

Sulfite Reduction in Mycobacteria[∇]

Rachel Pinto,¹ Joseph S. Harrison,¹ Tsungda Hsu,² William R. Jacobs, Jr.,² and Thomas S. Leyh^{1*}

Department of Biochemistry¹ and Howard Hughes Medical Institute and Department of Microbiology and Immunology,² Albert Einstein College of Medicine, 1300 Morris Park Ave., Bronx, New York 10461-1926

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***Mycobacterium tuberculosis* places an enormous burden on the welfare of humanity. Its ability to grow and its pathogenicity are linked to sulfur metabolism, which is considered a fertile area for the development of antibiotics, particularly because many of the sulfur acquisition steps in the bacterium are not found in the host. Sulfite reduction is one such mycobacterium-specific step and is the central focus of this paper. Sulfite reduction in *Mycobacterium smegmatis* was investigated using a combination of deletion mutagenesis, metabolite screening, complementation, and enzymology. The initial rate parameters for the purified sulfite reductase from *M. tuberculosis* were determined under strict anaerobic conditions [$k_{\text{cat}} = 1.0 (\pm 0.1)$ electron consumed per second, and $K_{m(\text{SO}_3^{2-})} = 27 (\pm 1) \mu\text{M}$], and the enzyme exhibits no detectible turnover of nitrite, which need not be the case in the sulfite/nitrite reductase family. Deletion of sulfite reductase (*sirA*, originally misannotated *nirA*) reveals that it is essential for growth on sulfate or sulfite as the sole sulfur source and, further, that the nitrite-reducing activities of the cell are incapable of reducing sulfite at a rate sufficient to allow growth. Like their nitrite reductase counterparts, sulfite reductases require a siroheme cofactor for catalysis. Rv2393 (renamed *che1*) resides in the sulfur reduction operon and is shown for the first time to encode a ferrochelatase, a catalyst that inserts Fe^{2+} into siroheme. Deletion of *che1* causes cells to grow slowly on metabolites that require sulfite reductase activity. This slow-growth phenotype was ameliorated by optimizing growth conditions for nitrite assimilation, suggesting that nitrogen and sulfur assimilation overlap at the point of ferrochelatase synthesis and delivery.**

The human genome does not encode the sulfur reduction and cysteine-biosynthetic enzymes found in many pathogenic bacteria. The species specificity of these enzymes and the essential metabolic nature of sulfur recommend them as potential targets for antimicrobial development. A considerable literature links sulfur metabolism to the pathogenicity and antibiotic susceptibility of *Mycobacterium tuberculosis*. Specific sulfolipids correlate well with the virulence of *M. tuberculosis* (16, 17, 19, 20, 26) and are reported to inhibit phagosome-lysosome fusion (18), which is critical for the survival of the bacterium in macrophages (9, 15). More-recent literature calls into question whether the segregation of sulfolipid across virulent and avirulent strains is a manifestation, rather than a root cause, of virulence (33, 41), and efforts to trace this lineage toward its root are under way (11, 27). Mycothiol, the mycobacterial equivalent of glutathione, utilizes a cysteine thiol to provide the antioxidant protection that the organism needs to survive, particularly during oxidative stress (35), and lower mycothiol levels correlate with enhanced susceptibility to antibiotics, including rifampin and isoniazid (32). The cysteine-biosynthetic pathway, a primary means of assimilating sulfur, has been linked to the survival of the organism during the chronic phase of infection, the basis of which may lie in its resistance to reactive oxygen and nitrogen species (39). The current work explores the reduction of sulfite, an essential step

in the biosynthesis of cysteine, in a model organism, *Mycobacterium smegmatis*.

Mycobacterial assimilation of sulfate begins with its active transport into the cell, whereupon it is chemically activated by the enzyme ATP sulfurylase (*cysDN*) to form activated sulfate (adenosine 5'-phosphosulfate [APS]) (Fig. 1). APS is then either phosphorylated by APS kinase (*cysC*) to form the universal sulfuryl group donor 3'-phosphoadenosine 5'-phosphosulfate (PAPS) or reduced by APS reductase (*cysH*) to form sulfite (12, 38). If sulfate assimilates through the sulfuryl transfer branch of the pathway, the sulfuryl group is transferred from PAPS, via sulfotransferases, to metabolic recipients whose activities are regulated by the modification. If, on the other hand, sulfate is drawn into the reductive branch of the pathway by the action of APS reductase (7, 8), the resulting sulfite is reduced further, in a six-electron reduction, to sulfide by the enzyme sulfite reductase (*sirA*). Sulfide is then incorporated into cysteine by *O*-acetyl-L-serine sulfhydrylase (*cysKI*) (30), and from there the sulfur atom, originally present in sulfate, flows into a myriad of reduced-sulfur-containing metabolites.

Nitrite and sulfite reductases have a number of similarities, including conserved catalytic architectures designed to guide an essential siroheme cofactor to function as the active center for a six-electron reduction of substrate to product (12, 13, 22). These enzymes are sufficiently similar that they will often catalyze the reduction of both sulfite and nitrite (12), suggesting that sulfur and nitrogen metabolism may be redundant at the point of sulfite/nitrite reduction and/or in the provision of the siroheme cofactor. Such redundancy is particularly pertinent in the case of *M. tuberculosis*, which must persist amid high levels of destructive reactive nitrogen species, which it elicits from activated T cells during infection (40).

* Corresponding author. Mailing address: Department of Biochemistry, Albert Einstein College of Medicine, 1300 Morris Park Ave., Bronx, NY 10461-1926. Phone: (718) 430-2857. Fax: (718) 430-8565. E-mail: leyh@aecom.yu.edu.

