Diisonitrile Lipopeptides Mediate Resistance to Copper Starvation in Pathogenic Mycobacteria

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ABSTRACT Bacterial pathogens and their hosts engage in intense competition for critical nutrients during infection, including metals such as iron, copper, and zinc. Some metals are limited by the host, and some are deployed by the host as antimicrobials. To counter metal limitation, pathogens deploy high-affinity metal acquisition systems, best exemplified by siderophores to acquire iron. Although pathogen strategies to resist the toxic effects of high Cu have been elucidated, the role of Cu starvation and the existence of Cu acquisition systems are less well characterized. In this study, we examined the role of diisonitrile chalcolipophores of pathogenic mycobacteria, synthesized by the enzymes encoded by the virulence-associated nrp gene cluster, in metal acquisition. nrp gene cluster expression is strongly induced by starvation or chelation of Cu but not starvation of Zn or excess Cu. Mycobacterium tuberculosis and Mycobacterium marinum strains lacking the nrp-encoded nonribosomal peptide synthetase, the fadD10 adenylate-forming enzyme, or the uncharacterized upstream gene ppe1 are all sensitized to Cu, but not Zn, starvation. This low Cu sensitivity is rescued by genetic complementation or by provision of a synthetic diisonitrile chalcolipophore. These data demonstrate that diisonitrile lipopeptides in mycobacteria are chalcolipophores that facilitate survival under Cu-limiting conditions and suggest that Cu starvation is a relevant stress for M. tuberculosis in the host.

IMPORTANCE Bacterial pathogens and their hosts engage in intense competition for nutrients, including metals. Mycobacterium tuberculosis, the cause of tuberculosis, lives within host macrophages and is subject to diverse stresses, including metal excess and metal limitation. In this study, we demonstrated that the nrp gene cluster, required for M. tuberculosis virulence and which directs synthesis of diisonitrile lipopeptides, mediates copper acquisition. Copper, but not zinc, deprivation strongly induces diisonitrile biosynthesis, and M. tuberculosis strains lacking the nrp gene, or the associated genes fadD10 or ppe1, are all sensitized to copper chelation or copper deprivation. These results establish a copper binding, or chalcolipophore, system in M. tuberculosis and indicate that resistance to copper restriction plays an important role in the ability of this global pathogen to cause infection.

KEYWORDS Mycobacterium tuberculosis, chalcolipophore, copper, metal resistance, nutritional immunity

Bacterial pathogens express complex virulence programs to ensure their survival within the host, including factors that confer resistance to or modify host immunity. Metal acquisition and resistance are one set of mechanisms used by phylogenetically diverse bacterial pathogens (1). Intracellular pathogens live in a nutrient- and metal-limited niche and require active acquisition of metals critical for enzymatic function. For example, host cells impose iron limitation on pathogens, a host strategy that is counteracted by microbial siderophores, high-affinity iron-binding molecules critical for virulence across pathogens (2). Mycobacterium tuberculosis lacking the iron siderophore mycobactin due to either genetic ablation of mycobactin...
biosynthesis (3), interruption of export (4), or chemical inhibition (5) is attenuated in infection models, as is the case for multiple other bacterial pathogens (6).

Despite the extensive characterization of iron acquisition systems, the relevance of acquisition of other trace metals is less well defined. Zinc and copper are essential nutrients and serve as a enzymatic cofactors, but scavenging systems for these metals are less well characterized (7). There is emerging evidence that pathogens use Zn acquisition systems during infection, through either the elaboration of Zn-scavenging molecules, transport systems for Zn (8), or import of host-derived Zn containing molecules (9–11). It was recently shown that yersiniabactin, previously characterized as an iron siderophore, also binds Cu (12, 13) and Zn (14) and that the yersiniabactin system is active during infections (12, 14–17).

In addition to metal limitation, the host also deploys copper and zinc as antimicrobial effectors to limit pathogen growth (18). Macrophages deposit Cu and Zn into the phagosome, and there is substantial evidence that M. tuberculosis experiences high Cu and Zn stress during infection (18–22). Recently, we identified a new pathway in M. tuberculosis that jointly controls nitric oxide and Cu resistance governed by the Rip1 intramembrane protease and the cytoplasmic PdtaS/PdtaR two-component system (19). This study identified the nrp gene cluster (rv0097 to rv0101) as a target of Rip1/PdtaS/R regulation and demonstrated that expression of the nrp locus was the critical determinant of NO resistance. However, despite the important role of Rip1/PdtaS/R in controlling resistance to Cu toxicity, the nrp cluster was not important for resistance to high Cu (19).

