

Proteomic Analysis of Survival of *Rhodococcus jostii* RHA1 during Carbon Starvation

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Rhodococcus jostii RHA1, a catabolically diverse soil actinomycete, is highly resistant to long-term nutrient starvation. After 2 years of carbon starvation, 10% of the bacterial culture remained viable. To study the molecular basis of such resistance, we monitored the abundance of about 1,600 cytosolic proteins during a 2-week period of carbon source (benzoate) starvation. Hierarchical cluster analysis elucidated 17 major protein clusters and showed that most changes occurred during transition to stationary phase. We identified 196 proteins. A decrease in benzoate catabolic enzymes correlated with benzoate depletion, as did induction of catabolism of alternative substrates, both endogenous (lipids, carbohydrates, and proteins) and exogenous. Thus, we detected a transient 5-fold abundance increase for phthalate, phthalate ester, biphenyl, and ethyl benzene catabolic enzymes, which coincided with at least 4-fold increases in phthalate and biphenyl catabolic activities. Stationary-phase cells demonstrated an ~250-fold increase in carbon monoxide dehydrogenase (CODH) concurrent with a 130-fold increase in CODH activity, suggesting a switch to CO or CO₂ utilization. We observed two phases of stress response: an initial response occurred during the transition to stationary phase, and a second response occurred after the cells had attained stationary phase. Although SigG synthesis was induced during starvation, a $\Delta sigG$ deletion mutant showed only minor changes in cell survival. Stationary-phase cells underwent reductive cell division. The extreme capacity of RHA1 to survive starvation does not appear to involve novel mechanisms; rather, it seems to be due to the coordinated combination of earlier-described mechanisms.

n most natural environments, and particularly in soil, microbes spend most of their existence starving (46). Organic compounds are the primary nutrients for heterotrophic organisms, providing both the reductant and carbon source for catabolism. Therefore, adaptation to deprivation of organic substrates must have been one of the key selective pressures during evolution of soil bacteria. Limitation of organic substrates is frequently referred to as C-limitation, but reductant for respiration, not C, is probably the limiting factor in these situations. To date, diverse strategies to exploit scarce nutrients and survive starvation have been discovered in different organisms. They include reductive cell division, scavenging endogenous substrates, dormancy, and spore formation, as well as programmed cell death and increased mutation rate (52). Some of these major modifications to cell physiology are mediated by different regulatory responses, including (i) a nonspecific, general stress response (regulated by SigB in many bacteria) providing cells with multiple stress resistance mechanisms; (ii) the ppGppdependent stringent response including decrease of growth-related functions (transcription, translation, nucleotide biosynthesis, cell envelope metabolism, and cell division); (iii) specific regulation of catabolic processes (central metabolism, alternative carbon source utilization, and use of endogenous storage compounds) and modified expression of transporters (5, 63).

Physiological responses to nutrient starvation have been studied in a number of organisms; however, most studies have focused on *Escherichia coli* (52). Far less is known of the molecular basis of starvation survival in organisms that better tolerate starvation, such as actinomycetes, although this group has attracted some recent attention. For example, models using oxygen, organic substrate, or nitrogen deprivation and stationary-phase survival demonstrated that mycobacteria are able to survive the extended periods in a nongrowing state by activating a SigBand SigH-regulated general stress response as well as a ppGppregulated stringent response (7, 22, 28, 39, 56, 61, 67). One central hypothesis about starvation resistance is focused on the role of lipid metabolism, which generally maintains mycobacterial viability in the absence of growth and particularly allows persistence in lung lesions (10).

Rhodococci are mycolate-containing soil actinomycetes, related to mycobacteria, that are abundant in soil environments. Rhodococci have great catabolic diversity, which permits them to play important roles in nutrient cycling and to have numerous important commercial applications (3, 69). Other rhodococci have evolved for pathogenicity in humans, animals, and plants (18, 45). Despite the critical role of starvation survival in these important processes mediated by rhodococci, very few studies are reported (17, 64), and the molecular mechanisms of nutrient starvation survival in rhodococci are not defined. To better understand the physiology and molecular basis of starvation survival in rhodococci, we have selected Rhodococcus jostii RHA1. The diverse catabolic capabilities of RHA1 were studied extensively (20, 24, 30, 49, 55, 59, 60, 66), and its genome was sequenced and annotated (44). Previous studies of RHA1 have investigated global responses to heat shock, high salinity (15), and desiccation (35).

The physiological responses to a stress condition primarily rely

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TABLE 1 Strains and plasmids used in this study

Strain or plasmid	Relevant genotype/comments	Source
Plasmids		
pKD46	λ -RED (gam bet exo) araC rep101(Ts) Amp ^r	10
pIJ773	λ -RED (gam bet exo) araC rep101(Ts) Amp ^r	23
pUZ8002	<i>tra neo</i> RP4, Km ^r	57
pUC-Hy	Source of <i>hyg</i> gene	38
RF00131J24	Fosmid clone carrying <i>sigG</i> , Cm ^r	71
RFMD5	RF00131J24 with cat replaced with hyg, Hmr	This study
RFMD6	RFMD5 with <i>sigG</i> replaced with Apra cassette, Hm ^r Apra ^r	This study
Strains		
E. coli BW25113	K-12 derivative: Δ <i>araBAD</i> , Δ <i>rhaBAD</i> Amp ^r	23
E. coli DH10B	Host for pUZ8002 and RFMD4, RFMD6, RFMD8, RFMD10, Km ^r	23
RHA1	Rhodococcus jostii RHA1, wild type	65
RHA1_013	Derivative of RHA1; $\Delta sigG$; Apr ^r	This study

on changes in protein synthesis. Therefore, to characterize the molecular basis of starvation tolerance in RHA1, we employed a global quantitative proteomic approach, utilizing high-resolution, large-format two-dimensional gel electrophoresis (2DGE) in combination with mass spectrometry (MS)-based protein identification. This approach allowed monitoring of temporal abundance profiles of a large number of cytosolic proteins and thus identification of the RHA1 carbon starvation proteome and its temporal behavior. These studies were followed by targeted gene disruption and resting-cell biotransformation and enzyme activity assays. The results provide new insights into the strategies rhodococci employ to survive carbon starvation.

MATERIALS AND METHODS

Chemicals. Pharmalyte 3-10 and Immobiline Dry-Strips were purchased from GE Healthcare (Baie d'Urfé, Canada). Iodoacetamide and 3-[(3-cholamidopropyl)-dimethylammonio]-1-propanesulfonate (CHAPS) were from Acros Organics (Fair Lawn, NJ) and MP Biomedicals (Aurora, OH), respectively. Mini Complete protease inhibitor cocktail was purchased from Roche (Laval, Canada). Sypro Ruby was from Bio-Rad Laboratories Ltd. (Mississauga, Canada). 3-[*N,N*-Dimethyl(3-myristoylaminopropyl)-ammonio]propanesulfonate (ABS-14) and dithiothreitol (DTT) were from Fluka (Switzerland) and Sigma, respectively. Nucleotides were purchased from Qiagen (Mississauga, Canada). All chemicals were of analytical grade and used without further purification.