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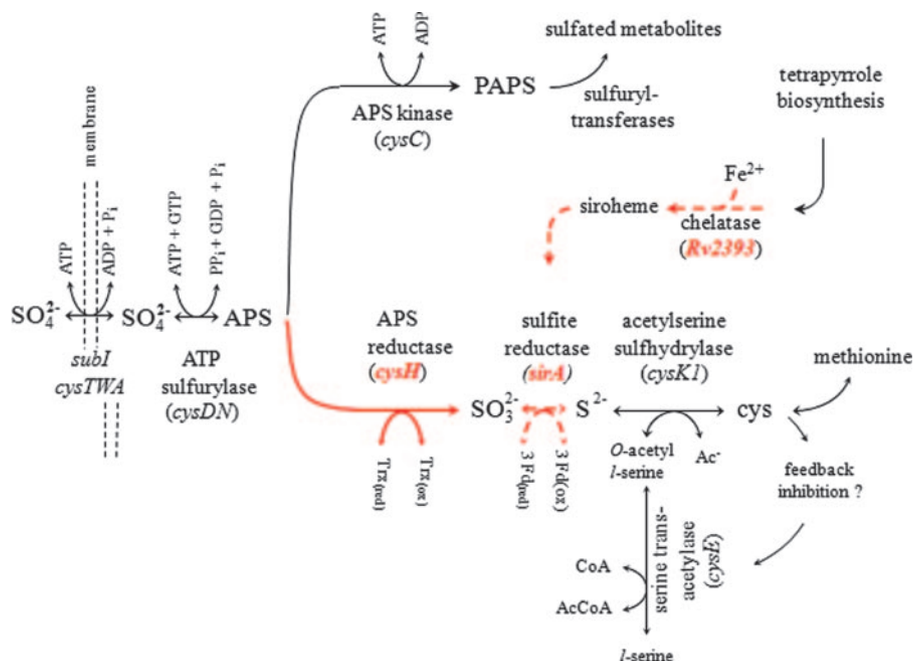


FIG. 1. Segment of sulfur metabolism in mycobacteria. Red arrows and mnemonic terms identify the activities and genes associated with the sulfur reduction operon. Dotted arrows highlight steps that are the primary focus of this study. The cysteine feedback inhibition loop is hypothesized based on similar inhibition in *Salmonella enterica* serovar Typhimurium (25) and *E. coli* (21) and complete conservation of the residues involved in the binding of cysteine (28, 29) across numerous organisms, including *M. tuberculosis* (29). Ac⁻, without acetyl; CoA, coenzyme A; AcCoA, acetyl coenzyme A; 3Fd(ox), oxidized ferredoxin; 3Fd_(red), reduced ferredoxin; Trx_(ox), oxidized thioredoxin; Trx_(red), reduced thioredoxin.

The genes encoding APS reductase (*cysH*) and sulfite reductase (*sirA*) are located in the sulfur reduction operon (see below) in both *M. smegmatis* and *M. tuberculosis*, along with a third coding region, Rv2393, which encodes a protein of unknown function. In the current study, the sulfite reduction step in *M. smegmatis* was explored by deletion mutagenesis, metabolite screening, enzymatic characterization, and complementation.

MATERIALS AND METHODS

Bacterial strains and cultures. The bacterial strains used in this study are listed in Table 1. The *Escherichia coli* XL1-Blue and HB101 strains, used for cloning, and the BL21(DE3) Rosetta pLysS strain, used for expression, were propagated in Luria-Bertani (LB) broth or on LB agar at 37°C. *M. smegmatis* strain mc²155 was grown at 37°C in 7H9 liquid medium (Difco) supplemented with 0.2% (vol/vol) glycerol, 0.2% glucose, and 0.05% Tween 80 or on Middlebrook 7H10 solid medium supplemented as described above. L-Cysteine was obtained from the Sigma Chemical Co. The following antibiotics were used at the

TABLE 1. Bacterial strains used in this study

Strain	Description	Source or reference
<i>E. coli</i> strains		
XL1-Blue	<i>recA1 endA1 gyrA96 thi-1 hsdR17 supE44 relA1 lac</i> [F' <i>proAB lacI</i> ^q Δ <i>M15</i> Tn10 (Tet ^r)]	Stratagene
BL21(DE3) Rosetta pLysS	F' <i>ompT hsdS_B</i> (r _B ⁻ m _B ⁻) <i>gal dcm</i> (DE3)pLysSRARE ² (Cam ^r)	Novagen
JR01	Rosetta pLysS carrying plasmid pJR1	This work
302Δa	<i>E. coli cysG</i> Nir ^s Lac ⁺ CysG ⁻	31
cysG pER247 strain	Strain 302Δa carrying the plasmid pER247 (pACYC184- <i>lac</i> ^q - <i>Ptac</i> - <i>P. denitrificans cobA</i>)	31
RP7	<i>E. coli cysG</i> pER247 containing plasmid pET23a, <i>M. tuberculosis</i> Rv2393	This work
RP8	<i>E. coli cysG</i> pER247 containing plasmid pET23a, <i>M. smegmatis</i> Rv2393	This work
HB101	<i>E. coli</i> K-12 F ⁻ Δ(<i>gpt-proA</i>)62 leuB1 glnV44 ara-14 galK2 lacY1 hsdS20 rpsL20 xyl-5 mtl-1 <i>recA13</i>	4
<i>M. smegmatis</i> strains		
mc ² 155	<i>ept-1</i>	43
RPMS001	mc ² 155Δ <i>sirA</i> :: <i>hyg sacB</i>	This work
RPMS002	mc ² 155ΔRv2393:: <i>hyg sacB</i>	This work
RPMS004	RPMS001 carrying plasmid pRP15 (<i>M. smegmatis sirA</i> complementation vector [Table 2])	This work
RPMS005	RPMS002 carrying plasmid pRP16 (<i>M. smegmatis che1</i> complementation vector [Table 2])	This work

TABLE 2. Plasmids used in this study

Plasmid	Description	Reference or source
p0004S	Cosmid containing the Hyg ^r -SacB cassette	2
phAE159	Shuttle phasmid, TM4ts::pYUB328	J. Kriakov and W. R. Jacobs, unpublished
pMV261	Kan ^r ColE1 <i>oriM aph P_{hsp60}</i>	44
pMV361	Integrative vector Hyg ^r	44
pMS2391	Recombinant cosmid derived from p0004S containing the <i>M. smegmatis</i> <i>sirA</i> AES	This study
pMS2393	Recombinant cosmid derived from p0004S containing the <i>M. smegmatis</i> Rv2393 AES	This study
pSKB4-9His-GST	Derived from pGEX-6P-1; contains an N-terminal His tag upstream of a GST tag	1
pJR1	pSKB4-9His-GST carrying the <i>M. tuberculosis</i> <i>sirA</i> gene	This study
pJR2	pJR1 with <i>E. coli</i> <i>cysG</i> cloned downstream of <i>sirA</i>	This study
pRP15	pMV261 carrying the <i>M. smegmatis</i> <i>sirA</i> gene	This study
pRP16	pMV261 carrying the <i>M. smegmatis</i> Rv2393 gene	This study
pET23a	Cloning vector containing an N-terminal His tag	Novagen
pRP17	pET23a carrying <i>M. tuberculosis</i> Rv2393	This study
pRP18	pET23a carrying <i>M. smegmatis</i> Rv2393	This study
pKK <i>E. coli</i> <i>cysG</i>	pKK223.3, an overexpression vector derived from pBR322 with the <i>tac</i> promoter, carrying <i>E. coli</i> <i>cysG</i>	31

concentrations indicated: ampicillin (100 µg/ml) (Fisherbiotech), kanamycin A (25 µg/ml) (Sigma), hygromycin B (50 µg/ml with *E. coli* and 50 to 150 µg/ml with *M. smegmatis*) (Roche).