The nrp gene cluster (rv0097 to rv0101), named for the nonribosomal peptide synthetase (NRPS) encoded by rv0101, has been implicated in M. tuberculosis virulence in multiple studies (19, 23, 24). M. tuberculosis lacking nrp is attenuated for growth in the lung during early infection but has no persistent virulence defect at later time points. Although these studies implicate the biosynthetic product of the nrp gene cluster in pathogenesis, the identity of this natural product was unknown. Recent studies using heterologous expression of the orthologous nrp gene clusters from Streptomyces thioluteus, Streptomyces coeruleorubidus, and Mycobacterium marinum (without Mmar_0261/ppe1) revealed a series of diisonitriles (compounds 1 to 5) that differed in structure based on the origin of the NRPS (Fig. 1A, B) (25, 26). Examination of the metal binding specificity of the diisonitrile products from heterologous expression (26) or synthesis (27) revealed tight binding to Cu. Further, products of the S. thioluteus nrp cluster expressed in Streptomyces lividans can directly mediate Cu uptake, indicating that diisonitriles function as chalkophores (26). Most recently, the structure of the correspondingly M. tuberculosis NRPS diisonitrile product (compound 7 in Fig. 1B) was identified, and it has several structural differences from the Streptomyces and M. marinum diisonitriles. Whereas diisonitriles 1 to 5 have lysine-based cores, M. tuberculosis diisonitrile 7 has a shorter ornithine-based core, which is also found in the natural product SF2369 (compound 6) (28), a C-terminal phenylalanine extension, attributed to an additional NRPS module found only in Rv0101, and very long (C14 to C20) β-isocyano acyl chains (Fig. 1) (24). However, the metal binding specificity of this M. tuberculosis diisonitrile is not known.

The physiologic roles of the diisonitrile products of the nrp locus in mycobacteria, in particular their roles in metal acquisition and detoxification, are unknown. In mycobacteria, the nrp cluster includes the ppe1 gene (rv0096) as a conserved feature, but the functional importance of ppe1 is unknown. In this study, we used genetic deletion of the nrp gene and the upstream genes ppe1 and fadD10, coupled with assays of metal starvation, to deduce the primary function of the nrp cluster. We show that these elements of the nrp gene clusters are required for resistance to starvation of Cu but not other metals, establishing the products of these gene clusters as chalkophores of pathogenic mycobacteria. Further, we show that a synthetic diisonitrile can rescue growth of nrp gene cluster mutants under Cu restriction, indicating that mycobacteria can accept this nonnative diisonitrile as a chalkophore.