Strains and media. Rhodococcus jostii strain RHA1 was isolated prior to 1995 from soil contaminated with lindane in Japan (41) and kindly provided by M. Fukuda. RHA1 and the mutant RHA1_013 (Table 1) were grown at 30°C on W minimal salt medium containing KH₂PO₄ (1.7 g/liter), Na₂HPO₄ (9.8 g/liter), (NH₄)₂SO₄ (1.0 g/liter), MgSO₄ · 7H₂O (0.1 g/liter), FeSO4 · 7H₂O (0.95 mg/liter), MgO (10.75 mg/liter), CaCO₃ (2.0 mg/liter), ZnSO \cdot 7H₂O (1.44 mg/liter), CuSO₄ \cdot 5H₂O (0.25 mg/liter), CoSO₄ · 7H₂O (0.28 mg/liter), H₃BO₃ (0.06 mg/liter), and concentrated HCl (51.3 µl/liter) (32) supplemented with 10 mM benzoate and 20 mM ammonium sulfate. These amounts of exogenous carbon and nitrogen were adjusted to ensure that it is carbon limitation that causes RHA1 to enter stationary phase. For proteomic studies, RHA1 cells were first precultured in 100-ml Erlenmeyer flasks to mid-log phase (optical density at 600 nm $[OD_{600}]$, 1.0), then used to inoculate (1%) fresh medium in 500-ml flasks, and further cultured to a growth phase of interest and harvested. We selected seven time points representing different growth phases (Fig. 1). Growth data were collected using cultures inoculated with normalized quantities of cells based on OD and were based on at least 3

independent experiments, each consisting of 3 biological replicates. For gene replacement, *Escherichia coli* and *Rhodococcus* strains were grown using media and culture conditions described previously (23). *E. coli* BW25113 was used to propagate pKD46 and fosmid RF00131J24. *E. coli* DH10B/pUZ8002 was the fosmid donor strain for intergeneric conjugation. Strains were grown on LB broth supplemented with ampicillin (100 μ g/ml), apramycin (Apra, 50 g/ml), chloramphenicol (12.5 μ g/ml), kanamycin (50 μ g/ml), and hygromycin (50 and 150 μ g/ml for RHA1 and *E. coli*, respectively) as required.

Preparation of cell extracts. Cell extracts were prepared essentially as described previously (59) with some modifications. Briefly, cell pellets were washed once with 0.14 M NaCl and once with Tris-EDTA (TE) buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8.0) and then stored as aliquots at -80° C. The pelleted cells were suspended in lysis buffer (2% ABS-14, 4% CHAPS, 30 mM DTT, 30 mM Tris [pH 7.5], Mini Complete protease inhibitor cocktail [1:100, vol/vol]) and disrupted using a Fast Prep Bio 101 Thermo Savant bead beater (5 cycles of 20 s). Protein concentration was determined using the 2D Quant kit (GE Healthcare). For enzyme assays, cell-free protein extracts were prepared in the same manner, except that the cell pellets were washed using 20 ml of 50 mM Tris-HCl, pH 7.5, and suspended in 1.0 ml of the same buffer, and the cell extracts were used immediately. The protein concentrations of the cell extracts were quantified using the bicinchoninic acid (BCA) method.

Proteomic analyses. The cytosolic proteomes were resolved using 2DGE essentially as described previously (59) with minor modifications. Briefly, aliquots of cell extracts containing 90 µg protein were separated in the first dimension by isoelectric focusing (IEF) for a total of 88 kVh at 20°C using 24-cm, nonlinear immobilized pH gradient (IPG) strips and a pH gradient of 3 to 7. The rehydration solution contained 9 M urea, 3 M thiourea, 35 mM DTT, 4% CHAPS, and Pharmalyte pH 3-10. Proteins were separated in the second dimension using 12% SDS-polyacrylamide gels and the ETTAN DALTtwelve System (GE Healthcare). Gels were stained using Sypro Ruby and digitally imaged using a Typhoon 9400 (GE Healthcare). Spot detection, matching, abundance quantification, and normalization were performed using Progenesis Workstation software (Nonlinear Dynamics, Durham, NC). The relative abundance of each protein was determined from the signal intensity and normalized against total signal intensity detected on a gel. Only spots with a minimum normalized volume of 0.002 or greater were analyzed. Protein patterns of each growth condition were based on gels from three independent cultures. Protein spots of interest were identified using a Voyager DESTR matrixassisted laser desorption ionization-time of flight (MALDI-TOF) mass spectrometer (Applied Biosystems, Foster City, CA) based on peptide mass fingerprint (PMF) analysis combined with the MASCOT search engine (Matrix Science) and the RHA1 protein database at the Proteomics Centre, University of Victoria. Identified proteins fulfilled four criteria: the MASCOT search score was above 55, a minimum of 4 peptides were matched, the protein sequence coverage was at least 20%, and the predicted molecular weight (MW) and pI values were consistent with the experimentally observed ones. We clustered the detected proteins based on their temporal abundance profiles. For this, the normalized signal intensities of the proteins were subjected to hierarchical cluster analysis by Gene Spring using Pearson's correlation coefficient.

CODH assay. The carbon monoxide dehydrogenase (CODH) enzyme activity assay was performed using a Varian Cary IE UV-visible spectrophotometer equipped with a cuvette holder with a thermostat. The CODH activity of RHA1 cells grown to M70 or S120 was measured by monitoring the CO-dependent reduction of 2-(4-indophenyl)-3-(4-ni-trophenyl)-2H-tetrazolium chloride (INT) ($\varepsilon = 17.981 \text{ mM}^{-1} \text{ min}^{-1}$) as described in reference 34 with some modifications. Briefly, samples of 20 ml were harvested from three 100-ml cultures by centrifugations at 4°C at 10,000 × *g*. The cell extracts were prepared as described above. We added 20 µl of each cell extract to 100 µl of CO-saturated phosphate buffer per assay, and A_{562} was followed over time for 30 min. One unit of enzyme



FIG 1 (A) Growth of *R. jostii* RHA1 and entry into benzoate-limited stationary phase: changes in OD₆₀₀ (\blacklozenge); in benzoate (mM) (\bigcirc) and muconate (mM) (\blacklozenge) concentration; in cell viability (CFU/ml) (\diamond); and in total cell protein concentration (μ g/ml) (\triangle). (B) Description of the time points representing different growth phases when the samples were taken for proteomic analyses.

activity was defined as the amount of enzyme required to reduce 1 nmol of INT per minute at 30°C.

