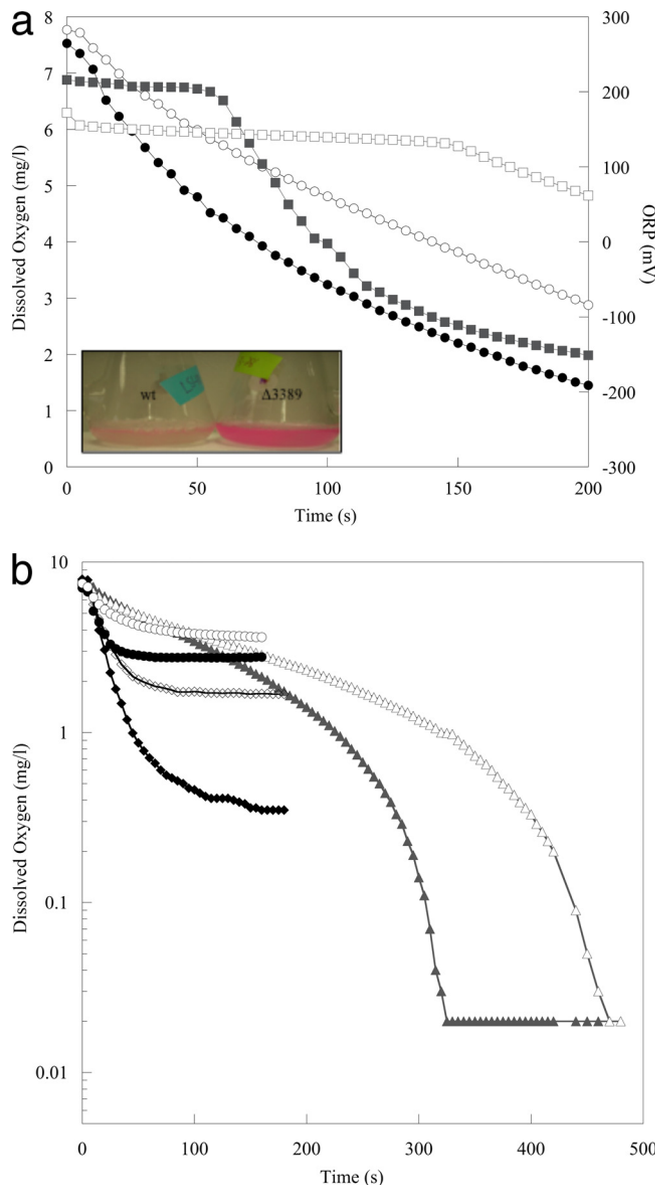


FIG. 4. (a) Dissolved oxygen measurements for WT (●) and Δ SO3389 (○) cells and oxidative-reductive potential (mV) measurements for WT (■) and Δ SO3389 (□) cells versus time when shaken aerobic cultures were incubated statically. The inset image shows the resazurin reduction-oxidation indicator for wild-type (left) and mutant (right) cultures postincubation. (b) The decrease in DO (mg/liter) was measured for cultures growing in lactate medium for wild-type (filled symbols) and mutant (empty symbols) cells at 0 rpm (▲, △), 150 rpm (◆, ◇), and 200 rpm (●, ○). The data represent an average of results from multiple experiments.

elevated levels of heme in comparison to wild-type cells from membrane fractions (0.15 ± 0.01 versus 0.27 ± 0.06 ng μ g⁻¹ protein).

Gene expression. A possible explanation for elevated heme content in the mutant is the overexpression of heme-containing proteins in order to compensate for the decline in oxygen affinity (at the cellular level) and/or the mutant failed to down-express certain cytochromes typically expressed during aerobic growth once transferred to anoxia. The later explanation is