RESULTS

The nrp gene cluster is transcriptionally induced under copper starvation or chelation but does not respond to Zn starvation. Our prior experiments determined that the nrp gene cluster is induced by nitric oxide and is a target of the Rip1/PdtaS/R regulatory cascade that controls NO resistance (19). However, although this Rip1/PdtaS/PdtaR system controls
Cu resistance, the nrp gene cluster was not involved in the Cu resistance arm of the pathway (19), despite the high Cu affinity of related diisonitrile products produced by Streptomyces (26, 27), suggesting an alternative role in Cu homeostasis. To understand the function of the nrp gene cluster in the mycobacterial metal response, we measured the levels of the mRNAs encoding multiple proteins in the cluster from rv0096 through rv0101 (NRPS) under conditions of metal limitation. We first treated Mycobacterium tuberculosis with tetrathiomolybdate (TTM), a Cu chelator, and measured each mRNA by real-time quantitative PCR (RT-qPCR). All genes in the cluster were induced by TTM 10- to 100-fold compared to untreated cells (Fig. 2A). The ppe1-5′ RNA at the 5′ end of the PPE1 open reading frame (ORF), previously identified as induced by nitric oxide (19), was the most highly induced by TTM (Fig. 2A). To measure the PPE1 protein, we engineered an M. tuberculosis strain with a functional (Fig. S5B) C-terminal green fluorescent protein (GFP) fusion to the chromosomal copy of PPE1 and treated it with TTM. TTM treatment strongly induced PPE1 protein expression (12-fold) compared to vehicle-treated cells (Fig. 2B). Treatment with the Zn selective chelator N,N,N′,N′-tetrakis(2-pyridylmethyl)ethylenediamine (TPEN) at a concentration known to impose Zn limitation on M. tuberculosis (29) did not alter PPE1-GFP expression, suggesting that the effect is specific to Cu chelation (Fig. 2A). To measure the PPE1 protein, we engineered an M. tuberculosis strain with a functional (Fig. S5B) C-terminal green fluorescent protein (GFP) fusion to the chromosomal copy of PPE1 and treated it with TTM. TTM treatment strongly induced PPE1 protein expression (12-fold) compared to vehicle-treated cells (Fig. 2B). Treatment with the Zn selective chelator N,N,N′,N′-tetrakis(2-pyridylmethyl)ethylenediamine (TPEN) at a concentration known to impose Zn limitation on M. tuberculosis (29) did not alter PPE1-GFP expression, suggesting that the effect is specific to Cu chelation (Fig. 2A). To confirm that the effect of TTM was due to Cu limitation, we depleted mycobacterial 7H9 medium of all metals using Chelex ion exchange resin and then reconstituted it with defined metal concentrations, leaving out either Cu or Zn. We confirmed that our Zn-depleted medium imposed Zn limitation by measuring the induction of ctpV, previously reported to be one of the most strongly induced genes in M. tuberculosis exposed to low Zn (29). Low Cu weakly induced ctpV, but, consistent with prior reports, low
Zn strongly induced ctpV RNA, indicating that our zinc-depleted medium imposed zinc starvation (see Fig. S1 in the supplemental material). When bacteria were grown in Cu-depleted medium, PPE1 protein was strongly induced, but Zn starvation alone had no effect (Fig. 2C). We observed that starvation of Zn did further induce PPE1 when Cu was limiting, suggesting that Zn influences expression of the 

nrp locus only when Cu is limiting (Fig. 2C).

The *M. tuberculosis* regulon induced by high Cu has been extensively studied (20, 21, 30), and the *nrp* gene cluster has not been identified as responsive to high Cu. To confirm these prior results, we treated cells with 500 μM CuSO₄ and measured the mRNA of each gene in the *nrp* cluster. Consistent with prior data, high Cu did not induce any gene in the *nrp* cluster, whereas *mymT*, previously identified as regulated by the Cu-responsive repressor RicR (20, 31), was highly induced (Fig. 2D). These data indicate that *nrp* gene cluster expression is

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**FIG 2** The chalkophore gene cluster is induced under low-copper conditions. (A) Normalized mRNAs from the chalkophore cluster genes, indicated on the x-axis, measured by RT-qPCR in *M. tuberculosis* treated with DMSO (vehicle [blue]) or 20 μM tetrathiomolybdate (TTM). Each point represents a biologic replicate, and values are normalized to sigA under the DMSO condition. (B) Immunoblotting with anti-GFP or anti-RpoB antibodies of protein lysates from a strain with a functional C-terminal translational fusion of GFP to the C terminus of PPE1 (PPE1 stop::GFP Hyg), treated with DMSO, 10 μM TTM, or 10 μM TPEN. Quantitation of replicate immunoblots is given in the bar graph. GFP signal was normalized to RpoB in the same lane and then relative to DMSO, which is set at 1. AU, arbitrary units. (C) Immunoblotting of biologic duplicates as in panel B in cells grown in metal-replete medium or medium lacking Zn, Cu, or Cu/Zn and corresponding quantitation. GFP signal was normalized to RpoB in the same lane and then relative to replete medium, which is set at 1. (D) mRNAs of the chalkophore cluster are not induced by high Cu. The indicated mRNAs were quantitated as in panel A in wild-type *M. tuberculosis* grown in 3.5 or 500 μM Cu. Each point is the average of technical triplicates for 3 biologic replicates. (E) Chalkophore transcription is autoregulated. Shown are results for mRNA quantitation by RT-qPCR of *ppe1* S*, ppe1*, or *fadD10* in wild-type *M. tuberculosis* (WT), *M. tuberculosis Δnrp*, or *M. tuberculosis Δnrp* plus *nrp* (comp) with or without treatment with TTM. Error bars indicate SEM.
primarily responsive to Cu but not Zn limitation, imposed by either chelation or metal starvation, but not Cu excess.