To obtain the growth curves associated with Fig. 4, *M. smegmatis* was grown to late exponential phase at 37°C; washed six to eight times in M9 minimal salts without a sulfur source (Na₂HPO₄ [42 mM], KH₂PO₄ [24 mM], NaCl [9.0 mM], NH₄Cl [19 mM], glucose [0.5%, wt/vol], Tween 80 [0.05%, vol/vol]); suspended in prewarmed M9 medium supplemented with a specific source of sulfur, SO₄²⁻ (1.0 mM), Na₂S (1.0 mM), Na₂SO₃ (1.0 mM), or cysteine (200 µM); and cultured further by shaking at 37°C. Bacterial growth was monitored at 600 nm. To normalize the wild-type and mutant genetic backgrounds in the strains used in Fig. 4C, the wild-type strain was transformed with pMV361, which codes for Hyg^r and integrates into the genome (44), and pMV261, which confers kanamycin resistance. The mutant strains associated with Fig. 4C were transformed using derivatives of pMV261 that contained either the wild-type *sirA* gene (pRP15) or Rv2393 (pRP16).

The cell growth studies presented in Fig. 6 were accomplished by shaking cells overnight at 37°C in the following M9 minimal medium: NaNO₃ (20 mM), cysteine (200 µM), Na₂HPO₄ (42 mM), KH₂PO₄ (24 mM), NaCl (9.0 mM), glucose (0.5%, wt/vol), Tween 80 (0.05%, vol/vol). The cells were then washed as described above in prewarmed M9 medium lacking a sulfur source. The growth studies were initiated by suspending the washed cells in prewarmed M9 medium with either cysteine or sulfate.

Ferrocyclase complementation. To assess whether Rv2393 encodes a ferrocyclase, an *E. coli* cysteine auxotroph that requires ferrocyclase function for growth on SO₄²⁻ (strain 302Δa, a *cysG* deletion strain containing pER247, a P15

origin plasmid that expresses uroporphyrinogen III methyltransferase) was transformed with ColE1 origin plasmids that express Rv2393 either from *M. tuberculosis*(pRP17) or *M. smegmatis*(pRP18) and then tested for the ability to grow on SO₄²⁻ as a sole source of sulfur. Conversion to prototrophy was assessed on minimal agar containing either SO₄²⁻ or cysteine as the sulfur source. Minimal agar contained Na₂HPO₄ (42 mM), KH₂PO₄ (24 mM), NaCl (9.0 mM), NH₄Cl (19 mM), agar (15 g/liter), CaCl₂ (0.10 mM), MgCl₂ (1.0 mM), glucose (2.0%), and either MgSO₄ (2.0 mM) or cysteine (280 µM). 302Δa carrying pKK (which expresses *E. coli* CysG) was used as the positive control for growth; the negative-growth-control strain was 302Δa, which carries pER247 and pET23a (the empty Rv2393 expression vector). All plates were incubated at 37°C for 24 to 48 h.

DNA manipulation. Restriction enzymes (REs) were purchased from New England Biolabs, and digestions were performed according to the manufacturer's recommendations. The purification of DNA from agarose gels and the isolation of plasmid DNA were done using QIAquick gel extraction and QIAprep spin miniprep kits (QIAGEN) according to the manufacturer's protocols. Isolation of *M. tuberculosis* and *M. smegmatis* chromosomal DNA was carried out as described previously (5). Standard heat shock protocols were used for the transformation of *E. coli* strains (34).

Plasmid construction. All of the plasmids and primers used in this study are listed in Tables 2 and 3, respectively.

***sirA* expression vector.** The expression of catalytically competent SirA requires the coexpression of CysG, which produces the quantities of the siroheme cofactor needed for stoichiometric incorporation into SirA. The construction of the coexpression plasmid pJR2 was accomplished in two steps. First, a 1.7-kb fragment containing *sirA* was PCR amplified from *M. tuberculosis* genomic DNA

TABLE 3. Primers used in this study

Primer ^a	Sequence (5'–3')	Amplified sequence
TBsirA F	GGAATTCATATGTCCGCGAAGGAGAACCC	<i>M. tuberculosis</i> <i>sirA</i>
TBsirA R	AAGGAAAAAGCGGCCGCTCATCGCAGGTCGCTCCTCGGCCCGGAT	
colicysG F	TTTGCGGCGCAAACGGCTGCCGGTTAATTAATAAGGGGTTTTTAC	<i>E. coli</i> <i>cysG</i>
colicysG R	GGCCTAGGTTAATGGTTGGAGAACCAGTTCAGTTTATC	
smegsirA F	TACAGCTGATGCTCGAAGACGAGTACTTCAT	<i>M. smegmatis</i> <i>sirA</i>
smegsirA R	CCCAAGCTTTCACGTTGCCTACCTCAAATCCGCTTCGTC	
smegRv2393 F	CGCAGCTGGTGACGCTCGTCTGACCGCAC	<i>M. smegmatis</i> Rv2393
smegRv2393 R	CCCAAGCTTTCAGCGCATGGCCCTGGCCCGTGCAT	
1	TTTTTTTTCCATAAATTTGGGTAGATGCCGAAGGGCCCGCCCGGTGTGGG	3'-end-flanking region of
1'	TTTTTTTTCCATTTCTTTGGGCCGACGGCGTCGAGGCCGCTTCCAGATCTC	<i>M. smegmatis</i> <i>sirA</i>
2	TTTTTTTTCCATAGATTGGGGCCCGCAGGAGGGTTTCCAGGTGCATCTG	5'-end-flanking region of
2'	TTTTTTTTCCATCTTTTGGGCCCGCATTGCGTCTGGACAGCCCGGCC	<i>M. smegmatis</i> <i>sirA</i>
3	TTTTTTTTCCATAAATTTGGGTGCGCAACTTCGTGAAACAACGCGAGGAC	3'-end-flanking region of
3'	TTTTTTTTCCATTTCTTTGGAGATTCGGTTCGTTCTGCTCACAGAAC	<i>M. smegmatis</i> Rv2393
4	TTTTTTTTCCATAGATTGGTTCGCGCTAGGACGAGGATCGAGG	5'-end-flanking region of
4'	TTTTTTTTCCATCTTTTGGTATCGGTTTCGTGTTCCAGCACTATGCGGC	<i>M. smegmatis</i> Rv2393

^a F and R refer to forward and reverse primers.