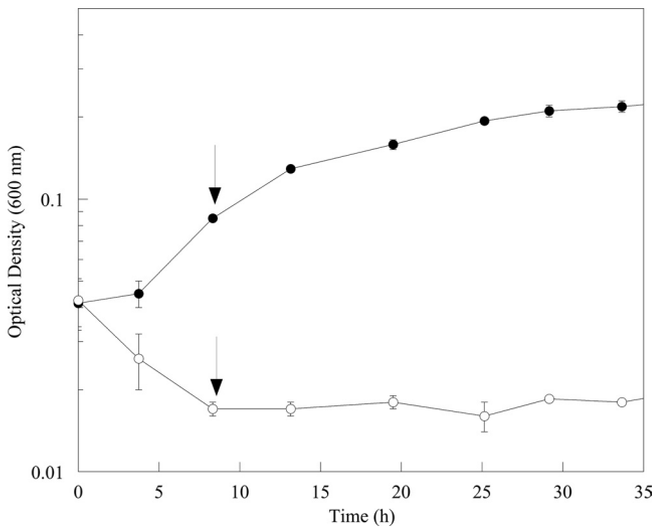


FIG. 5. Growth curves of wild-type MR-1 (●) and Δ SO3389 (○) cells in anaerobic, defined minimal medium with lactate and fumarate. Aerobic cultures were grown in shake flasks and used to inoculate anoxic cultures to similar initial ODs. The arrows depict the sampling time point for transcriptomic analyses. Error bars represent standard deviations.

more likely and coincides with the microarray data for numerous cytochromes that were significantly down-expressed in wild-type but not mutant cells upon transfer to anoxia (e.g., SO3285, SO3286, SO1427) (Table 2). During the transition from aerobic to anoxic conditions, wild-type cells up-expressed

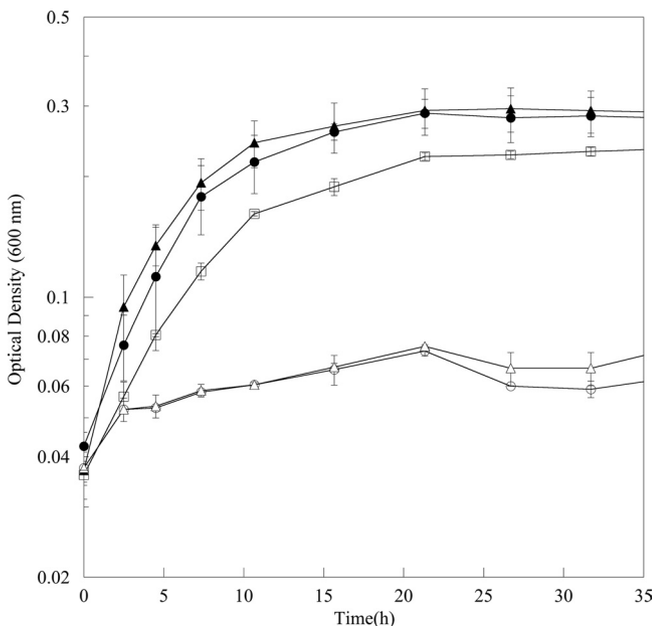


FIG. 6. Growth curves of wild-type MR-1 (●), Δ SO3389 (○), wild-type pBBR1MCS-2 (▲), Δ SO3389 pBBR1MCS-2 (△), and Δ SO3389::pBBR1MCS-2-SO3389 (□) cells in defined minimal medium with lactate as the electron donor and fumarate as the electron acceptor. Wild-type^{kn} and Δ SO3389^{kn} were constructed by mating with WM3064 empty plasmid pBBR1MCS-2. Error bars represent standard deviations.

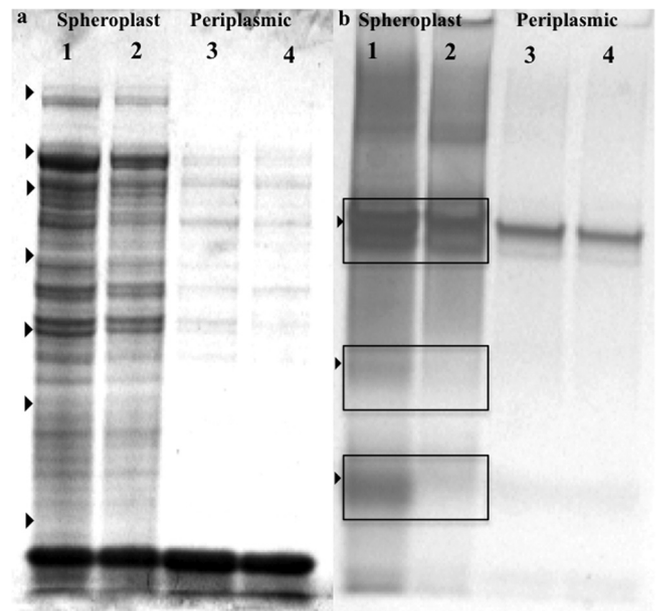


FIG. 7. SDS-PAGE gels of spheroplast (lanes 1 and 2) and periplasmic-shock (lanes 3 and 4) fractions of wild-type and Δ SO3389 cells stained with Coomassie blue for total protein (a) or cytochrome *c* heme stain (b). Equal amounts of total protein for wild-type (lanes 1 and 3) and Δ SO3389 (lanes 2 and 4) were loaded per well, and molecular weight markers are denoted with arrows in panel a: 150, 100, 75, 50, 35, 25, and 15 kDa.