**Chalkophore biosynthesis is transcriptionally autoregulated.** Many bacterial regulons are subject to feedback regulation such that the biosynthetic end product, or an intermediate, represses expression of the genes that lead to synthesis of that product. To determine whether expression of the chalkophore cluster is affected by the production of the diisonitrile product, we measured the RNAs of *ppe1-5*, *ppe1*, and *fadD10* in an *M. tuberculosis* strain lacking the *nrp* gene (Fig. S2), which should not produce chalkophores, with and without TTM. In wild-type cells, TTM induced expression of all three mRNAs, as reported above (Fig. 2E). In the Δ*nrp* strain, all three mRNAs were induced without Cu chelation to a level similar to that in wild-type cells with chelation (Fig. 2E). Treatment with TTM had a small incremental inductive effect on each RNA, and complementation with the wild-type *nrp* gene restored the inducibility pattern of wild-type cells, indicating that these phenotypes are due to loss of *nrp* and chalkophore biosynthesis. These data strongly indicate that the expression of chalkophore biosynthesis machinery is repressed by the diisonitrile chalkophore product, either directly or indirectly.

**Loss of *nrp* confers sensitivity to copper chelation or starvation but not zinc chelation or starvation.** To determine the contribution of isonitrile lipopeptide biosynthesis to mycobacterial growth under metal-limiting conditions, we tested *M. marinum* (*nrp::tn*) or *M. tuberculosis* (Δ*nrp*) lacking *nrp* for sensitivity to TTM. Wild-type *M. marinum* growth was unaffected by 20 μM TTM, whereas growth of *M. marinum* lacking *nrp* arrested after several doublings (Fig. 3A). Complementation of *M. marinum* *nrp::tn* with *M. marinum* *nrp* restored TTM resistance to wild-type levels.

Bathocuproinedisulfonic acid (BCS) is a copper chelator with an association constant for 

Cu(I) of 10^{19.8} M^{-2} (32) in 2:1 stoichiometry. Addition of BCS did not affect growth of wild-type *M. marinum* but arrested growth of *M. marinum* lacking *nrp* (Fig. S3). This is consistent with a previous study that showed that a synthetic diisonitrile chalkophore, related to the predicted structure of the *M. marinum* natural product diisonitrile, has much higher affinity for Cu(I) than does BCS (27). To confirm that the effect of BCS is mediated by Cu starvation, we supplemented BCS with Cu, Zn, Mn, and Fe and observed that only Cu could reverse the growth limitation of *nrp*-deficient *M. marinum* by BCS (Fig. S3). In addition to limiting growth, TTM treatment was lethal for *nrp*-deficient *M. marinum*, a phenotype that was also rescued by genetic complementation (Fig. 3B).

For *M. tuberculosis*, growth of the wild-type Erdman strain was also unaffected by TTM, but *M. tuberculosis* Δ*nrp* ceased replication after several doublings, a phenotype that was complemented by reintroduction of *nrp* (Fig. 3C). We confirmed that *M. tuberculosis* lacking *nrp* was sensitized to both BCS and TTM on agar media (Fig. S4). To test whether chelation of zinc had a similar effect, we repeated this experiment with the Zn chelator TPEN. Addition of 10 μM TPEN had no effect on growth of either wild-type *M. marinum* or *M. marinum* *nrp::tn*, indicating that the *M. marinum* NRPS (mmaA) function is not required to resist Zn chelation (Fig. 3D).

To extend these results, we tested whether isonitrile lipopeptide biosynthesis is required for growth under Cu-limiting conditions using the media preparation strategy described above. Although wild-type cells grew in media depleted of both copper and zinc, both *M. marinum* (Fig. 3E) and *M. tuberculosis* lacking *nrp* (Fig. 3F) were unable to grow in medium without added Cu and Zn, a phenotype that was complemented by wild-type *nrp* in both species. Supplementation of this medium with Zn did not restore growth of *nrp*-deficient cells but supplementation with Cu did, indicating that *nrp* is required for growth in low Cu but not low Zn (Fig. 3E and F), recapitulating the results with Cu and Zn chelation. These experiments suggest that *nrp* function is uniquely required for bacterial growth under Cu-limiting conditions and that mycobacterial chalkophores function in Cu but not Zn acquisition.