TABLE 4. Primary sequence similarity of SirA from *M. smegmatis* and *M. tuberculosis*^a

Enzyme	Coding region	Identity (%)	Similarity (%)
Sulfite reductase	<i>sirA</i>	82	89
APS reductase	<i>cysH</i>	70	78
Hypothetical protein	Rv2393	58	70

^a CLUSTAL W comparison metrics for homologous proteins encoded by the *M. smegmatis* and *M. tuberculosis* sulfur reduction operons (45).

using primers TBsirA F and TBsirA R and inserted into the NdeI and NotI sites of pSKB4 (1), yielding pJR1. In the second step, *E. coli cysG* was PCR amplified (primers colcysG F and colcysG R) and cloned into pJR1 linearized with NotI and BsaAI. The resulting plasmid, pJR2, was used to transform the *E. coli* BL21(DE3) Rosetta pLysS strain for expression and purification (see below). This strain was named JR01.

Complementation plasmids. *M. smegmatis sirA* and Rv2393 were PCR amplified from genomic DNA using the primer pairs smegsirA F/smegsirA R and smegRv2393 F/smegRv2393 R, respectively. In both cases, the forward and reverse primers introduced PvuII and HindIII restriction sites, respectively, which were used to subclone the PCR products into pMV261 (cleaved with the same enzymes), producing plasmids pRP15 (*sirA*) and pRP16 (Rv2393). pMV261 contains the mycobacterial *hsp60* promoter, which facilitates constitutive expression of *M. smegmatis* SirA and *M. smegmatis* Rv2393 (44). All constructs were sequenced (AECOM DNA sequencing facility) to confirm the fidelity of the clones.

Construction of *M. smegmatis* gene deletion mutants. The *M. smegmatis* gene deletion mutants were constructed in three stages: recombinant cosmids containing the DNA sequences needed for allelic exchange (allelic-exchange substrates [AESs]) were prepared, high-titer mycobacteriophages needed to isolate genomic deletion events (which occur at low levels) were obtained, and mycobacterial DNA was transduced and selected for allelic exchange. The protocols used to create mycobacterial gene deletion mutants have been described previously (5).

Cosmid construction. DNA flanking the 5' and 3' regions of *sirA* was PCR amplified, using primer pairs 1/1' and 2/2' (see Fig. 3A and Table 3). The flanking regions were subcloned directionally, using the Van911 RE, on either side of the hygromycin resistance-*sacB* gene cassette found in cosmid p0004S. The resulting recombinant cosmid, pMS2391, contained the AES needed to construct the bacteriophage. Using an identical protocol, primer pairs 3/3' and 4/4' were used to generate pMS2393, which contains the Rv2393 AES.

Mycobacteriophage construction. The AES cosmids and purified phagemid DNA (phAE159) were digested separately with PacI, ligated together, and in vitro packaged (using the GIGAPackIII Gold packaging extract [Stratagene]) to produce the transducing λ bacteriophage. *E. coli* HB101 was then transduced with the phage, and transductants were selected on medium containing hygromycin (150 μ g/ml). Phasmid construction was confirmed by PacI digestion prior to electroporation of the phasmid DNA into *M. smegmatis*. Electroporated cells were plated on 7H10 medium and incubated at the permissive temperature, 30°C, for 3 days. All transducing phages were plaque purified, and high-titer phage lysate (10^{10} to 10^{11} PFU/ml) was prepared. *M. smegmatis* was then transduced by mixing late-log-phase cells with phage lysate (1:1, vol/vol) overnight at 37°C. Transductants were plated onto 7H10 medium plates containing hygromycin (75 μ g/ml) and cysteine (40 μ g/ml) and incubated at the nonpermissive temperature, 37°C, for 5 days. *M. smegmatis* deletion mutants were screened by Southern blot analysis (see below).

Southern blotting. Five micrograms of *M. smegmatis* genomic DNA was digested completely with either the BanII or the MscI RE and separated by electrophoresis on a 0.7% DNA agarose gel. The gel was immersed in 0.25 M HCl for 5 min, rinsed in H₂O, and soaked in 0.40 M NaOH for 10 min. DNA fragments were transferred onto a nylon membrane (Amersham) overnight by the capillary action of a 0.40 M NaOH transfer solution. The membrane was cross-linked (Stratagene) to allow immobilization of DNA fragments on the membrane and then incubated for 10 min in 5 ml of Rapid-hyb buffer (Amersham). Changes in DNA fragment size caused by the deletions were probed using ³²P-labeled probes that spanned the 5' end of *sirA* and 3' end of Rv2393. The probes were generated by PCR using the 1/1' and 4/4' primer pairs (Table 3 and see Fig. 3A).

Expression and purification of *M. tuberculosis* SirA. LB medium was inoculated with an overnight culture of JR01 at an A_{600} of 0.01, and the cells were grown at 37°C until the optical density at 600 nm reached ~0.7, at which point SirA expression was induced by adding isopropyl- β -D-thiogalactopyranoside (IPTG) to a final concentration of 0.80 mM. The incubation temperature was then shifted to 17°C, and cells were harvested by centrifugation 17 h later. The cell pellet was suspended in 4.2 ml/g cell paste of lysis buffer (KPO₄ [50 mM, pH 8.0], KCl [0.4 M], phenylmethylsulfonyl fluoride [290 μ M], pepstatin A [1.5 μ M], lysozyme [0.10 mg/ml]), and the solution was stirred for 1 h at 4°C prior to sonication on ice (Branson sonifier). Cellular debris was removed by centrifugation (20 min at 31,000 \times g), and the supernatant was loaded onto a 5-ml-chelating-Sepharose fast-flow column charged with Ni²⁺ and equilibrated with buffer A (KPO₄ [50 mM, pH 7.3], KCl [0.4 M]). The column was washed with 10 bed volumes of buffer B (KPO₄ [50 mM, pH 7.3], KCl [0.4 M], imidazole [10 mM]). The His-glutathione S-transferase (GST)-tagged fusion protein was then eluted from the Ni²⁺-chelated Sepharose column with 10 bed volumes of buffer C (KPO₄ [50 mM, pH 7.3], KCl [0.4 M], imidazole [250 mM], KCl [0.40 M]). The N-terminal His-GST tag was removed by digestion with PreScission protease (GE Healthcare) during overnight dialysis against HEPES-K⁺ (25 mM, pH 7.5), KCl (100 mM), glycerol (5%), and β -mercaptoethanol (5.0 mM) at 4°C. To remove the tag and any uncut fusion protein, the dialyzed proteolysate was passed back over the Ni²⁺ affinity column and washed extensively with buffer B to remove the SirA, which exhibited an affinity for the resin. Typically, 10 mg of protein (>95% pure, as judged by eye from Coomassie blue-stained sodium dodecyl sulfate-polyacrylamide gels) was obtained per liter of culture.

