Oleoyl Coenzyme A Regulates Interaction of Transcriptional Regulator RaaS (Rv1219c) with DNA in Mycobacteria*

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Background: RaaS mediates mycobacterial survival in nonpermissive growth conditions by controlling expression of ATP-dependent efflux pumps.

Results: Oleoyl-CoA regulates binding of the RaaS transcription factor to DNA and thus expression of the RaaS regulon and RaaS-mediated persistence.

Conclusion: The activity of bacterial efflux is regulated by metabolites that are produced during active growth.

Significance: Disregulation of efflux pumps results in killing of persisting mycobacteria with low metabolic activity.

Tuberculosis remains a major infectious disease, with 8.6 million new cases estimated in 2012 claiming 1.3 million lives (1). Moreover, the causative agent of tuberculosis, Mycobacterium tuberculosis, is believed to latently infect one-third of the world’s population (2). Treatment of tuberculosis requires the administration of a combination of four drugs (isoniazid, rifampin, ethambutol, and pyrazinamide) for at least 6 months. The remarkable ability of M. tuberculosis to survive in the infected host for years is well documented and involves the activation of complex regulatory pathways, as well as the production of specialized enzymes and transcriptional regulators (3, 4).

We have recently demonstrated that treatment of mycobacteria with antimicrobial agents targeting cell wall biosynthesis (cereuloen, isoniazid, or ethambutol) improves bacterial survival in non-permissive growth conditions (5). This phenomenon is mediated by the transcriptional regulator RaaS, which controls expression of putative ATP-dependent efflux pumps (Bcg_1278c/Beg_1277 in Mycobacterium bovis) bacllus Calmette-Guérin (BCG) and Rv1218c/Rv1217c in M. tuberculosis). Deletion of raaS from M. tuberculosis or M. bovis BCG genomes has no effect on mycobacterial growth in logarithmic phase. However, the deletion mutants in both mycobacteria are impaired in long-term survival at stationary phase. Moreover, the M. bovis BCG deletion mutant displays a survival defect during macrophage infection in murine lungs and spleen, whereas the M. tuberculosis deletion mutant does not persist during macrophage infection (5). These results demonstrate that RaaS controls mycobacterial survival in various models of M. tuberculosis disease. In the present study, we conducted bioinformatic analyses and predicted that fatty acid derivatives of CoA are putative ligands of RaaS. We confirmed this experimentally by demonstrating specific binding of oleoyl-CoA to RaaS and the role of oleic acid, a precursor of oleoyl-CoA, in RaaS-mediated mycobacterial survival. We propose that fatty acid metabolites produced during active growth are also involved in controlling expression of genes encoding mycobacterial efflux pumps.

* The abbreviations used are: BCG, bacillus Calmette-Guérin; ITC, isothermal titration calorimetry.
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Oligonucleotides used in this study

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*a qRT-PCR, quantitative RT-PCR.

EXPERIMENTAL PROCEDURES

Organisms and Media—The M. bovis BCG Glaxo strain was grown in Sauton’s liquid medium supplemented with albumin-dextrose complex as described previously (5). Ethambutol and oleic acid were added, using a 1-ml syringe fitted with a 25-gauge needle, 30 days after inoculation at final concentrations of 98 and 200 μM, respectively. An equivalent volume of sterile water was added to control cultures. Bacterial viability was assayed by counting colony-forming units on 7H10 agar.

Transcriptional Profiling—Total RNA was isolated from 30 ml of early logarithmic (7 days) or stationary phase (31 days) M. bovis BCG cultures after a 24-h exposure to 200 μM oleic acid using the guanidinium thiocyanate/TRIzol method (6). DNA contamination was removed with Turbo DNA-free DNase (Ambion) before cDNA was generated using SuperScript reverse transcriptase II (Invitrogen) as described previously (7). Quantitative PCR was performed in a Corbett Rotor-Gene 6000 real-time thermocycler (Qiagen) by application of ABsolute qPCR SYBR Green mixture (Thermo Scientific) and normalized to 16 S rRNA using the guanidinium thiocyanate/TRIzol method (6).