**Loss of *FadD10* confers sensitivity to low copper.** Although the *nrp* gene cluster can direct isonitrile chalkophore biosynthesis in heterologous bacteria (25, 26), the individual roles of the genes in the chalkophore cluster for diisonitrile biosynthesis are not known. *FadD10* (encoded by *rv0099*) is an upstream adenylate-forming enzyme (33) that is involved in synthesis and coupling of the characteristic β-isocyanol fatty acid chains in the natural...
To determine whether FadD10 (35) is required for resistance to copper starvation, we tested *M. marinum* and *M. tuberculosis* mutants lacking fadD10. *M. marinum fadD10::tn* was severely growth restricted in the presence of 10 μM TTM, a phenotype that was complemented by the wild-type *M. marinum* or *M. tuberculosis fadD10* gene (Fig. 4A). *M. marinum fadD10::tn* also could not grow in Cu-depleted medium, a phenotype that was also genetically complemented by *M. marinum* or *M. tuberculosis fadD10*, whereas loss of fadD10 had no effect on growth without Zn (Fig. 4B). We observed a similar effect with *M. tuberculosis*, in which a ΔfadD10 mutant was deficient for growth in TTM or low Cu but not low Zn (Fig. 4C and D). To determine whether the activity of FadD10 is required for low-Cu growth, we mutated aspartate 408 to alanine (D408A), as this residue directly interacts with the ribose 3'-hydroxyl of the cognate fatty acyl-AMP reaction intermediate in the FadD10

![Image](FIG 3)

**FIG 3** Loss of *nrp* confers sensitivity to copper chelation or limitation. (A) Wild-type *M. marinum* (circles), *M. marinum nrp::tn* (squares), *M. marinum nrp::tn* plus *M. marinum nrp* (triangles) treated with vehicle DMSO (blue) or 10 μM TTM (red). Dilution-corrected OD of a continuous growth curve is shown, with each point the mean of triplicate cultures and error bars displaying SEM. **, *P < 0.01 by unpaired test for each time point of *nrp::tn* versus wild-type or complemented strain. (B) Bacterial killing assays of the same strains as in panel A, treated with 10 or 20 μM TTM as described in Materials and Methods. Bacterial survival was quantitated on agar medium without TTM. Each dot represents a biologic replicate. ***, *P < 0.001; **, *P < 0.01. (C) Extended growth curves of the indicated *M. tuberculosis* strains with (red) or without (blue) TTM. **, *P < 0.01; *, *P < 0.02. (D) Growth of the same *M. marinum* strains as in panel A with 10 μM Zn chelator TPEN. (E) Growth of *M. marinum* strains in medium lacking Cu and Zn (blue), Cu alone (red), or Zn alone (yellow). **, *P < 0.0001. (F) Growth of *M. tuberculosis* strains in medium lacking Cu (red) or Zn alone (yellow). Error bars for all panels indicate SD and when not visible are within the symbol.
FIG 4  FadD10 activity is required for low copper tolerance. (A) Wild-type *M. marinum* (circles), *M. marinum fadD10::tn* (squares), *M. marinum fadD10::tn plus M. marinum fadD10* (triangles), and *M. marinum fadD10::tn plus M. tuberculosis fadD10* (inverted triangles) treated with the vehicle DMSO (blue) or 10 μM TTM (red). Dilution-corrected OD of a continuous growth curve (see Materials and Methods) is shown, with each point the mean of triplicate cultures and error bars displaying SEM. **, P < 0.01 by unpaired test for each time point of fadD10::tn versus the wild type. (B) FadD10 is required for growth in low Cu but not low Zn. The indicated strains were grown in medium lacking either Cu (red) or Zn (yellow). **, P < 0.01 by unpaired test for each time point of fadD10::tn versus the wild type. Error bars display SEM. (C) *M. tuberculosis* wild type (circles) or *M. tuberculosis ΔfadD10* (squares) treated with either DMSO (blue) or TTM (red). **, P < 0.01 by unpaired test for each time point of ΔfadD10 versus the wild type. (Continued on next page)
structure (35). To confirm protein expression, we inserted an ALFA tag (36) at the C termini of both wild-type FadD10 and FadD10(D408A). The two proteins were expressed to similar levels as detected by an ALFA-specific nanobody (Fig. 3E). Wild-type FadD10 ALFA fully rescued M. tuberculosis ΔfadD10 under conditions of low Cu growth and TTM chelation (Fig. 4F and G). However, FadD10(D408A) could not rescue growth under either condition, suggesting that FadD10 substrate binding is required for chalkophore biosynthesis.