Sulfite reductase assay. *M. tuberculosis* SirA was assayed for sulfite and nitrite reductase activity under anaerobic conditions using methyl viologen (MV) as an electron donor, which was reduced with Zn metal immediately prior to use (42). The assay mix contained *O*-acetyl-serine (5.0 mM), sodium sulfite (13 μ M), reduced MV (250 μ M), *O*-acetyl-serine sulfhydrylase (1.5 μ M), SirA (1.3 μ M), and 50 mM HEPES-K⁺ (pH 8.0) (25 \pm 3°C). The assay was carried out under anaerobic conditions; argon gas scrubbed with reduced MV and ascorbic acid was used to degas all buffers and to maintain all samples under positive pressure in a vacuum manifold. A custom optical cuvette that allowed maintenance of positive pressure while the mixture was stirred was used to optically monitor reactions. The assay was initiated by the addition of the sodium salts of sulfite or nitrite. The enzymatic consumption of either sulfite or nitrite was monitored by following the decrease in absorbance at 684 nm ($\epsilon = 4.8 \times 10^3$ M⁻¹ cm⁻¹) caused by the oxidation of MV (24).

RESULTS

Sulfur reduction operon—an overview. The *M. smegmatis* and *M. tuberculosis* sulfur reduction operons are quite similar (Fig. 1A; Table 4). At their 5' termini, they begin with *sirA* (sulfite reductase) followed by *cysH* (APS reductase) and Rv2393 (a hypothetical protein). In both organisms, each of the three coding regions appear to be translationally coupled (i.e., the stop codon of the 5' gene overlaps the start codon of its downstream partner). The homologous coding regions in the two strains are similar in both length and sequence (Table 4; Fig. 2) (45). The metabolically linked, sulfate transport operons, which include an ABC transporter (encoded by *cysTWA*) and a sulfate binding protein (encoded by *subI*), are antipar-

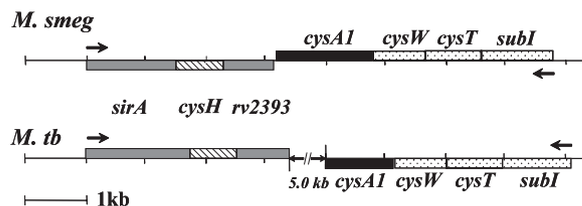


FIG. 2. Sulfur reduction operons of *M. smegmatis* and *M. tuberculosis*. (A) Schematic representation of the *M. smegmatis* (*M. smeg*) and *M. tuberculosis* (*M. tb*) *sirA* operons. The double hatch mark in the *M. tuberculosis* diagram indicates a 5-kb intergenic region between the operons.

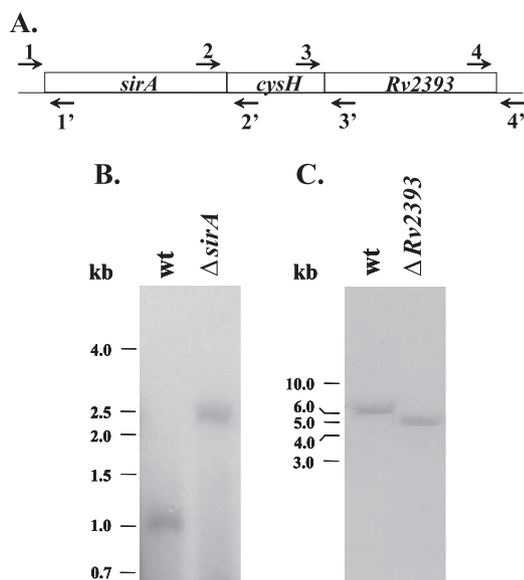


FIG. 3. Insertion of a hygromycin resistance cassette into *sirA* and *Rv2393* in *M. smegmatis* mc²155. (A) Primer sets used for the construction of Southern blot probes and AESs. (B) Southern blot of genomic DNAs from wild-type (wt) and Δ *sirA* strains. *Ban*II-digested genomic DNA was probed using a PCR fragment generated using primers 1 and 1'. The blot revealed that insertion of the resistance cassette caused the expected shift from 1.1 to 2.5 kb. (C) Southern blot of genomic DNAs from the wild-type (wt) and Δ *Rv2393* strains. *Msc*I-digested genomic DNA was probed using a PCR fragment generated using primers 4 and 4'. Insertion of the resistance cassette resulted in the expected shift from 6.3 to 5.0 kb.

allel to the sulfur reduction operon in both cases but differ in that they are separated by \sim 5 kb in *M. tuberculosis* and only 10 bp in *M. smegmatis*.

Construction of *sirA* and *Rv2393* deletion mutants. To begin to assess the role of *sirA* and *Rv2393* in vivo, each gene in *M. smegmatis* was deleted using allelic exchange. First, allelic-exchange cosmids containing a selectable gene cassette ($\gamma\delta$ *res hyg sacB res* $\gamma\delta$) flanked by \sim 900-bp regions of DNA that spanned the 3' and 5' edges of *sirA* or *Rv2393* were constructed (see Materials and Methods). The AESs were then incorporated into the conditionally replicating phasmid pHA159, in vitro packaged into λ phage heads, and transduced into *E. coli* HB101. Recombinant shuttle cosmids were purified from Hyg^r *E. coli* transductants and converted into mycobacteriophage-packaged DNA molecules by transfecting them into *M. smegmatis* cells. These cells were plated for phage plaques at the permissive temperature of 30°C. High-titer transducing lysates were obtained by propagation of the mycobacteriophage in *M. smegmatis*. The lysates were used to transduce *M. smegmatis* at the nonpermissive temperature (37°C) and plated on selective medium. Restriction of phage replication results in a double-crossover event between the homologous DNA arms flanking the disrupted gene (5). Fifteen transductants were obtained in the case of *sirA* and six in the case of *Rv2393*. Six isolates from each set were tested by Southern blot analysis, all of which had undergone the expected allelic replacement. A single representative from each set is shown in Fig. 3B. The observed shifts in DNA fragment size caused by insertion of the Hyg^r-*SacB* gene cassette into the genomic copies of *sirA* and *Rv2393* are

in excellent agreement with the expected values—1.4 and 1.3 kb, respectively.