54 genes and down-expressed 103 genes (\log_2 values of ≥ 2.0 and *z* scores of ≥ 2.0). In comparison, under the tested transitions from aerobic to anoxic conditions, the mutant cells did not have any significant expression changes that were greater than 2.0 (*z* scores of ≥ 2.0).

SO1427 was previously shown to be a DMSO reductase subunit (18), and the expression data suggested that SO3389 could be involved in coordinating the expression of reductases in the absence of oxygen. Interestingly, SO0054 displayed increased expression in the wild-type cells compared to that in the mutant cells upon transfer to anoxia (Table 2). SO0054 is assigned to a family of conserved hypothetical proteins that belong to COG2081 (predicted flavoproteins) and IPR013027 (FAD-linked oxidoreductases). Future work is needed to elucidate a possible role for SO0054 during transitions to anoxic growth conditions.

S. oneidensis MR-1 has over 40 putative cytochrome *c* genes, and two putative monoheme cytochrome *c* genes (SO4047 and SO4048) located in an operon were up-expressed in wild-type cells but not in mutant cells after transfer to anoxia (Table 2). The predicted molecular weights are 38 and 21 kDa, respectively. A putative decaheme cytochrome *c* gene (SO4360) has a predicted molecular weight of 33.1 kDa, and expression for SO4360 was lower in the mutant culture transferred to anoxia than in the wild type ($\log_2 R = -0.91$; *z* score = -2.10). Recently, SO4360 was predicted to be a *mirA* paralog, but a specific function for the cytochrome is not known (7).

Recent work demonstrated that *S. oneidensis* MR-1 can utilize both L- and D-lactate, and it was hypothesized that SO1518 to SO1520 encodes an L-lactate dehydrogenase and SO1521 encodes a D-lactate dehydrogenase (40). In our study, these

TABLE 2. Comparison of log₂ expression levels for selected genes when wild-type or mutant cells were compared between aerobic and anoxic conditions, respectively^a

Gene	Gene annotation	Log ₂ expression (z score) when transferred to anoxia	
		Wild-type culture	ΔSO3389 culture
SO0054	Conserved hypothetical (oxidoreductase)	1.55 (2.23)	0.06 (0.10)
SO0399 (<i>frdB</i>)	Succinate dehydrogenase/fumarate reductase	<u>-1.35 (-2.09)</u>	0.17 (0.31)
SO0583 (<i>bfd</i>)	Bacterioferritin-associated ferredoxin	2.89 (3.15)	0.67 (1.01)
SO0768	Flavin reductase	2.90 (3.36)	0.63 (1.38)
SO1329 (<i>cyaA</i>)	Adenylate cyclase	2.11 (2.88)	0.41 (0.73)
SO1427	Decaheme cytochrome <i>c</i> DMSO reductase subunit	<u>-3.30 (-2.64)</u>	0.79 (1.16)
SO1519	L-Lactate dehydrogenase (subunit)	<u>-0.67 (-1.63)</u>	0.15 (0.40)
SO1520	FeS cluster protein	<u>-1.05 (-1.83)</u>	-0.38 (-0.75)
SO1521	D-Lactate dehydrogenase	<u>-2.17 (-2.51)</u>	-0.76 (-1.42)
SO1522	Lactate permease	<u>-2.37 (-3.26)</u>	<u>-1.46 (-1.70)</u>
SO1927 (<i>sdhC</i>)	Succinate dehydrogenase	1.29 (1.59)	0.24 (0.63)
SO1928 (<i>sdhA</i>)	Succinate dehydrogenase/fumarate reductase	1.06 (1.12)	0.12 (0.22)
SO1929 (<i>sdhB</i>)	Succinate dehydrogenase/fumarate reductase	0.77 (1.78)	0.34 (1.09)
SO1930 (<i>sucA</i>)	2-Oxoglutarate dehydrogenase (E1)	2.01 (2.98)	0.90 (1.72)
SO1931 (<i>sucB</i>)	2-Oxoglutarate dehydrogenase (E2)	1.43 (2.59)	0.63 (1.34)
SO1932 (<i>sucC</i>)	Succinyl-CoA synthase (β)	1.33 (2.16)	0.79 (1.45)
SO2097 (<i>hydC</i>)	Ni/Fe hydrogenase	<u>-3.22 (-2.41)</u>	-0.53 (-0.89)
SO2098 (<i>hyaB</i>)	Ni/Fe hydrogenase	<u>-3.46 (-2.49)</u>	1.22 (1.04)
SO3285 (<i>cydB</i>)	Cytochrome oxidase	<u>-3.24 (-2.97)</u>	-1.14 (-1.68)
SO3286 (<i>cydA</i>)	Cytochrome oxidase	<u>-2.69 (-3.02)</u>	<u>-0.68 (-1.17)</u>
SO4047	Mono-heme cytochrome <i>c</i>	0.86 (2.03)	0.45 (1.29)
SO4048	Mono-heme cytochrome <i>c</i>	0.91 (2.16)	0.20 (0.70)