Resting-cell assay. RHA1 cells were collected by centrifugation at different growth phases and washed with, and resuspended in, W medium to an OD_{600} of 1.0. The resultant cell suspensions (1 ml) were supplied with 0.2 mM biphenyl or 0.2 mM phthalate and incubated for 120 min at 30°C with shaking in sealed 4.5-ml glass vials. To stop the reaction, 0.1 ml of 6 N HCl was added to the mix. As an internal standard, 30 ppm of 2,6-dichlorobiphenyl or 40 ppm terephthalic acid dimethyl ester was added. To extract biphenyl or phthalate, we added hexane or ethyl acetate, respectively, and the mixtures were mixed vigorously using a vortex mixer for 1 min. The supernatant was recovered and dehydrated with sodium sulfate. One microliter of extract was analyzed by gas chromatography-MS. The analysis was done in triplicate. Controls were performed using heat-killed cells.

HPLC. The high-pressure liquid chromatography (HPLC) system consisted of a Waters 2695 separations module and 2996 photodiode array detector (Waters, MA). Chromatographic data were collected and

compiled using Empower software (Waters). Cell culture supernatants were filtered through 0.2-µm syringe filters. Chromatographic separation of benzoate was performed using an Aqua C18 125-Å (250 by 4.6 mm, 5-µm particle size) column (Phenomenex, CA) at a flow rate of 1.0 ml/ min at room temperature. The mobile phase consisted of two solutions: A, 0.5% phosphoric acid; and B, 100% methanol. The gradient profile was 0 to 5 min 25% (vol/vol) B; 5 to 6 min linear gradient to 60% (vol/vol) B; 6 to 11 min 60% (vol/vol) B; 11 to 12 min linear gradient to 25% (vol/vol) B; and 12 to 18 min 25% (vol/vol) B. The wavelengths of UV detection were set at 230 nm, 273 nm, and 260 nm to detect benzoic acid, catechol, and *cis,cis*-muconic acid, respectively.

Gene replacement. The *sigG* gene was deleted and replaced by an apramycin cassette using the λ -RED-based methodology as described previously (59). Fosmid RF00131J24 from the RHA1 fosmid library, which contained 36.5 kb of RHA1 genomic DNA, including *sigG*, was used. The cassette used to replace the *sigG* gene was constructed by PCR amplification and contained the Apra^r gene flanked by flippase recognition target (FRT) sequences. The fosmid with the replaced gene was trans-

INDEL 21 OK PIII	iers used in this study	
Primer	Sequence (5'-3')	Comments
SigGapraF	CCG ACA TCT GTC GGA CGG CCG CAG TAA GCT CCA ACC ATG att ccg ggg atc cgt cga cc	Apra ^r cassette amplification for <i>sigG</i> replacement, forward
SigGapraR	CAT GTA CGG ACG CGT TTT CCG CGC CAT CGG TAG GAG TCA tgt agg ctg gag ctg ctt c	Apra ^r cassette amplification for <i>sigG</i> replacement, reverse
SigGtestF	CGGAATGAAGACTCTGCTC	External sigG primer, forward
SigGtestR	GTGTGGGACACGGTAGAAC	External <i>sigG</i> primer, reverse
SigGintF	GTGTGGGACACGGTAGAAC	Internal <i>sigG</i> primer, forward
SigGintR	ATCAGGGTGCCGATCGAAG	Internal <i>sigG</i> primer, reverse

 TABLE 2 PCR primers used in this study^a

^{*a*} The 5' end of SigGapraF contained 39 nucleotides (uppercase letters) corresponding to the sense strand upstream of *sigG* ending in the start codon. The 5' end of SigGapraR contained 39 nucleotides (uppercase letters) corresponding to the antisense strand ending in the stop codon. The 3' ends of the PCR primers (lowercase letters) matched the 20-nucleotide and 19-nucleotide extensions, respectively, of the *aprA*-containing disruption cassette sequence.

ferred to RHA1 by intergeneric conjugation as described in reference 31. Apra^r Hm^s exconjugants were selected as double-crossover mutants. The mutant genotypes were verified using PCR with three sets of primers: SigGtestF and SigGtestR, SigGtestF and SigGapraR, and SigGintF and SigGintR (Table 2).

Phylogenetic analysis. Sequences of various CODHs were aligned by using ClustalX (version 2.0.9) with all parameters set at their default values. Phylogenetic analyses were performed using the neighbor-joining algorithm of PHYLIP (version 3.66). Neighbor joining was performed by using a random order of sequences on input. For bootstrap analyses, the Seqboot program of the PHYLIP package was used to generate 100 data sets, which were jumbled three times and analyzed as described above. Trees generated were visualized and printed by using TreeView (version 1.6.6).

RESULTS

Benzoate-limited growth of RHA1 and reductive division. As benzoate was depleted from the medium, RHA1 ended logarithmic growth, passed through a short transition phase (TP), and entered stationary phase (ST) (Fig. 1). The point of entry into ST was defined as the point when the biomass stopped increasing, which corresponded to the beginning of a plateau in both total protein and optical density. Although both protein and OD measurements stopped increasing in ST, CFU continued to increase until late stationary phase (S150). During TP, a metabolite of benzoate, muconate, was detected, and its concentration decreased

gradually throughout stationary phase. The low level of muconate in the medium (less than 1 mM) was unlikely to support the increase in CFU during stationary phase, and this increase was probably primarily due to reductive division. Accordingly, scanning electron microscopy showed a decrease in cell length from 7.5 μ m (±2 μ m) for exponentially growing cells to 3.5 μ m (±1.5 μ m) during ST (Fig. 2). Further, the total protein calculated per CFU increased during mid-log phase (ML) and then sharply decreased in TP and stationary phases (not shown).

To estimate a survival capacity of RHA1, cells continued to be incubated for a total of about 2 years. After plateauing, protein, OD, and CFU remained essentially constant for 83 days. After 125 days, both OD and CFU decreased 2-fold, and after 500 days we detected about 30% of the initial maximum number of cells remaining viable. Finally, after 700 days of incubation 10% of the cells were still viable (data not shown).

Changes in the RHA1 proteome during starvation. To understand the RHA1 starvation response at the molecular level, we classified more than 1,600 proteins based on their abundance profile. This constitutes 18% of the total number of predicted proteins encoded by the RHA1 genome. Considering that the starvation response develops over time and depends on the severity of starvation, we monitored protein abundance profiles over seven time points representing different growth phases: ML, transition after



S120



1µm

FIG 2 Scanning electron micrographs of R. jostii RHA1 cells collected during mid-log (ML) and stationary (S120) phases of growth.



FIG 3 Sections of 2D gels showing parts of the cytosolic proteomes of *R. jostii* RHA1 grown on benzoate and collected at different time points: ML, T78, T83, S88, S120, S150, and S447. CO dehydrogenase subunits CoxM2, CoxF3, CoxM3, and CoxL3 are shown with arrows.