Purification of Recombinant RaaS—The M. tuberculosis raaS gene was cloned into the Ndel and NheI sites of the pET15-Tev plasmid to generate a hexahistidine-tagged recombinant protein (4). Protein expression in Escherichia coli BL21(DE3) was induced by isopropyl β-D-thiogalactopyranoside at a final concentration of 0.2 mM. Recombinant RaaS was purified using a HiTrap 1-ml IMAC HP column (Amersham Biosciences). Site-directed mutants of RaaS were generated using a GeneArt mutagenesis kit (Invitrogen) according to the manufacturer’s instructions.

Fluorescence Anisotropy—The synthetic oligonucleotides, containing the imperfect direct repeats (Pr14F and Pr14R in Table 1), were covalently labeled with ATTO 647N succinimidyl ester dye (Invitrogen). Steady-state fluorescence anisotropy binding titrations were carried out on a Tecan Safire2 micro-plate reader using a 635-nm light-emitting diode for excitation and a monochromator set at 680 nm (bandwidth of 20 nm) for emission in buffer containing 50 mM Tris-HCl (pH 8.5) and 150 mM NaCl.

Isothermal Titration Calorimetry (ITC)—RaaS protein and oleoyl-CoA (Sigma) were diluted in 50 mM Tris-HCl (pH 8.5) and 150 mM NaCl. RaaS (15 μM in a 1.4-ml cell) was then titrated at 25 °C by 5-μl injections of the ligand (250 μM in the syringe) using a VP-ITC calorimeter (MicroCal). Raw data were normalized and corrected for heats of dilution of the ligand. Binding stoichiometries, enthalpy values, and equilibrium dissociation constants were determined by fitting the corrected data to a bimolecular interaction model using Origin 7 software (OriginLab).

Small-angle X-ray Scattering Experiments and Data Analysis—Synchrotron x-ray scattering data were collected on the SWING beamline of the SOLEIL Synchrotron (Gif-sur-Yvette, France) using a PCCD-170170 detector at a wavelength of 1.03 Å. The scattering patterns were measured by merging 10–20 data recordings with 1-s exposure time each for several solvent concentrations at 1 and 3 mg/ml. To check for radiation damage, all successive exposures were compared, and no changes were detected. Using a sample-detector distance of 1.8 m, a range of momentum transfer of 0.0065 < q < 0.6 Å−1 was covered (s = 4 min/θ/λ, where 2θ is the scattering angle, and λ = 1.5 Å is the x-ray wavelength). The data were processed using standard procedures and extrapolated to infinite dilution using the program PRIMUS (8). The forward scattering, I(0), and the radius of gyration, Rg, were evaluated using Guinier approximation, assuming that at very small angles (s < 1.3/Rg), the intensity is represented as I(s) = I(0)exp(−s2Rg2/3).