^a Bolding, up-expressed genes; underlining, down-expressed genes ($|z \text{ score}| > 1.6$). The time of sampling for anoxic culture is denoted by the arrow in Fig. 5.

genes were down-expressed in wild-type but not mutant cells when transitioned to anoxia compared to in aerobic cells. The Pinchuk et al. (40) study reported that these genes functioned under both aerobic and anaerobic conditions, and the decreased expression observed in our study might be a result of a decline in lactate consumption under anoxic conditions compared to that under aerobic conditions. The results suggested that SO3389 plays a potential role in down-expressing genes involved in lactate consumption when oxygen becomes limited.

The *fccA* (SO0970) gene in *S. oneidensis* has been shown to be a major soluble fumarate reductase during anaerobic growth (39), but *fccA* expression was not significantly up-expressed at the transcriptional level in wild-type or mutant cultures under the tested conditions. However, recent work has shown that FccA might be involved in multiple periplasmic electron transfer networks (48). It should be noted that gene expression was done with cultures that had been grown aerobically and then transferred to anoxic conditions; therefore, gene expression for some genes may differ from those reported in previous studies that have used cultures grown in pre-reduced, anaerobic medium for longer time periods. The current study was focused on the transition from aerobic to anoxic conditions, for which the mutant was deficient.

Wild-type cells had slightly elevated expression levels for a succinate dehydrogenase/fumarate reductase operon (*sdh*; SO1927 to SO1929) compared to those of the mutant, and the mutant also did not up-express components of the 2-oxoglutarate dehydrogenase (*sucAB*) or succinyl coenzyme A (succinyl-CoA) synthase (*sucC*) (Table 2). These tricarboxylic acid cycle enzymes were up-expressed in wild-type cells transferred to anoxia compared to in aerobic cells and may be a result of carbon flow from the added glutamate. However, mutant cells

did not up-express the same genes when transferred to anoxia and further indicated that the mutant was deficient in the ability to respond to anoxic conditions.

A presumptive bacterioferritin-associated ferredoxin (SO0583), a flavin reductase (SO0768), and a presumptive adenylate cyclase (SO1329) were also up-expressed in wild-type but not mutant cells when transferred to anoxia (Table 2). The class III adenylate cyclase was recently shown to regulate anaerobic respiration in *S. oneidensis* MR-1, and separate studies have shown that cyclic AMP (cAMP) levels were involved in the regulation of anaerobic respiration (8, 44). Hence, SO3389 may be a sensor involved in the global regulation of different modulators for the up- and down-expression of genes related to shifts between aerobic and anoxic metabolism.

Fumarate reductase activity. Because the transition to anoxic conditions was done in the presence of lactate and fumarate, fumarate reductase activities were measured. During the transition, wild-type cells increased fumarate reductase activity more than mutant cells as detected with activity gels, and this further corroborated the observation that the reductase activity necessary for anoxic growth was not up-expressed in mutant cells during the transition to anoxia. A small amount of activity was detected in the mutant cultures, but this activity was similar to the levels observed in aerobic cultures. In addition, fumarate reductase activity was quantified in wild-type and mutant cultures spectrophotometrically during the transition to anoxia. After transfer to anoxic medium, activity was not detected for the mutant, compared to 0.76 ± 0.09 mM benzyl viologen $\text{min}^{-1} \mu\text{g}^{-1}$ for wild-type cultures.