78 and 83 h of incubation (T78 and T83, respectively), and stationary phase after 88, 120, 150, and 447 h of incubation (S88, S120, S150, and S447, respectively) (Fig. 1). The total number of detected proteins remained approximately constant throughout all the time points, while the relative abundance of a large proportion of proteins changed (Fig. 3). Compared to the ML proteome, the total proportion of proteins with increased abundance (including the unique spots) was 21% in the T78 proteome and increased to 41% in the S120, S150, and S447 samples. The total proportion of proteins with decreased abundance (including the absent spots) was 16% in P78 cells and increased to 40% in the T83 proteome and further to 47 and 53% in the S150 and S447 proteomes, respectively. Assuming that proteins displaying similar patterns of abundance often participate in related processes, we clustered the detected proteins based on their temporal expression patterns. This analysis clustered 1,090 proteins, which constitute about 66% of the total detected proteins, and elucidated 17 clusters, of which the most distinct and represented 10 are shown in Fig. 4. The highest number (30%) of the abundance changes (including increase and decrease) among the clustered proteins occurred during the relatively brief posttransition phase (T78 to T83) of growth, indicating that the switch between exponentially growing and nongrowing states requires more significant adjustments in cell physiology.

Based on MALDI-TOF, we identified 196 proteins differentially expressed in at least one of the seven proteomes, each representing one time point of growth. These constitute 16% of the proteins detected in an average proteome and represent each of the elucidated clusters. The averaged normalized signal intensity of each identified protein at each time point, as well as protein MS identification parameters, is presented in Table S1 in the supplemental material. Of these proteins, 55 are shown in Table 3.

Catabolism of aromatics. Fourteen enzymes responsible for the catabolism of benzoate (Ben), catechol (Cat), and protocatechuate (Pca) were identified (Table 3). The abundance of the upper benzoate pathway enzymes (BenABCD) started declining in T83 (cluster 1), whereas the abundance of the lower benzoate pathway enzymes (CatAB and PcaIGJH) decreased in S120 (cluster 5). This decline correlated with the depletion of benzoate and muconate (Fig. 1). The abundance of PcaBLF and the regulatory protein PcaR remained essentially constant until a reduction of at least 2-fold in the S447 cells (cluster 9). This supports differences observed earlier in PcaBLFR and PcaIGJH abundance profiles (59) and further suggests their independent regulation.

Once benzoate and its intermediates were depleted, starving RHA1 cells increased capacity to use alternative exogenous substrates. We observed the activation of several aromatic catabolic pathways. The abundance of enzymes responsible for the catabolism of phthalate (PadAd1) and phthalate esters (PatE) increased up to 5-fold in T83 (cluster 3). These abundances correspond to 5 to 10% of those previously measured in phthalate-grown ML phase cells (59). Later, during S120 and S150, six Bph and Etb enzymes responsible for utilization of biphenyl, ethylbenzene, benzene, and styrene appeared (cluster 6). The abundances of the large catalytic subunits BphAa and EtbAa1 reached 28 to 35% of those in biphenyl-grown ML phase cells (60). Further, we identified 6 other catabolic enzymes whose abundance increased in starving RHA1 cells (ro08833, ro02718, ro08019, ro02204, ro00441, and ro06057) (see Table S1 in the supplemental material). Two of them (propane monooxygenase and propanediol dehydrogenase) were up to 40-fold more abundant during S150 than ML (cluster 2) (Table 3).

To test whether the abundance increase correlates with the catalytic activities of the Bph and Pad enzymes, we measured phthalate and biphenyl degradation in resting cells of benzoatelimited RHA1. As summarized in Table 4, phthalate degradation activity was greatest during T83, when PadAd1 expression reached its maximum. Biphenyl degradation activity was greatest in S150, in agreement with BphAa and EtbAa1 abundance profiles. Al-though increased, the maximum biphenyl degradation rate of benzoate-limited cells was 150-fold lower than that of ML cells grown on biphenyl (data not shown).

CO oxidation. In addition to aromatic catabolism, we observed the activation of CO oxidation systems during S120 and S447 phases. The RHA1 genome encodes three CODHs, whose structural and accessory genes are organized in 3 separate chromosomal clusters (Fig. 5). Each CODH enzyme is comprised of

Cluster	Growth phase		Proteins	Proteins					
Number	ML	T78	T83	S88	S120	S150	S447	In Cluste	r Examples (Table 3)
1	· ·							133	BenABCD
2	-							69	PrmA, SigG, GroL1
3	8 0.1							126	PadAd1, PatE
4								26	CitA4, SdhA2
5	- : WW						X II I	67	CatAB, PcaIGJH
6							W W	21	BphAaB2D, CoxM2F3M3L3
7								41	SucDC, FabF
8	•••							64	USP, DnaK1
9	••	1						65	PcaFBLR, GroL2, GroEL
10						4		71	ClpP

FIG 4 Hierarchical cluster analysis of protein spot signal intensities for the RHA1 proteomes. Protein signal intensities from averaged triplicate samples were obtained by using Progenesis and exported into GeneSpring software for cluster analysis. Proteins with similar signal intensity patterns were grouped together into 17 clusters, of which 10 are shown. The thick lines represent the average profiles.

three subunits: small (CoxS), medium (CoxM), and large (CoxL). We observed the expression of four of these proteins (Fig. 5). Expression of these proteins increased up to 245-fold in S150 compared to the expression of those in ML cells (Fig. 3, cluster 6). We also measured the CODH activity in ML and S120 cells, which increased 130 times during starvation (not shown).

The three CODH-encoding gene clusters in the RHA1 genome differ in the organization of structural as well as accessory genes (Fig. 5). Furthermore, the comparative sequence analysis revealed that the large subunits, CoxL, of the three CODHs share only 33 to 43% sequence identity and cluster with distinct sets of homologs. Phylogenetic analysis using the best-studied CODHs confirmed that the RHA1 CODHs belong to three distinct subclades (Fig. 6). CoxL3 (Rjo_ro05232) is most similar to CODHs from actinomycetes *Mycobacterium tuberculosis* and *Arthrobacter* sp. FB24, whereas CoxL1 (Rjo_ro03489) and CoxL2 (Rjo_ro04976) cluster with CODHs from alphaproteobacteria and betaproteobacteria, respectively. The fact that CoxL3 closely clusters with CODHs from *Mycobacterium* species was also demonstrated in a recent study (68).