EMSA—Annealed Pr14F and Pr14R (Table 1), containing the RaaS-binding site or the raaS upstream region (174 bp), were used for EMSAs as described previously (5). Briefly, the reaction buffer (10 μl) contained 50 mM Tris HCl (pH 8.0), 1 mM EDTA, 50 mM NaCl, 4% (v/v) glycerol, and 1 g of sheared salmon sperm DNA per reaction. 32P-Labeled annealed oligonucleotides and purified recombinant RaaS were added to the reaction at concentrations of 60 and 215 nM, respectively, except where stated otherwise. Oleoyl-CoA, palmitoyl-CoA, and dodecyl-CoA were added to samples as indicated. The samples were incubated for 10 min at room temperature before running on a polyacrylamide gel as described above. Following electrophoresis, gels were fixed in 20% (v/v) methanol and 10%...
RaaS is annotated as a member of the TetR family (1.4 E-value) and is conserved in the actinobacteria, with sequence identities ranging from 99% (M. tuberculosis) to 74% (M. avium) based on the sequence identity to the regulator domain of YsiA (14%) (Fig. 1). More distant homologs are also found in other actinobacteria, with sequence identities between 56% (M. avium) and 48% (Streptomyces sulphuris). Extensive fold recognition analysis was performed using the @TOME-2 server (10), and a structural alignment with proteins sharing low sequence identity (<20%) with RaaS was generated. These distantly related proteins share a strong hydrophobic ligand-binding pocket within the C-terminal domain. One of these proteins, YsiA (FadR), is a transcriptional regulator of the fatty acid degradation pathway in Bacillus subtilis, and its structure has been solved in complex with stearoyl-CoA (Protein Data Bank ID 3WHB) (11). Comparative modeling of RaaS suggested that it has features compatible with the binding of the CoA moiety of the acyl-CoA ligand, despite a low sequence identity to the regulator domain of YsiA (14%) (Fig. 1).

In particular, the C-terminal domains (involved in ligand binding) of both proteins contain conservation clusters around a short motif DXX (Protein Data Bank ID 3WHB). We examined the diphosphate group of acyl-CoA and its structure has been solved in complex with stearoyl-CoA. Arg-144 is conserved in the RaaS homologs and in FadR. In addition, for the RaaS models generated, Arg-144 is compatible with acyl-CoA binding. Further analysis of the dimeric structure of YsiA highlights the possible salt bridge with the conserved arginine. Arg-140 is then well positioned to directly interact with the effector, but instead, it is involved in N-terminal cap of an α-helix and in the formation of a salt bridge with the conserved arginine. Arg-140 is then well positioned to directly interact with the acyl-CoA ligand (ID 3WHB). We examined residues in the neighborhood position of the DXX motif. The Gln-154 lateral chain in FadR points toward the ligand in the YsiA structure. This residue is Arg-144 in RaaS. The substitution of glutamine to arginine suggests a possible salt bridge with the diphosphate group of acyl-CoA. Arg-144 is conserved in the close RaaS homologs and in FadR. In addition, for the RaaS models generated, Arg-144 is compatible with acyl-CoA binding. Further analysis of the dimeric structure of YsiA highlighted the possible role of Tyr-174 (corresponding to Tyr-174 in RaaS). This residue in the monomer is in contact with the ligand molecule bound to the other monomer. The aromatic ring of the tyrosine is stacked with the adenine ring of the acyl-CoA ligand. Thus, according to our dimer model, Tyr-174 in RaaS was postulated to play the same role as Tyr-174 in YsiA to stabilize the bound acyl-CoA. On the basis of these observa-
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Oleoyl-CoA Regulates RaaS Binding to DNA—Mycobacterial growth media and eukaryotic cells (a natural niche for mycobacterial replication and persistence) are rich sources of oleate (12). Mycobacteria convert oleate to oleoyl-CoA by fatty acyl-CoA ligases (13), and oleoyl-CoA is an abundant fatty acid precursor for the synthesis of mycobacterial cell wall components. Therefore, we tested the effect of oleoyl-CoA on the RaaS-DNA interaction using EMSAs. As shown in Fig. 2A, oleoyl-CoA was able to completely abolish the RaaS-mediated band shift at concentrations ≥15 μM. Using ITC, we demonstrated direct binding of oleoyl-CoA to RaaS and a dissociation constant ($K_d$) for oleoyl-CoA binding to RaaS of 3.65 ± 0.28 μM (Fig. 2B).

We next investigated whether other acyl-coenzyme derivatives have any effect of the RaaS-DNA complex. Lauroyl-CoA (C12:0), palmitoyl-CoA (C16:0), and palmitoleoyl-CoA (C16:1) had no effect on the band shift in the EMSA (Fig. 3, A–C). Stearoyl-CoA (C18:0) showed a dose-dependent inhibition; however, it did not eliminate RaaS-DNA binding even at the highest concentration tested, 100 μM (Fig. 3D). These results suggest that RaaS preferably binds C18 fatty acid derivatives (C18:1 oleoyl-CoA or, to some extent, C18:0 stearoyl-CoA). Longer acyl-CoA metabolites (>C18) could not be tested in our assay due to their poor solubility.