Structural similarity to PAS O₂/redox sensors. A role in O₂/redox sensing for SO3389 coincides with the strong evidence that PAS domains can be sensors for gases (e.g., O₂)



FIG. 8. Alignment of *M. capsulatus* MmoS PAS1 (amino acids 85 to 201), *M. capsulatus* MmoS PAS2 (amino acids 209 to 329), *A. vinelandii* NifL PAS (amino acids 21 to 140), *E. coli* Dos PAS (amino acids 16 to 133), *B. japonicum* FixL PAS (amino acids 151 to 257), SO3389 PAS1 (amino acids 204 to 270), and SO3389 PAS2 (amino acids 327 to 441). Hydrophobic or small residues are colored red, acidic residues are colored cyan, basic residues are colored magenta, and residues with hydroxyl or amine are colored green. Important residues for FAD ligand interactions in MmoS and NifL are denoted by underlining and bolding, and similar residues are underlined in SO3389 PAS2. Approximate location of β -sheets and α -helices are denoted based upon the MmoS crystal structure (3EWK) (58).

and/or intracellular redox potentials (53). For example, the PAS protein NifA has been shown to respond to O₂ via a flavin adenine dinucleotide (FAD) (6), and the *E. coli* Dos protein (EcDos) is a heme-containing gas sensor that has N-terminal PAS domains (12). PAS domain families comprise a protein clan with divergent sequences (15); however, a recent study characterized the structures of PAS domains with known and unknown cofactors and showed a broadly conserved structure that is comprised of a five-stranded antiparallel β -sheet and several α -helices for all PAS domains characterized to date (34).

The NifL sensor has been shown to be a reversible, FAD-based O₂/redox sensor that helps control the expression of nitrogen fixation genes in *Azotobacter* (21, 26). More recently, the MmoS in *Methylococcus capsulatus* (strain Bath) has been shown to be an FAD-based PAS sensor that coordinates responses to Cu via changes in redox (55). The SO3389 PAS1 domain did not have significant (E value of >0.1) amino acid sequence identity with characterized PAS domains but had 50% amino acid sequence identity (E value of 4.7e-12; SMART database) with a hypothetical protein (VP0092) in *Vibrio parahaemolyticus*. The SO3389 PAS2 domain had the highest peptide sequence identity (42% [E value of 4e-23]) with the MmoS PAS1 domain from *M. capsulatus* (followed by 38% [3e-12] with EcAer PAS [*E. coli*], 31% [2e-20] with AzNifL PAS [*Azotobacter vinelandii*], 28% [1e-13] with MmoS PAS2 [*M. capsulatus*], and 23% [4e-06] with EcDos PAS [*E. coli*]). Important residues for FAD binding were previously predicted from the crystal structure of MmoS (55), and the SO3389 PAS2 peptide sequence shared 6 of these MmoS residues and also 5 of the 10 conserved amino acids for FAD binding in the *A. vinelandii* NifL (Fig. 8).

In order to compare SO3389 to known PAS domain structures, models were predicted with CHPModels and I-Tasser tools. Both models were similar, and the I-Tasser model was

used for analysis. The I-Tasser model for the SO3389 PAS1/2 domains had a C score of 0.8, a TM score of 0.7 ± 0.1, and an experimental RMSD (root mean square deviation) of 5.6 ± 3.5 (positive C scores and TM scores of >0.5 indicate models with correct topology) (43). Moglich et al. (34) used z scores from structural superpositions to construct a dendrogram based upon the degree of structural diversity between PAS domains from different proteins, and PAS domains with known cofactors (i.e., FMN, FAD, and heme) formed separate clusters. Therefore, we compared the predicted model of SO3389 PAS1/2 to known PAS sensors to gain insight into possible functions of SO3389 in conjunction with the presented phenotypic data.

The I-Tasser model (<http://zhanglab.cmb.med.umich.edu/I-TASSER>) predicted the closest match with 3EWK from *M. capsulatus* (91% coverage; z score of 4.91) (Fig. 9). When the DALI Protein Structure Database (version 3.0; http://ekhidna.biocenter.helsinki.fi/dali_server/start) was used, the closest matches with the SO3389 PAS1/2 model were the FAD-binding MmoS PAS1 from *M. capsulatus* (3EWK) and the NifL PAS1 (2GJ3) from *A. vinelandii* (RMSD of 0.6 Å, z score of 24.9; RMSD of 1.4 Å, z score of 15.4, respectively). In addition, the Phyre server (<http://www.sbg.bio.ic.ac.uk/~phyre/index.cgi>) predicted that 2GJ3 was the closest match based upon protein sequence and structure (E value of 2.3e-15; 30% sequence identity). Both 2GJ3 and 3EWK are known FAD-based redox PAS sensors (26, 51, 55).