**Chemical rescue of rnp deficiency with synthetic diisonitrile.** The data above indicate that loss of two genes in the rnp operon, rnp (rv0101) itself and the upstream fadD10 (rv0099), encoding an adenylate-forming enzyme, confers sensitivity to low copper or Cu chelation. These phenotypes are genetically complemented and are therefore due to deficiency in NRPS or FadD10 function. Prior work demonstrated that the rnp gene cluster from Streptomyces or M. marinum, when expressed in Escherichia coli, directs synthesis of a diisonitrile lipopeptide that bind Cu with high affinity (26). Total synthesis of the Streptomyces thioluteus-derived diisonitrile 7F2768 and related linear congeners confirmed that these molecules bind Cu(I) in a 2:1 stoichiometry with a Cu affinity that exceeds that of BCS, but they failed to demonstrate strong Zn binding (27). To test whether the low-Cu phenotype of mycobacterial cells lacking rnp or fadD10 is due to deficiency of the diisonitrile product, we attempted to rescue the low-Cu phenotype in trans with a synthetic chalkophore.

Because mycobacterial chalkophore products have not been synthesized, we used a lysine-derived congener (compound 4 in reference 27 and compound 3 in Fig. 1B) with the caveat that this molecule may not be used efficiently by M. marinum or M. tuberculosis, particularly given that it has 4-carbon β-isonitrile side chains, much shorter than those attributed to the mycobacterial diisonitriles (24, 25). Despite this caveat, addition of 10 μM synthetic diisonitrile partially rescued TTM-imposed growth restriction of M. marinum lacking rnp (Fig. 5A). The TTM-imposed growth restriction of M. tuberculosis lacking fadD10 was also substantially reversed by the synthetic chalkophore (Fig. 5B). The synthetic chalkophore also fully rescued M. marinum lacking rnp from growth restriction by BCS (Fig. 5C). These results demonstrate that the sensitivity of mycobacteria lacking fadD10 or rnp to Cu starvation is due to diisonitrile chalkophore deficiency and further indicate that the mycobacterial chalkophore system can utilize nonnative diisonitriles as functional chalkophores.

**PPE1 is required for growth in low copper.** The first gene in rnp gene cluster is ppe1 (rv0096c) (Fig. 1A). The data above indicate that the PPE1 protein and the small RNA ppe1-5′, which is at the 5′ end of the ppe1 locus (19), are both strongly induced by copper starvation (Fig. 2). PPE1 is predicted to be a multipass transmembrane protein, but prior heterologous-expression studies that identified the diisonitrile products of the M. marinum rnp locus did not include PPE1, suggesting that it is not required for diisonitrile biosynthesis (25). To determine whether PPE1 is also required for resistance to Cu starvation, we tested M. marinum ppe1::tn under low-Cu and low-Zn conditions. M. marinum lacking ppe1 grew identically to wild-type M. marinum in replete medium (Fig. 6A). However, chelation of Cu with TTM impaired growth of M. marinum lacking ppe1, a phenotype that was rescued by reintroduction of ppe1 (Fig. 6B). PPE1 is also required for M. marinum growth under Cu-limiting conditions but not Zn-limiting conditions, a phenotype that resembles loss of fadD10 or rnp (rv0101) (Fig. 6C). Loss of ppe1 severely sensitized M. tuberculosis to both BCS and TTM on agar medium (Fig. 5A). Combined with data above showing that ppe1 is coregulated by low Cu with the other genes in the chalkophore cluster, these functional data indicate that PPE1 is an integral component of the chalkophore system required for resistance to Cu starvation.

**FIG 4 Legend (Continued)**

test for each time point of ΔfadD10 versus the wild type. Error bars display SEM. (D) M. tuberculosis wild type (circles) or M. tuberculosis ΔfadD10 (squares) grown in medium lacking Zn (yellow) or Cu (red). **, P < 0.01 by unpaired test for each time point of ΔfadD10 versus the wild type. Error bars display SD. (E) Whole-cell protein lysates of M. tuberculosis ΔfadD10 with either plasmid vector or plasmids encoding C-terminally ALFA-tagged FadD10 or FadD10(D408A) were separated by SDS-PAGE and probed with antibodies to RpoB as a loading control or an anti-ALFA tag nanobody. (F) Low-Cu growth curves of the indicated strains from panel E. **, P < 0.01 by unpaired test for each time point of ΔfadD10:tn+ vector versus ΔfadD10:tn+fadD10 or ΔfadD10:tn+fadD10 versus ΔfadD10:tn+fadD10(D408A). Error bars display SEM. (G) Growth of the induced strains as in panel F with or without TTM. Significance comparisons are the same as in panel F. Error bars display SEM.
DISCUSSION