Role of Conserved Amino Acids in DNA and Ligand Binding—Our thorough sequence structure analysis (see above) suggested that three amino acids could be responsible for acyl-CoA binding in RaaS: Arg-140, Arg-144, and Tyr-174. Using site-directed mutagenesis, we generated three RaaS mutants in which these residues were replaced with alanine: R140A, R144A, and Y174A.

FIGURE 2. Oleoyl-CoA regulates RaaS binding to DNA. A, purified RaaS (215 nM) was mixed with radiolabeled annealed Pr14F/Pr14R oligonucleotides (60 nM); oleoyl-CoA was added as indicated. B, characterization of the interaction between RaaS and oleoyl-CoA by ITC. Samples were prepared as described under "Experimental Procedures." Oleoyl-CoA bound to RaaS with a $K_d$ of 3.65 ± 0.28 μM in an enthalpy-driven reaction ($\Delta H = -6.42 \pm 0.14$ kcal/mol). The band shift identifying RaaS binding to DNA is marked with an arrow.

FIGURE 3. Effect of acyl-CoA derivatives on RaaS-DNA binding. Lauroyl-CoA (A), palmitoyl-CoA (B), palmitoleoyl-CoA (C), stearoyl-CoA (D), and oleoyl-CoA (E) were added at 0, 5, 10, 25, 50, and 100 μM, respectively.

We first confirmed the structural integrities of the RaaS proteins by performing small-angle x-ray scattering experiments.
Guinier plot analysis yielded radii of gyration of 30.4 ± 1, 29.7 ± 1.8, and 30.7 ± 1.3 Å for the wild-type and mutant R144A and Y174A proteins, respectively. The R140A mutant was unstable at high concentrations, and therefore, it was not possible to obtain reliable data for this protein. The apparent molecular masses calculated from the I(0) values were 40, 44, and 40 kDa for wild-type RaaS, R144A, and Y174A, respectively, confirming that all proteins were present as folded homodimers in solution.

We next investigated their binding to DNA. All three mutations caused substantial increases in the dissociation constant measured by fluorescence anisotropy, ranging from 129 ± 70 nM for R144A to 611 ± 120 and 654 ± 300 nM for Y174A and R140A, respectively, compared with 31 ± 6 nM determined for wild-type RaaS (Fig. 4). The reduced DNA binding affinity resulting from these mutations was consistent with the results of EMSAs. As shown in Fig. 5A, the mutant proteins did not cause substantial DNA band shifts under the conditions used. The observed changes in the RaaS-DNA binding affinity in the wild-type and mutant proteins could be explained by altered RaaS conformation and/or flexibility, as observed in other TetR-like regulators (14). We therefore could not use EMSA to investigate the influence of oleoyl-CoA on the interaction of mutant RaaS with its DNA-binding site. The mutants also required higher oleoyl-CoA concentrations for titration in ITC experiments compared with wild-type RaaS. In fact, the RaaS-oleoyl-CoA binding affinities for the mutant proteins could not be measured by ITC because oleoyl-CoA aggregates at the high concentrations required (>300 µM), reducing data quality. We therefore employed native gel electrophoresis to investigate oleoyl-CoA binding to RaaS mutants, measuring the effect of oleoyl-CoA binding on migration of RaaS proteins. As shown in Fig. 5B, the addition of 30 or 60 µM oleoyl-CoA resulted in more...
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rapid migration of RaaS and an apparent reduction in size (indicated with an arrow), which could be explained either by dissociation of the dimer to a monomeric form or by an altered protein charge or flexibility. Treatment of the mutant proteins with 30 μM oleoyl-CoA had no pronounced effect on their migration patterns, whereas the addition of 60 μM oleoyl-CoA had a partial effect, resulting in formation of two forms of RaaS (Fig. 5B). These results suggest that the conserved amino acids in the ligand-binding domain are indeed involved in binding oleoyl-CoA.