The SO3389 ORF also contains putative GGDEF and EAL domains (originally named for the conserved amino acid residues) (9, 24, 38). The two domains are thought to interact with cyclic-diguanosine monophosphate (c-di-GMP), and c-di-GMP was recently proposed to be a global secondary messenger involved in regulating cell-cell and cell-surface interactions (9). The GGDEF domain was recently shown to be associated with the synthesis of c-di-GMP via diguanylate cyclase (DGC)

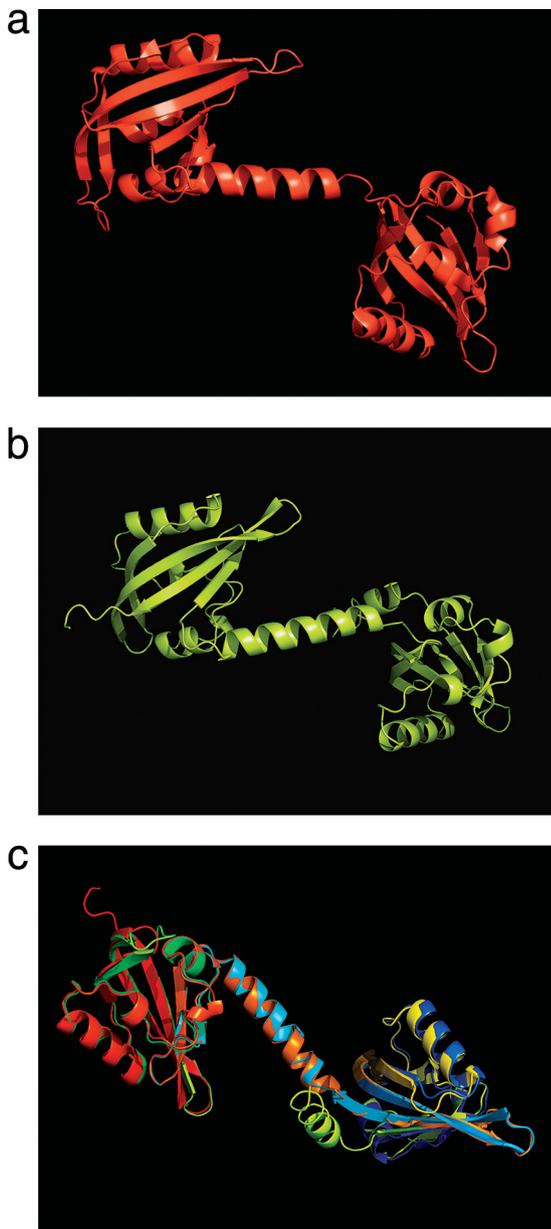


FIG. 9. Structure of the *M. capsulatus* MmoS (a) (PDB:3EWK) and SO3389 (b) PAS domains and the structural superposition (c). The SO3389 PAS domain model was predicted with I-Tasser, and the structural alignment was visualized with PyMol (www.pymol.org).

and the EAL domain to contain phosphodiesterase (PDE) activity in *Yersinia*, *Caulobacter*, and *Salmonella* (9, 16, 27, 24, 38) and more recently in *Shewanella* (54). In addition, the involvement of DGC and PDE domains in cell metabolism and cell behavior has recently been shown in *Burkholderia*, *Xanthomonas*, *Streptococcus*, *Vibrio*, and *Pseudomonas* (for examples, see references 5 and 30). Interestingly, sequence comparisons suggested that SO3389 was a phosphodiesterase and not a cyclase based upon conserved amino acids predicted for active phosphodiesterases (data not shown) (47).

Our data indicated that SO3389 was involved in sensing DO/redox levels that affected both aerobic growth and transi-

tions to anoxia. The resultant phenotypes were a consequence of the inability to regulate genes that included cytochromes and reductases. The sequence and structure comparisons suggested that SO3389 could be a FAD-based O₂/redox sensor, and future work includes the characterization of ligand interactions for SO3389. It is interesting to speculate that the orthologs identified in the other genera (e.g., *Vibrio*, *Cyanothecce*, *Endoriftia*) (Fig. 2) could function in a similar manner to help cells control growth associated with O₂ and/or anoxia.

The most similar gene sequences to that of SO3389 were annotated as conserved hypothetical and sensory box proteins, and thus SO3389 encodes a novel protein for which a function has not been previously described. Numerous systems biology studies have demonstrated the often realized but unknown importance of hypothetical and conserved hypothetical proteins in microorganisms (13), and the elucidation of function for these proteins provides great insight into the ability of microorganisms to sense external stimuli and optimize metabolism.

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