Central metabolism. Four central metabolic pathways, i.e., glycolysis, tricarboxylic acid (TCA), glyoxylate shunt, and pentose phosphate pathways, responded to starvation (Table 3). Two glycolysis enzymes (6-phosphofructokinase and pyruvate kinase) decreased abundance in S120 (cluster 5). Dihydrolipoyl dehydrogenase (pyruvate dehydrogenase complex) was reduced in T78 and

not detected during starvation (cluster 1). The abundance of seven TCA cycle enzymes (two homologs of citrate synthase, isocitrate dehydrogenase, α and β subunits of succinate-CoA ligase, succinate dehydrogenase, and malate dehydrogenase) decreased in S88 and S150 (clusters 4 and 7). In contrast, the abundance of the glyoxylate shunt enzyme (isocitrate lyase) and fumarate hydratase as well as two enzymes of the pentose phosphate pathway (glucose-6-phosphate 1-dehydrogenase and phosphogluconate dehydrogenase) increased in T83 (cluster 2).

Protein, lipid, and carbohydrate metabolism. Consistent with reductive cell division, the abundance of the cell division initiation protein (Ro01080) responsible for cell division and chromosome partitioning was increased up to 56-fold during starvation (cluster 6) (Table 3); however, ribosomal proteins (RpsAIB, AlaS, and Tsf) were less abundant upon entering T83 phase (see Table S1 in the supplemental material). Further, ClpP protease (Ro01371, cluster 10) and peptidase (Ro01171, cluster 8) (Table 3) were more abundant. Biosynthesis of individual amino acids also showed significant changes (see Table S1). This includes IlvC, involved in biosynthesis of branched amino acids, which increased in abundance 500-fold during stationary phase.

Further, several enzymes required for incorporation of inorganic nitrogen into cell material changed abundance. Glutamate dehydrogenase and glutamine synthetase GlnA1 decreased in abundance (see Table S1 in the supplemental material); however,

					Normali	ized signal i	ntensity ^b				
Protein name	name	Ro no.	Genetic context	Cluster	ML	T78	T83	S88	S120	S150	S447
Benzoate catabolism											
Benzoate 1,2-dioxygenase alpha subunit	benA	02384	ben	1	1.70	1.30	0.61	0.43	0.15	0.11	0.07
Benzoate 1,2-dioxygenase beta subunit	benB	02385	ben	1	1.09	0.97	0.59	0.46	0.18	0.17	0.18
Benzoate 1,2-dioxygenase reductase subunit	benC	02386	ben	1	0.98	0.81	0.35	0.21	0.02	Z	Z
cis-1,6-Dihydroxycyclohexa-3,5-diene-1-carboxylate dehydrogenase	benD	02387	ben	1	0.56	0.50	0.33	0.20	0.02	0.04	0.01
Catechol 1,2-dioxygenase	catA	02373	cat	5	3.50	3.28	2.40	1.82	0.67	0.36	0.19
Muconate cycloisomerase	catB	02372	cat	J.	0.42	0.35	0.23	0.11	0.02	0.01	0.01
3-Oxoacid CoA-transferase, subunit A	pcal	01334	pca	J.	0.35	0.32	0.30	0.27	0.10	0.04	0.02
Protocatechuate dioxygenase alpha subunit	pcaG	01336	pca	5	0.26	0.28	0.25	0.18	0.04	0.05	0.05
3-Oxoacid CoA-transferase	pcaJ	01333	pca	5	0.10	0.12	0.10	0.02	0.02	0.02	0.02
Protocatechuate dioxygenase beta subunit	pcaH	01335	pca	IJ.	0.14	0.20	0.08	0.08	0.02	0.01	0.01
Acetyl-CoA C-acyltransferase	pcaF	01340	pca	6	0.28	0.39	0.42	0.39	0.30	0.20	0.02
3-Carboxy-cis,cis-muconate cycloisomerase	pcaB	01337	pca	6	0.10	0.11	0.07	0.08	0.08	0.07	z
3-Oxoadipate enol-lactone hydrolase/4-carboxymuconolactone	pcaL	01338	pca	6	0.21	0.20	0.10	0.06	0.08	0.06	0.03
decarboxytase Transcriptional ramilator TelD familu	hcaD	01330	5.5 4	o	0.03	0.04	0.05	0.06	0.05	0.03	0.00
	Vinud	CCC10	pru		0.0	F0.0	00.0	00.0	C0.0	000	70.0
Other catabolic enzymes											
Biphenyl 2,3-dioxygenase alpha subunit	bphAa	08060	p p h	9	Z	Z	Z	Z	0.48	0.54	0.15
cis-3-Phenylcyclohexa-3,5-diene-1,2-diol dehydrogenase	bphB2	10126	bph/etb	9	Z	Z	Z	z	0.03	0.03	0.13
2,6-Dioxo-6-phenylhexa-3-enoate hydrolase	bphD1	10136	bph/etb	9	Z	Z	z	z	0.10	0.08	0.03
Ethylbenzene dioxygenase alpha subunit	etbAal	10133	bph/etb	6	Z	Z	Z	z	0.47	0.45	0.26
2-Hydroxy-6-oxohepta-2,4-dienoate hydrolase	etbD1	08044	bph/etb	6	0.01	0.01	0.02	0.01	0.24	0.29	0.18
2,6-Dioxo-6-phenylhexa-3-enoate hydrolase	etbD2	10146	bph/etb	6	0.01	0.01	0.02	0.02	0.27	0.26	0.10
Phthalate 3,4-dioxygenase, ferredoxin reductase subunit	padAd1	08162/12013	pad	3	0.02	0.03	0.07	0.09	0.03	0.02	z
Phthalate ester hydrolase	patE	08169	pat	3	0.04	0.04	0.09	0.12	0.07	0.06	0.03
Propane monooxygenase hydroxylase large subunit	prmA	00441	Cluster with phenol hydroxylase and GroEL	2	0.02	0.03	0.03	0.04	0.13	0.68	0.38
Probable 1,3-propanediol dehydrogenase		06057	ATP-dependent protease	2	0.05	0.03	0.15	0.24	0.24	0.56	0.29
Other catabolic enzymes											
CODH, medium subunit	coxM2	04975	CODH	9	0.09	0.11	0.08	0.12	0.22	0.20	0.19
CODH, maturation factor ^c	coxF3	05229	CODH	9	Z	Z	Z	Z	0.17	0.38	0.49
CODH, medium subunit ^c	coxM3	05230	СОДН	9	0.04	0.07	0.03	0.04	0.18	0.20	0.43
CODH, large subunit c	coxL3	05232	CODH	9	Z	Z	0.05	0.04	0.17	0.20	0.46
Central metabolism											
6-Phosphofructokinase	nfkA	06479	A midotran sfera se <i>oat</i>	LC.	0.02	0.04	0.03	0.03	0.01	z	z
Pyruvate kinase	pvkl	01007	Glutamate synthase	ы го 1	0.29	0.29	0.29	0.27	0.30	0.18	0.19
, Dihydrolipoyl dehydrogenase ^e	dldH1	02140	Hypothetical	1	0.09	0.04	Z	z	Z	Z	z
Citrate synthase		04998	Peptidylprolyl isomerase	7	0.03	0.05	0.05	0.05	0.03	0.02	z
Citrate synthase	citA4	04993, 08812	Phosphoserine transaminase	4	0.09	0.09	0.03	0.01	0.02	0.01	0.004
Isocitrate dehydrogenase ^e		06238	Arabinose efflux	4	0.16	0.10	0.02	z	Z	z	z
Succinate dehydrogenase	sdhA2	06246	sdh	4	0.60	0.45	0.33	0.31	0.32	0.19	0.27
Succinate-CoA ligase, alpha subunit	sucD	05573	suc	7	1.00	0.85	0.85	0.80	0.67	0.58	0.42
Succinate-CoA ligase, beta subunit	sucC	05574	suc	7	1.07	0.82	0.79	1.03	0.92	0.58	0.50
Malate dehydrogenase	fabF	06000	Single gene	7	0.20	0.24	0.25	0.16	0.06	0.02	0.06
Isocitrate lyase		02122	Lipid metabolism	2	0.20	0.22	0.74	0.75	0.59	0.38	0.46
Fumarate hydratase ^{e}	į	08824	Succinate dehydrogenase	2	z	z	0.01	0.01	0.01	0.02	0.02
Glucose-6-phosphate 1-dehydrogenase	zwj4	07184	Pentose phosphate pathway	5 5	0.03	0.02	0.03	0.04	0.05	0.05	0.11
Phosphogluconate dehydrogenase	gnd3	07246	RNA helicase	2	0.08	0.06	0.15	0.23	0.39	0.43	0.50