***sirA* is essential for growth in minimal media.** Upon entering the macrophage, *M. tuberculosis* maintains the expression of SirA, a sulfite reductase that catalyzes the six-electron reduction of sulfite (37). Frequently, both sulfite and nitrite can be reduced by either nitrite or sulfite reductases, which are classified on the basis of their substrate preference (12). Thus, a strain lacking *sirA* might obtain the sulfide needed for cysteine biosynthesis from the reduction of sulfite by other reductases, which would render *sirA* nonessential and perhaps diminish its efficacy as an antimicrobial target. To establish whether the organism depends essentially entirely on SirA for the provision of sulfide, the *M. smegmatis* *sirA* deletion mutant was tested for its ability to grow, relative to the wild type, on sulfur metabolites that straddle either side of the point of action of *sirA* in the cysteine-biosynthetic pathway. When the mutant and wild type were fed downstream metabolites (i.e., S²⁻ or cysteine), their growth rates were virtually indistinguishable; however, when fed an upstream metabolite (SO₄²⁻ or SO₃²⁻), the mutant showed no detectible growth over a 50-h incubation, while the wild type grew with a doubling time of 7.1 (\pm 0.4) h (Fig. 4A). *sirA* is clearly essential for growth on SO₄²⁻ or SO₃²⁻ and central to the sulfur-reducing metabolism of the organism.

SirA—the enzyme. The kinetic parameters and efficiency of the enzyme toward sulfite reduction have not yet been determined. For this reason, the enzyme from *M. tuberculosis* was expressed in *E. coli* and purified to homogeneity (see Materials and Methods), and the kinetic constants were determined. Obtaining pure SirA with the expected levels of siroheme cofactor (22) required coexpression of *E. coli* CysG, a trifunctional protein that catalyzes the last three steps in siroheme biosynthesis (see Materials and Methods). The reduction reaction was monitored continuously, under anaerobic conditions, by following the change in absorbance at 684 nm associated with the oxidation of the electron donor, MV (24). Sulfide produced by SirA was removed rapidly by serine sulfhydrylase (from *E. coli*), which converts *O*-acetyl-L-serine and sulfide to cysteine (46). The removal of the product ensures that product inhibition will not contribute significantly to the reaction, which simplifies extracting kinetic constants from the reaction progress curve (1). Under such conditions, slopes taken over sufficiently small regions (\sim 3%) of the progress curve provide initial rate measurements over an essentially continuously varying range of substrate concentrations. The double-reciprocal plot of the SirA-catalyzed reduction of sulfite is shown in Fig. 5. The kinetic constants obtained by fitting the data are as follows: $K_m(\text{SO}_3^{2-})$ was equal to 27 (\pm 1) μ M, the k_{cat} was equal to 0.17 (\pm 0.01) electron consumed s⁻¹ (sulfite reduced to sulfide) or 1.0 electron consumed s⁻¹, and the catalytic efficiency (k_{cat}/K_m) was equal to $3.7 \times 10^4/\text{M}^{-1} \text{s}^{-1}$. It should be noted that, consistently with previous work (38), no enzymatic turnover was observed when sulfite was replaced by nitrite at concentrations as high as 3.0 mM.

Growth phenotype of the *Rv2393* deletion mutant. *Rv2393* is situated at the C terminus of the sulfur reduction operon. This locale suggests that *Rv2393*, whose function is unknown, may be important for cysteine biosynthesis. To explore the ways in which *Rv2393* might function in the cysteine-biosynthetic path-

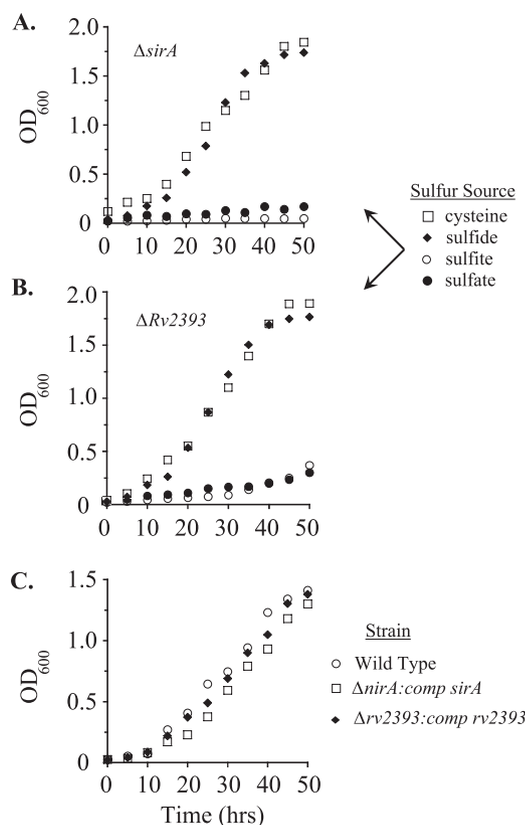


FIG. 4. Growth of *M. smegmatis* mutants on various sulfur nutrients. (A) Growth of the $\Delta sirA$ mutant on minimal media containing cysteine (\square), sulfide (\blacklozenge), sulfite (\circ), or sulfate (\bullet) as the sole sulfur source. Generation times for growth on cysteine and sulfide were 7.5 and 6.7 h, respectively; no growth was observed in media containing sulfite or sulfate. (B) Growth of the $\Delta Rv2393$ mutant on the same sulfur sources as those mentioned for panel A. The generation time for the mutant was 10 h when the medium contained cysteine or sulfide. Both cells fed sulfate and cells fed sulfite exhibited steady-state growth with a generation time of 19 h. (C) To demonstrate that the growth effects of the deletions were due solely to a lack of expression of the intended coding regions, the growth on sulfate of the deletion strains complemented with plasmids that express the wild-type gene was compared to that of the wild-type strain containing an empty vector. The doubling times of the three strains were extremely similar: for the wild type (\circ), 8.8 h; for the $\Delta sirA$ strain compared with the *sirA* strain (*nirA::comp sirA*) (\square), 8.6 h; and for the $\Delta Rv2393$ strain compared with the *Rv2393* strain ($\Delta rv2393::comp rv2393$) (\blacklozenge), 8.5 h. Growth protocols are described in Materials and Methods. Each data point represents the average of results from three experiments; single standard deviation units (not shown) are comparable to the diameters of the symbols representing the averaged values. OD₆₀₀, optical density at 600 nm.