nrp, fadD10, and ppe1 are required for copper acquisition. Our data strongly support a model in which M. tuberculosis and M. marinum use diisonitrile lipopeptides as a copper acquisition system under low-Cu conditions. Loss of chalkophore biosynthesis through genetic deletion of the nrp (rv0101) gene, fadD10, or ppe1 confers sensitivity to Cu chelation or Cu starvation. These phenotypes are rescued by a synthetic diisonitrile chalkophore, indicating that the phenotypes of nrp cluster mutants are due to deficiency of the diisonitrile product. PPE1 is predicted to be a membrane protein, and we speculate that its role may be in chalkophore transport, as other members of this gene family have been implicated in transport of other nutrients in mycobacteria, including glycerol, phosphate, and heme (37–39). The role of chalkophores in low-Cu growth is thematically similar to the iron acquisition role of siderophores, which are widely utilized by bacterial pathogens to acquire iron within the host. Multiple groups have reported that M. tuberculosis lacking nrp is attenuated in mouse models of infection (19, 23), strongly suggesting that M. tuberculosis experiences Cu-limiting environments in the host and that diisonitrile chalkophores serve to counter this host-imposed Cu limitation.

A recent study by Mehdiratta et al. examined the function of the NRPS encoded by nrp (rv0101) and its natural product in M. tuberculosis (24). The investigators identified the diisonitrile product produced by M. tuberculosis and demonstrated the accumulation of this lipopeptide in infected mouse lung, with a time course of accumulation that matches the pattern of attenuation of M. tuberculosis lacking nrp. Importantly, this study identified the previously unknown structure of the M. tuberculosis diisonitrile, which differs from the M. marinum diisonitrile due to differences between the biosynthetic modules of the M. tuberculosis NRPS protein and the M. marinum NRPS. However, our findings and conclusions about the role of diisonitriles in metal acquisition differ substantially from those of Mehdiratta et al., who concluded that the major function of the product of the nrp cluster is both to acquire Zn and to resist Zn toxicity. In contrast, our data do not indicate a role of the nrp gene cluster in Zn acquisition but indicate an important role in Cu acquisition.
are several methodologic differences that could contribute to our different conclusions. In the work of Mehdiratta et al., the metal limitation assays were performed in Chelex-treated Sauton’s medium, removing all metals, including those ordinarily present in this medium (iron and magnesium), with single metals then added back and bacterial growth measured. Under these conditions, only Zn was able to rescue the defective growth of \textit{M. tuberculosis} lacking \textit{nrp}; Cu did not. In contrast, our assays use a dropout approach in which all metals are depleted and then replaced, with the exception of individual metals. Thus, \textit{M. tuberculosis} requires \textit{nrp} to grow when Cu is lacking but can grow when Zn is the only metal provided.

Our data support a primary role for chalkophores in tolerance to copper limitation, rather than zinc. Prior data obtained using synthetic diisonitrile SF2768, and the same compound purified from heterologous expression, indicated that diisonitriles bind Cu, but not Zn, with high affinity. In addition, in \textit{Streptomyces}, SF2768 has been shown to mediate Cu uptake into cells directly (26). Our data support a similar role in \textit{M. tuberculosis} and \textit{M. marinum} based on the following observations:

1. The genes in the \textit{nrp} gene cluster are responsive to Cu limitation but not Zn limitation.
2. \textit{M. tuberculosis} and \textit{M. marinum} lacking either \textit{nrp}, \textit{fadD10}, or \textit{ppe1} are growth inhibited in Cu- but not Zn-deficient medium or in the presence of Cu but not Zn chelators.
3. The phenotypes of chalkophore-deficient \textit{M. tuberculosis} and \textit{M. marinum} are rescued by a synthetic diisonitrile, which binds Cu but not Zn. Although this synthetic diisonitrile differs in structure from the recently identified diisonitrile natural product from \textit{M. tuberculosis} and the predicted diisonitrile natural product from \textit{M. marinum} (24, 25), we show that it is utilized as a functional chalkophore by

![FIG 6](image-url)
mycobacteria and its defined metal binding specificity for Cu further supports a model in which diisonitriles function as chalkophores.