Glucose-6-phosphate 1-dehydrogenase Phosphogluconate dehydrogenase

Stress response											
Chaperonin GroL2 ^{c,e}	groL2	02146	Universal stress protein	6	0.83	0.69	0.60	0.60	0.65	0.63	Z
Chaperone protein GroEL ^{G,e}	groEL	06190	10-kDa chaperonin	6	0.81	0.67	1.20	0.93	0.71	0.63	0.34
Chaperonin GroL1 ^{G,e}	groLl	00448	Propane monooxygenase	2	Z	Z	0.61	0.37	0.11	1.45	Z
Heat shock protein Hsp70 DnaKl ^{c,e}	dnaKl	05497	Chaperones: grpE1, dnaJ2	8	1.01	1.10	0.71	0.72	1.07	1.99	1.36
Universal stress protein ^{c,e}		02263	Universal stress protein, oxidoreductase	6	0.03	0.03	0.06	0.06	0.05	0.03	0.03
Universal stress protein ^{c,e}		04893	Pseudouridylate synthase	8	0.08	0.06	0.06	0.12	0.12	0.19	0.18
Catalase ^{d,e}	katE	04309	Fe(3+) uptake regulator	7	0.02	0.05	0.05	0.07	0.04	0.01	0.02
Superoxide dismutase ^e	sodA	04009	Amidotransferase	2	1.08	06.0	1.10	1.19	1.40	1.38	1.62
Heme superoxide dismutase c		06860	Heme biosynthesis, oxidoreductase	10	0.19	0.19	0.18	0.26	0.25	0.31	0.39
ATP-dependent Clp protease ^{d,e}	clpP	01371	<i>clpX</i> , FKBP-type bacterial trigger factor	10	0.25	0.25	0.40	0.41	0.30	0.43	0.50
Prolyl amin opeptidase c,e		01171	panB, coenzyme metabolism	8	0.01	0.03	0.03	0.02	0.01	0.09	Z
Cell division initiation protein		01080	Cell division	6	0.002	0.005	0.01	0.07	0.11	0.08	0.03
Transcription											
RNA polymerase sigma factor G	sigG	05361	ArsR	2	0.07	0.07	0.11	0.13	0.17	0.14	0.18
" Protein identification was accepted if the MASCOT-generated probabili- the threshold, the protein ID was accepted only if the theoretical and expe	ty-based Mov erimental pI a	wse score in the l und MW agreed.	the theorem of the theorem of the theorem of the the the the theorem of the theo	0.05) and at l	east 3 peptic	les were ide	entified. W	/hen the l	AASCOT :	score was l	below

Proteins reported to be affected by starvation in references 4, 7, 14, 21, 25, 50, and 55.

Spot signal intensities were normalized and averaged over three biological replicates.

^c Proteins reported to be regulated by MglA (21). ^d Proteins reported to be regulated by SigB (4).

TABLE 4 Degradation activities in RHA1 cells during different growth phases^a

	Degradation	activity, 10 ⁻³ μι	nol/min/mg (SD)	
Substrate	ML	T83	S88	S150
Phthalate	0.23 (0.10)	0.58 (0.05)	0.26 (0.004)	0.36 (0.007)
Biphenyl	0.33 (0.12)	ND	0.15 (0.10)	1.37 (0.07)

^{*a*} The assays were performed in duplicate. The degradation activities were measured by monitoring the concentration of the substrates. ND, not detected.

the alternative pathway of L-glutamate production that includes prolyl aminopeptidase, 1-pyrroline-5-carboxylate dehydrogenase, and 5-oxoprolinase showed up to 300-fold-higher abundance.

A number of proteins responsible for metabolism of carbohydrates increased in abundance in starving cells (see Table S1 in the supplemental material). These included glycosyltransferases and sugar epimerases, as well as glucan and glycoside hydrolases, which increased in abundance up to 13-fold in stationary-phase cells.

We detected 21 proteins involved in lipid metabolism, whose abundance was changed during starvation (see Table S1 in the supplemental material). More abundant proteins included acyl carriers and carboxylases, likely involved in lipid degradation. Among proteins with decreased abundance, we identified acyl and mycolyl carriers, fatty acid synthases, and desaturases, as well as enzymes involved in *de novo* synthesis and modification of fatty and mycolic acids.

Sigma factor G. We identified 9 transcriptional regulators whose abundance changed during starvation (see Table S1 in the supplemental material). Sigma factor SigG (cluster 2) doubled in abundance during stationary phase (Table 3). To elucidate a pos-



FIG 5 Genetic organization of CO oxidation genes in *R. jostii* RHA1. The numbers above the arrows indicate CDS (Ro) numbers from the RHA1 genome assembly (http://www.rhodococcus.ca). These gene clusters are located on the chromosome. Patterns represent different types of genes. Structural CODH genes: diagonally striped, CODH large subunit; vertically striped, CODH medium subunit; horizontally striped, CODH small subunit (ferredoxin). Accessory CODH genes: diamonds, CODE maturation factor; hatched diamonds, CoXE homologs; squares, CoXG homologs. Other genes: stippled, transcriptional regulator; white, miscellaneous; and waved, hypothetical. Gray background and bold border indicate genes whose products were identified in this study, as listed in Table 3.