way of *M. smegmatis*, an *Rv2393* deletion strain was constructed (see Materials and Methods). The deletion removes the entire gene with the exception of short, 140- and 53-nucleotide stretches of sequence at the 5' and 3' edges of the coding region, respectively (Fig. 3A). Growth of the deletion strain on sulfide or cysteine is indistinguishable from that of the wild type (Fig. 4B and C), suggesting that *Rv2393* acts upstream of the enzymes that incorporate sulfide into cysteine (*O*-acetylserine sulfhydrylase [*cysKI*] and serine transacetylase [*cysE*]). In contrast, the doubling time of the mutant on sulfite or sulfate was approximately twofold slower than that of the wild

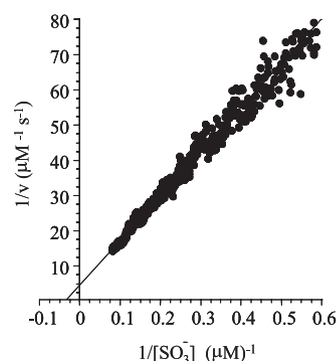


FIG. 5. Initial rate determination of the Michaelis constants for the SirA-catalyzed reduction of sulfite. Sulfite reduction was monitored continuously (under stringent, anaerobic conditions) via the change in absorbance at 684 nm associated with the oxidation of the electron donor, MV (see Materials and Methods). The assay conditions were as follows: sulfite (13 μM), reduced MV (250 μM), *O*-acetyl-L-serine (30 μM), SirA (1.3 μM), *O*-acetyl-L-serine sulfhydrylase (1.2 μM), and HEPES-K⁺ (50 mM, pH 8.0) (25 ± 3°C). The reaction rate is given in terms of sulfite reduced per unit time (μM/s⁻¹). The reduction of one equivalent of sulfite requires the oxidation of six equivalents of MV. v, initial velocity.

type (Fig. 4B and C). Thus, the point of action of *Rv2393* coincides with that of *SirA*: the reduction of sulfite.

The primary structure of *Rv2393* clusters with an orthologous group of proteins from ancient phylogenies (COG2381.1), many of which are type II chelataes—catalysts that insert divalent cations (Co²⁺, Fe²⁺, Mg²⁺, or Ni²⁺) into modified tetrapyrroles to produce redox-sensitive cofactors, including chlorophylls, vitamin B₁₂, heme, coenzyme F₄₃₀, and siroheme (6). Sulfite and nitrite reductases, which are often essential for the assimilation of sulfur and nitrogen, require siroheme to accomplish their redox chemistry. The linking of *Rv2393* to the chelatae family and to sulfite reductase function provides a rationale for the colocalization of *sirA* and *Rv2393* and the common metabolic point of action of the proteins that they encode, which is that *Rv2393* is the ferrochelatase that inserts Fe²⁺ into sirohydrochlorin to produce the siroheme cofactor that is necessary for *SirA* function.

To test the hypothesis that *Rv2393* is, in fact, a ferrochelatase, plasmid-borne *Rv2393* was used to complement an *E. coli* $\Delta cysG$ mutant (302Δa) (Table 1), in which ferrochelatase function is deleted. The genome of the mutant lacks *cysG*, which encodes siroheme synthase, a trifunctional protein that catalyzes the last three steps in siroheme biosynthesis: (i) methylation of uroporphyrinogen III (at C-2 and C-7) to produce dihydrosirohydrochlorin; (ii) oxidation of dihydrosirohydrochlorin to produce sirohydrochlorin; and (iii) insertion of Fe²⁺ into sirohydrochlorin to produce siroheme (31). The cysteine-biosynthetic pathway is disrupted in *cysG* deletion strains at the point of insertion of iron into sirohydrochlorin. The cysteine auxotrophy that results from the inability to produce siroheme provides a nutrient-based selection for methyltransferase and/or chelatae function. It should be noted that the dehydrogenase function needed by these strains is provided by endogenous activity (47); thus, selecting for ferrochelatase function exclusively requires restoration of the methyltransferase activity in 302Δa, which is accomplished by transforma-

tion with pER247, a P15 origin plasmid that expresses the *Pseudomonas denitrificans* CobA protein, a uroporphyrinogen III methyltransferase that does not exhibit chelatase function (3, 14). The 302Δa strain harboring pER247 and pET23a with or without the Rv2393 coding region were plated onto minimal medium plates containing either sulfate or cysteine as the sole source of sulfur (see “Ferrochelatase complementation” in Materials and Methods) and incubated at 37°C for 24 to 48 h. The methyltransferase-competent strain (302Δa/pER247/pET23a) showed no detectible growth on sulfate as a sole source of sulfur. However, when transformed with a pET23A plasmid containing Rv2393, either from *M. tuberculosis* or *M. smegmatis*, the growth on the sulfate-containing plates was comparable to that of the *cysG*-complemented strain. The observed Rv2393-dependent transformation of the strain to cysteine prototrophy strongly supports the idea that Rv2393 encodes a ferrochelatase that is capable of inserting Fe²⁺ into sirohychlorin to produce siroheme. Based on these findings, we have named Rv2393 *che1* to indicate both its chelatase function and that it does not appear to be the only chelatase in the *M. tuberculosis* genome.

Metabolic complementarity in sulfite and nitrite reduction.

Assimilatory nitrite and sulfite reductases, like those found in *M. smegmatis* and *M. tuberculosis*, share similar catalytic strategies, molecular architectures, and substrate specificities. These enzymes are named on the basis of their relative catalytic efficiencies (V_{\max}/K_m) for sulfite versus nitrite, and interestingly, despite a greater overall efficiency toward sulfite, sulfite reductases often turn over faster with nitrite [i.e., $V_{\max}(\text{NO}_2^-) > V_{\max}(\text{SO}_3^{2-})$] (12). Thus, the inability of the ΔsirA strain to grow on sulfite could reflect the inefficiency of the mycobacterial nitrite reductase toward sulfite, which has not been measured, and/or the level at which the enzyme is expressed in the organism when grown on ammonia (as was the case in the present study).