Oleic Acid Influences RaaS-mediated Biological Effects—Finally, to confirm the biological relevance of our findings with oleoyl-CoA, we investigated the effect of oleic acid on expression of the RaaS regulon and the drug-mediated improvement of M. bovis BCG survival. We have recently shown that RaaS is an antirepressor protein, and its regulon includes several genes encoding efflux pumps and conserved proteins of unknown function (5). In mycobacterial cells, oleic acid is converted to oleoyl-CoA by the coordinated action of fatty acyl-AMP ligases and acyl-CoA-synthesizing fatty acyl-CoA ligases (12), resulting in an increase in intracellular acyl-CoA concentration. An increase in oleoyl-CoA concentration should prevent the transcriptional repressor RaaS from binding DNA, leading to elevated expression of the RaaS efflux regulon in nonreplicating mycobacteria, which is detrimental to long-term survival. Expression levels of selected genes belonging to the RaaS regulon were measured using quantitative RT-PCR. We compared the effect of oleic acid treatment on expression of raaS, bcg_1277c, bcg_1278c, and drrC in logarithmic and stationary growth phases. Oleic acid had no statistically significant effect on expression of these genes in logarithmic phase (Fig. 6A). However, expression of RaaS-regulated genes (bcg_1277c, bcg_1278c, and drrC) was induced by 4.8-, 2.7-, and 3.7-fold, respectively, in stationary phase M. bovis BCG cultures 24 h after treatment with oleic acid (Fig. 6A). Expression of raaS (in the same operon as bcg_1277c and bcg_1278c) in stationary phase mycobacteria was also increased after treatment with oleic acid. Exposure of log phase mycobacteria to oleic acid did not result in growth alteration (Fig. 5B). Strikingly, the addition of oleic acid to 1-month-old stationary phase cultures completely abolished the antimicrobial survival-enhanced effect (Fig. 6C) mediated by RaaS, thus implicating oleoyl-CoA in the regulation of long-term mycobacterial survival.

DISCUSSION

Expression of bacterial operons is tightly controlled by transcription factors, repressors or activators, which alter transcript levels in response to physiological bacterial state or environmental cues (15). In some cases, as pertinent here for RaaS, repressors are cotranscribed with their regulon (14). RaaS is a transcriptional regulator that plays an important role in long-term mycobacterial survival in vitro and in vivo (5). Moreover, RaaS mediates an improvement in mycobacterial survival after exposure of nonreplicating bacteria to antimicrobial agents targeting cell wall biosynthesis. Here, we have demonstrated that the RaaS-DNA complex is regulated by acyl-CoAs, specifically oleoyl-CoA and, to a lesser extent, stearoyl-CoA (Figs. 2B and 3). The addition of oleic acid, a precursor of oleoyl-CoA, completely abolished the survival-promoting effects of ethambutol and dysregulated the RaaS regulon (Fig. 5), confirming the biological relevance of oleoyl-CoA as a regulator of mycobacterial persistence.