FIG 6 Phylogenetic analysis of CODH. The unrooted neighbor-joining tree was calculated from an alignment of the amino acid sequences of the large subunits of three predicted CODHs from *R. jostii* RHA1 (Ro03489, Ro04976, and Ro05232) and the following proteins from the PIRSF (strain followed by accession number with a subscript indicating the strain): Oligotropha carboxidovorans P19919_{Oca}; Hydrogenophaga pseudoflava P19913_{Hpc}; Sulfolobus solfataricus Q97V74_{Soc}; Pyrobaculum aerophilum Q8ZW74_{Pac}; Sulfolobus acidocaldarius Q4J7T6_{Sac}; Thermoproteus tenax Q703X0_{Tte}; Haloquadratum walsbyi Q18IU5_{Hwa}; Arthrobacter sp. FB24 A0JWJ9_{Asp}; Burkholderia xenovorans LB400 Q13IY1_{Bxe}; Ralstonia eutropha H16 Q0K601_{Reu}; Geobacillus kaustophilus Q53708_{Mti}; Clostridium tetani Q891A4_{Cte}; Rhodopseudomonas palustris Q6N0U4_{Rpa}; Desulfotalea psychrophila Q6AK66_{Dps}; Sinorhizobium meliloti Q92LP4_{Sme}; Silicibacter pomeroyi Q5LT95_{Spo}.

sible physiological role of SigG in RHA1 starvation response, we investigated the growth phenotype of a gene disruption mutant lacking *sigG* (Table 1). No drastic changes in the phenotype (OD, CFU, total cell protein) were observed in the $\Delta sigG$ strain RHA1_013 during growth on benzoate. The strain showed 40% loss in viability after sudden starvation and an ~5-h delay in, and 20% lower, recovery after benzoate was provided back to the medium (data not shown).

Stress response during starvation. We identified 23 proteins whose functions may be associated with stress response and whose abundance changed under starvation (Table 3; see Table S1 in the supplemental material). All three GroEL chaperones encoded by the RHA1 genome and chaperone DnaK were identified and belong to the 5 most abundant proteins in the detected proteomes. Interestingly, their abundance profiles differed but covered all the tested points of growth. GroL2 (ro02146) was highly abundant at all the time points except S447, where it was not detected (cluster 9). Chaperonin GroL1 (ro00448) was first detected in T83 cells, which coincided with the abundance 723-fold in S150 cells, which coincided with the abundance increase of DnaK1 (ro05497).

Two universal stress proteins (USPs) were detected. Similarly

to the GroEL proteins, the USPs had different abundance profiles, falling into 2 phases: ro02263 was 2-fold more abundant in T83 to S120 cells (cluster 3), and ro04893 was 2-fold more abundant in S120 to S447 cells (cluster 8). Three oxidative stress response proteins, superoxide dismutase (ro04009), heme superoxide dismutase (ro06860), and catalase KatE (ro04309), were more abundant during starvation. Notably, we again see 2 phases in the abundance profiles: KatE was more abundant during T78 to S120 phases of growth; in contrast, the dismutases were more abundant in S150 to S447.

DISCUSSION

This study demonstrated the remarkable ability of a soil actinomycete, *Rhodococcus jostii* RHA1, to survive carbon source starvation for at least 2 years, retaining 10% of cells viable. The detailed proteomics study identified two groups of processes responding to starvation. The first group included a transition from growing to nongrowing cells, a reductive division, a shutdown of benzoate catabolism, and a decrease in transcription, translation, *de novo* biosynthesis, glycolysis, and TCA cycle. The second group represented induction of the two-phase stress response and a switch to utilization of alternative endogenous and exogenous carbon sources including the activation of multiple aromatic catabolic pathways and CO utilization. These mechanisms, in combination with other detected rearrangements in cell physiology, likely contribute to the extreme capacity of RHA1 to survive long-term starvation.

The ability to survive long-term starvation groups RHA1 with mycobacterial strains known to survive for extended periods of time (51, 67, 72). We observed that RHA1 stationary-phase cells underwent reductive cell division, which generally results from completion of DNA replication and cell division in the absence of growth. It is also known as dwarfing. This strategy has been proposed to be beneficial during early starvation (33, 52). To understand the RHA1 starvation response at the molecular level, we characterized the temporal abundance profiles of seven cytosolic proteomes collected at different points of growth and starvation and showed that the highest number of protein changes occurred during the switch between exponentially growing and nongrowing states. We identified alternative catabolic pathways that transiently increased abundance levels at different time points during starvation, which may indicate that RHA1 cells attempt to exploit different alternative substrates at different times. Such a strategy may allow starving RHA1 a quicker recovery from a dormant state once substrates become available. Examples of this survival strategy have been seen in other bacteria. For example, adaptation of Lactobacillus lactis to carbon starvation involves induction of pathways for metabolism of alternative exogenous carbon sources, such as sugars, citrate, and glycerol (63). The choice of alternative carbon sources may be indicative of catabolic specialization of a starving organism.

A particularly intriguing result is the increased abundance of CODHs, which reversibly oxidize CO to CO_2 and permit use of CO as a reductant for energy metabolism (62). Organisms oxidizing CO usually can also use CO as the sole carbon source (62). The induction of CODHs during starvation suggests that RHA1 is switching from heterotrophic to autotrophic metabolism. Since CODH interconverts CO and CO_2 , it enables assimilation of CO₂ or CO into cellular building blocks (62). Interestingly, it has recently been shown that glycerol-limited *M. tuberculosis* is able to

fix CO_2 into biomass through carboxylation reactions (6). The phylogenetic differences between the three CODHs encoded by the RHA1 genome coincide with the detected differences in their abundance profiles. For example, the CODH related to CODH in actinomycetes (CoxF3M3L3) was not detected in ML to T83 but showed a significant abundance spike in S447, whereas CoxM2 was detected during all phases and doubled in abundance in stationary cells. Thus, there may be a meaningful link between the phylogeny and physiology of the individual RHA1 CODH isozymes.

Among central metabolic pathways, the enzymes of glycolysis, pyruvate oxidation, and the TCA cycle reduced in abundance during starvation; however, the glyoxylate shunt and pentose phosphate pathway enzymes became more abundant during transition phase. The glyoxylate shunt is required to assimilate fatty acids via acetyl-coenzyme A (acetyl-CoA), whereas the pentose phosphate pathway provides precursors for nucleic acids. Interestingly, in *M. tuberculosis*, glyoxylate shunt enzymes were shown to be essential for fatty acid catabolism, virulence, and survival in the host (47). Most recently, isocitrate lyase has been shown to be essential for *M. tuberculosis* growth under glycerol-limited conditions and proposed to be a key enzyme of a novel metabolic route for carbohydrate metabolism (6).