Given that sulfite reductase (SirA) is remarkably inefficient toward nitrite (see above), it appears from the *M. smegmatis* genome that nitrite reductase (*nirBD*) is the organism's sole means of obtaining ammonia from either nitrate or nitrite (10, 23). Ammonia is required for the biosynthesis of amino acids, and from there, nitrogen is drawn into the urea cycle, glutamate metabolism, pyrimidine biosynthesis, and ultimately every nitrogen-containing compound in the cell (23). To assess the ability of nitrite reductase to reduce sulfite *in vivo*, the growth of the ΔsirA mutant on sulfate was assessed in a medium selected to optimize the expression of nitrite reductase for the assimilation of nitrogen, M9 medium in which nitrate was substituted for ammonia at an equivalent concentration, 20 mM. Under this condition, all assimilated nitrogen must pass either through nitrite reductase or through other, as-yet-unidentified, reductases in the cell. The substitution of nitrate for ammonia did not result in detectible growth of the organism on sulfate over a 70-h period at 37°C; however, replacement of sulfate by cysteine produced normal growth with nitrite (Fig. 6A). Clearly, the nitrite reductase activity of the cell is sufficient to allow normal growth on metabolites that lie downstream of the point of action of SirA. Thus, nitrogen and sulfur metabolism are well isolated at the point of nitrite and sulfite reduction by what may ultimately prove to be pronounced

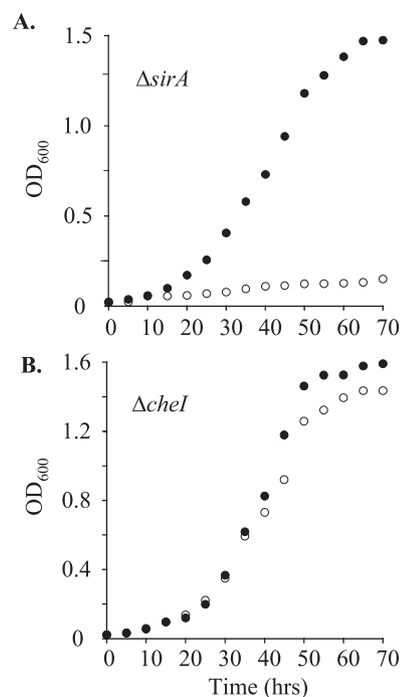


FIG. 6. Growth of *M. smegmatis* mutants on nitrate. (A) Growth of the ΔsirA mutant on minimal medium containing nitrate as the sole nitrogen source and either cysteine (○) or sulfate (●) as the sole sulfur source. The generation time for growth on cysteine was 7.1 h, as no growth was observed in medium containing sulfate. (B) Growth of the Δche1 mutant on the same nitrogen and sulfur sources as described for panel A. Generation times for growth on cysteine or sulfate were virtually identical, 7.3 h. Growth protocols are described in Materials and Methods. Each data point represents the average of results from three experiments; single standard deviation units (not shown) are comparable to the diameters of the symbols representing the averaged values. OD₆₀₀, optical density at 600 nm.

differences in the substrate specificities of the enzymes that catalyze these reactions.

Unlike the ΔsirA mutation, which causes a no-growth phenotype, the Δche1 mutation results in slow growth of the organism on sulfate or sulfite. Thus, the orthogonality seen in the sulfite reduction step clearly does not extend to the siroheme cofactor, which can be provided to SirA from alternative metabolic sources. Amelioration of slow growth by plasmid-born Che1 suggests that the endogenous siroheme pool(s) is in some way insufficient to achieve the levels of SirA activity needed for normal growth. Because nitrite reductase requires siroheme, it is plausible that the growth of the organism on nitrite, compared to that on ammonia, will up-regulate the cellular levels of siroheme; if so, the growth rate of the Δche1 mutant, when grown on sulfate or sulfite, will increase when the nitrogen source is switched from ammonia to nitrate. This is precisely what was observed (Fig. 6B). Clearly, the perturbation of nitrogen metabolism caused by the ammonia-to-nitrate substitution has impacted sulfur metabolism at the intersection of sulfite reduction and ferrochelatase function.

DISCUSSION

The structure of SirA was determined recently, and the enzyme was shown to reduce sulfite (at a single sulfite concen-

tration, 5.0 mM) under oxygen-depleting conditions (excess sodium dithionite) (38). In the current work, the initial rate of sulfite reduction was studied under strict anaerobic conditions as a function of sulfite concentration, which yielded kinetic constants for the enzyme [$k_{\text{cat}} = 0.17 (\pm 0.1)$ electron consumed s^{-1} (sulfite reduced) and $K_{m(\text{SO}_3^{2-})} = 27 (\pm 1) \mu\text{M}$] that are in line with previously published values for sulfite reductases from other organisms (12, 24). The value of k_{cat}/K_m , $3.4 \times 10^4 \text{M}^{-1} \text{s}^{-1}$, indicates that SirA catalyzes SO_3^{2-} reduction with modest efficiency, and the inability to detect the SirA-catalyzed reduction of nitrite at concentrations as high as 3.0 mM reveals that the efficiency of the enzyme toward NO_2^{-} is extremely low and suggests that this reaction is not likely to be physiologically relevant.

High-density transposon insertion mutagenesis in combination with microarray mapping of the *M. tuberculosis* genome has indicated that *sirA* is important for optimal growth of the organism on minimal medium containing sulfate as a sole source of sulfur (36). *sirA* was deleted in the present work and shown to be essential for the growth of *M. smegmatis* on sulfate or sulfite, regardless of whether nitrate or ammonia was used as a nitrogen source. These facts are of value not only for their contribution to the genetic and biochemical fundamentals of sulfur reduction in *M. smegmatis* and related organisms but also for what they reveal about the orthogonality of the sulfur and nitrogen-reducing pathways in these species. On this note, it is interesting that it may be possible to assess, and therefore more accurately annotate, the sulfite-versus-nitrite substrate preferences of ferredoxin-dependent reductases based on primary sequence (38).

The common metabolic point of action of Rv2393 and SirA, the siroheme requirement for SirA function, and the ability of Rv2393 to complement an *E. coli* ferrochelatase mutant argue strongly that Rv2393, a protein with previously unidentified function, is a ferrochelatase, which we have named *che1*. The fact that the slow-growth phenotype of the Δche1 mutant is rescued by plasmid-encoded Che1 or by growth on nitrate supports the idea that the organism's growth rate is limited by siroheme access. While nitrogen and sulfur metabolism are well isolated by what appear to be nonoverlapping substrate specificities of the nitrite and sulfite reductases, these pathways do overlap at the level of siroheme production. It is interesting to note that a second CbiX-like open reading frame (Rv0259c) is found quite near the genes that encode nitrite reductase in *M. smegmatis*. Perhaps the siroheme pool(s) in the mutant grown on rate-limiting sulfur metabolites is simply too small to satisfy the metabolic demands placed on it; alternatively, access of SirA to the pool(s) could be limited by intrinsic specificities that bias delivery of the metalated porphyrin to particular recipients. It is exciting to consider the possibility that the Δche1 strain, whose growth rate is limited by sulfite reduction, presents us with the opportunity to establish a linkage between sulfite reduction and metabolic sources of siroheme.

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