During the preparation of this manuscript, a crystal structure of RaaS (Rv1219c) was solved (16). The structure analysis confirmed our predictions concerning the importance of Arg-140 and Arg-144, but not Tyr-174, in ligand binding. However, the structure revealed the presence of a very large hydrophobic cavity of 880 Å compared with the calculated volume of the aliphatic chain of oleoyl-CoA of 345 Å (Fig. 7B). The RaaS cavity size significantly exceeds volumes of the binding pocket of YsiA (403 Å) (Fig. 7A) or other mycobacterial TetR regulators such as EthR (399 Å; Protein Data Bank ID 4DW6) and KstR (190 Å; ID 3MNL). This striking difference in the RaaS structure indicates that other longer chain acyl-CoAs or complex lipids containing fatty acids might potentially fit into the binding pocket and
regulate the DNA-binding activity of RaaS. Mycobacteria are able to produce and export a great variety of lipids containing fatty acids from oleic acid itself (17) to acyl-trehaloses (18). It is possible that the RaaS-regulated Rv1218c/Rv1217c pump is involved in the export of these oleate-containing lipids. Moreover, DrrC (which may also be regulated by RaaS) has been implicated in the export of the complex lipid phthiocerol dimycocerosate in *M. tuberculosis* (19).

Derivatives of fatty acids have been previously demonstrated to regulate the expression of mycobacterial lipid transporters (20). Our findings support the importance of acyl-CoA metabolites in the regulation of mycobacterial efflux and long-term persistence. As free acyl-CoA accumulates, RaaS is released from its DNA-binding site, and downstream genes encoding efflux pumps are expressed (Fig. 8A). Depletion of free acyl-CoA allows RaaS to bind to DNA, resulting in repression of the downstream transporter genes (Fig. 8B).

ATP-dependent efflux pumps are integral components of the energy and metabolic circuitry of growing cells, and their activity depends on the metabolic state of bacteria, which may be finely tuned by metabolites binding to transcriptional regulators. In non-growing cells, the levels of acyl-CoAs drop following the slowing down of metabolism. Under stressful conditions (for example, hypoxia or iron limitation), mycobacteria can redirect carbon flux from the tricarboxylic acid cycle to alternative metabolic pathways and to the synthesis of triacylglycerols (21, 22). This mechanism is important for long-term survival, as it generates storage compounds, removes toxic fatty acid derivatives, and depletes free fatty acids and their CoA derivatives (21, 22).

We hypothesize that under stressful conditions or to survive long-term during infection, free RaaS binds to its DNA recognition sequence, repressing the transcription of this cluster of efflux pump genes and itself in a classical feedback loop (Fig. 8).
We further propose that antimicrobial treatment potentiates this process, directly or indirectly inhibiting fatty acid synthases and rapidly depleting residual (C_{16–C_{18}} chain) fatty acid precursors (23–25).

We have shown that RaaS is a component of the complex regulatory mechanisms orchestrating a coordinated down-regulation of energy-consuming processes and the activation of long-term persistence. This is supported by our previous findings that *M. tuberculosis* employs the RaaS-mediated mechanism for *in vitro* persistence in prolonged stationary phase and during *in vivo* macrophage and mouse infections (5). Transposon inactivation of Rv1218c results in the highest fitness cost in macrophage culture (26), but the precise physiological role of the Rv1218c/Rv1217c (Bcg_1278c/Bcg_1277c) pump remains unknown. We propose that Rv1218c/Rv1217c likely transports lipid molecule(s) that are important for the initial stages of infection and may become toxic or are dispensable during non-replicating persistence. Our findings suggest that this putative lipid contains fatty acid moieties (oleic and possibly also stearic acids), which are highly abundant in growing mycobacteria (as CoA derivatives) and are rapidly depleted in nonpermissive growth conditions. Our sequence analysis extended to Rv1216c (Bcg_1276c and Bcg_1275c) highlights distant similarities to an *S*-adenosylmethionine-dependent membrane-embedded methyltransferase and a lipid esterase/transferase. Accordingly, these proteins might be predicted to modify a lipid precursor to be secreted by the Rv1218c/Rv1217c (Bcg_1278c/Bcg_1277c) efflux pump. Future experiments will focus on the nature of the molecules transported by Rv1218c/Rv1217c (Bcg_1278c/Bcg_1277c) and elucidation of their precise roles in mycobacterial growth, infection, and persistence. Our findings indicate that dysregulation of efflux pumps can be employed to kill non-growing mycobacteria.

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