The decreased production of protein synthesis machinery and the increased abundance of ClpP protease and peptidase suggest an arrest of *de novo* protein synthesis and a switch to protein turnover. This is consistent with the stringent response, which occurs in response to starvation in a wide range of bacteria, including *E. coli* (42, 73). In addition to protein scavenging, ClpP proteases degrade short-lived regulatory proteins as well as misfolded and damaged proteins, which occur more often under stress (19), thus maintaining cellular homeostasis (75). Biosynthesis of branched amino acids was significantly induced during stationary phase as exemplified by IlvC. Intriguingly, its 25% identical homolog in yeast was suggested to be a bifunctional protein, whose second role is to be involved in DNA packaging required for maintenance of stable mitochondrial DNA (76), which may be of higher demand under starvation stress.

Synthesis of glutamate and glutamine is required for incorporating inorganic nitrogen into cell material. Although carbonstarved RHA1 cells were provided with nitrogen, the enzymes responsible for nitrogen uptake decreased in abundance. This suggests that carbon starvation may trigger the nitrogen starvation response. Such coregulation of the responses to carbon and nitrogen starvation has been reported in *E. coli* (43, 48). Simultaneously, RHA1 increased the level of the alternative pathway of Lglutamate production using proline cleaved from peptides. This indicates that starving RHA1 reduces inorganic nitrogen uptake, but in order to provide amino groups to amino acid synthesis, it continues to maintain a cellular glutamate pool using cellular peptides.

Starvation increased the production of a series of enzymes necessary for metabolizing glycogen, a storage compound in RHA1 (26). In addition to glycogen, lipids (triacylglycerols, wax esters, and polyhydroxyalkanoates) are important storage compounds in actinomycetes (1). They can be deposited as insoluble inclusions in the cytoplasm, accounting for up to 70% of the cellular dry matter (1). Therefore, in the absence of an available carbon source, the products of lipid degradation, fatty acids, become an important source of carbon and energy. We detected a massive rearrangement in the biosynthesis of enzymes involved in lipid metabolism as well as metabolism of fatty and mycolic acids, major components of the cell envelope of the mycolic acid bacteria. It is noteworthy that lipid metabolism has been shown to play a central role in maintaining long-term mycobacterial viability in the absence of growth (10). Thus, starving RHA1 scavenges cellular glycogen and lipids as endogenous sources of carbon.

Overall, the detected reduction in biosynthesis and the switch to utilizing the endogenous sources of carbon and energy have been reported in other organisms as the first response to carbon and energy deprivation (52, 63). This allows cells to save and redirect the intracellular carbon and energy resources to essential and strategically important processes. Reductive cell division accompanied by dwarfing involves self-digestion, resulting in degradation of endogenous material including storage and membrane lipids, carbohydrates, and proteins.

We aimed to elucidate a possible physiological role for SigG in RHA1 starvation response. The protein belongs to the most phylogenetically divergent group 4 of σ 70 family of sigma factors with extracytoplasmic function (ECF), which regulates genes in response to environmental signals (58). SigG shares 64% amino acid sequence identity with SigG from *Mycobacterium tuberculosis* strain H37Rv, whose expression was also highly increased in stationary phase (40) and during macrophage infection (9). We think that the subtle changes in the phenotype of the *sigG* deletion mutant do not necessarily negate a role for SigG in the starvation response, and the role of SigG in RHA1 starvation remains unresolved. One possibility is that, similarly to the *M. tuberculosis* SigG, it may play a role in the SOS response (36).

We detected a significant stress response in starving RHA1. Three GroEL and DnaK chaperones were more abundant at different points during starvation, indicating their differential regulation. A high abundance of GroL2 in all except one proteome is consistent with the housekeeping role suggested for GroL2 homologs (Cpn60.2) in actinobacteria (37). GroEL and DnaK protein chaperones are commonly induced under stress and assist in proper assembly of misfolded or unfolded polypeptides generated under stress conditions (8, 13). The highest level of induction of the homologous chaperones in M. tuberculosis was detected in cells exposed to heat shock, but induction was also seen under hyperosmolarity, oxidative stress, and starvation conditions (29). Similarly, two USPs had different abundance profiles, falling into two phases. Intriguingly, the RHA1 genome encodes 21 predicted USPs (reference 44 and this study). This number stands out drastically from one USP predicted in the Xanthomonas campestris genome, 8 in the M. tuberculosis genome, and 12 in the Streptomyces coelicolor genome (56). The detected USPs, ro02263 and ro04893, share only 36% sequence identity. Although induction of USPs was repeatedly associated with stress response, their biochemical function remains largely unknown. In E. coli, UspA was shown to be required for survival of prolonged growth arrest, and its mutation caused global changes in protein synthesis (53, 54).

Starvation also induced a two-phase oxidative stress response. Oxidative damage was suggested to be the Achilles' heel of stationary-phase bacterial cells (52) and therefore requires a protective response. As a part of this response, respiratory activity is reduced during early starvation. This prevents uncontrolled drainage of endogenous carbon reserves and protects cells against the damaging effects of reactive oxygen species (ROS) produced by the electron transport chain (52). Higher resistance to oxidative stress was also demonstrated for Mycobacterium smegmatis (67). We also detected the increased abundance of several other stress-related proteins. They include methyltransferase of the type I restrictionmodification system (ro11151), known to protect bacterial cells against invasion of foreign DNA (74), and two subunits of succinate-semialdehyde dehydrogenase from the glutamate catabolic pathway, a homolog of which has been proposed to buffer redox changes in the cytoplasm of Saccharomyces cerevisiae (11). They further include two predicted DPS (DNA protection during starvation) family miniferritins (ro04028 and ro04043) known to protect bacterial DNA from oxidants during starvation (27) and diaminobutyrate-pyruvate aminotransferase, responsible for biosynthesis of ectoine, a primary compatible solute in RHA1, whose production had been shown to be induced during heat shock, osmotic, and desiccation stresses (15, 35). NADPH:quinone reductase was also more abundant and, similarly to the NADPH quinone reductase in Helicobacter pylori, may play an important role in coping with oxidative stress by managing the cell quinone pool (70).

Most of the detected stress-related proteins followed 2 phases of expression: the first phase was transiently induced during transition phase and included catalase KatE, GroEL chaperones, and USP ro02263. The second phase was induced later during stationary phase and included superoxide dismutases, DnaK chaperone, thiocyanate hydrolase, USP ro04893, ClpP, and a type I restriction-modification system. This two-phase stress response is consistent with the early stationary expression of general stress proteins (2) and catalase KatE (16) and the late-starvation-induced ROS-specific proteins MrgA (Dps homolog), catalase KatA, and alkyl hydroxyperoxide hydrolase AhpCF (12) observed in *Bacillus subtilis*.

Overall, this study reports the complex rearrangements in cytosolic protein biosynthesis in response to starvation in *Rhodococcus jostii* strain RHA1, a catabolically diverse soil actinomycete. We conclude that the exceptional ability of RHA1 to survive starvation does not involve novel mechanisms but rather is due to the coordinated combination of many previously described mechanisms.

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