It remains possible that the structure of the *M. tuberculosis* diisonitrile product confers some binding to Zn and an ability to mediate its uptake when Zn is the only metal available, a hypothesis that will require total synthesis of the *M. tuberculosis* natural product. Our data do indicate that expression of PPE1, although primarily responsive to low Cu, is enhanced by low Zn when Cu is also absent, suggesting some interaction between Zn and Cu starvation. Prior data (29), which we reproduce here, indicate that low zinc stimulates expression of *cptV*, which encodes a copper exporter under the control of CsoR (40–42). This low-zinc response would result in copper efflux, potentially lowering the cytoplasmic copper concentration and explaining an interaction between deprivation of zinc and deprivation of copper.

**Roles of copper acquisition by chalkophores.** The best-studied chalkophore systems are encoded by *Methanococcus*. The chalkophores in methanococci are ribosomally produced, posttranslationally modified peptides that bind copper with high affinity (43, 44). *Methanococcus* chalkophores mediate Cu import and eventual loading onto the membrane-bound Cu-dependent enzyme particulate methane oxygenase. Although the mycobacterial chalkophores differ in chemical structure and Cu binding chemistry, the mycobacterial diisonitriles appear to play similar roles in Cu acquisition. However, the critical cellular function of Cu that is maintained by the mycobacterial chalkophore system under conditions of Cu limitation, both in vitro and in the host, remains to be determined. Given the requirement for the chalkophore system in *M. tuberculosis* virulence, the findings reported here may suggest that Cu limitation is imposed by the host during infection. This conclusion adds complexity to the existing paradigm that high-Cu stress imposed on phagosomal pathogens in macrophages is the most relevant Cu component of nutritional immunity. Direct measurements of the metal concentrations in *M. tuberculosis* containing phagosomes of murine macrophages have revealed Cu concentrations that range from approximately 400 μM immediately after infection to 25 μM after 24 h (45), concentrations that are sufficient to induce the Cu-responsive CsoR regulon, indicating that *M. tuberculosis* experiences high Cu in macrophages (18). Zn concentrations are approximately 40 μM immediately after infection, rising to ~426 μM 24 h after infection (45), a concentration that is sufficient to induce the Zn efflux pump CtpC (18). The role of the *nrp*-dependent diisonitrile natural products in Cu acquisition, but not resistance to Cu toxicity, may suggest that Cu starvation paradoxically coexists with high Cu stress during infection. Additional support for the importance of the Cu starvation response in *M. tuberculosis* pathogenesis comes from analysis of *M. tuberculosis* lacking the SigC alternative sigma factor (46). *M. tuberculosis* lacking sigC is sensitive to Cu, but not Zn, limitation, underexpresses the chalkophore cluster genes in low Cu, and is attenuated in SCID mice (46). Low and high Cu stresses could be imposed on *M. tuberculosis* in distinct cellular or anatomic compartments, imposed by distinct effectors of host immunity, or in distinct phases of infection (i.e., acute versus chronic versus latent). Tissue-specific differences in Cu availability have been found in *Candida albicans* infection, in which infection of the kidney imposes a Cu limitation response and infection of the blood imposes a high Cu response (47). Similarly, although Zn and Cu intoxication is imposed on *M. tuberculosis* in macrophages (18, 20, 21), the casemum in which *M. tuberculosis* resides in nonhuman primate lung cavities is rich in calprotectin (29), which can bind multiple metals (48, 49), supporting the idea the metal excess and metal limitation may coexist in *M. tuberculosis* infection.

In summary, the data presented here show that diisonitrile lipopeptides produced by the *nrp* locus function as chalkophores in copper acquisition and will empower further investigation of the roles of chalkophores and Cu restriction in nutritional immunity to *M. tuberculosis*.

**MATERIALS AND METHODS**

*Reagents.* Middlebrook 7H10 agar, 7H9 broth, dextrose, Tween 80, bovine serum albumin (BSA), and UltraPure DNase/RNase-free distilled water were purchased from Fisher Scientific. Ammonium tetraethyldisulfoborate (TTM), N,N,N’,N’-tetraisopropylcyclotetrasiloxane (TPEN), dimethyl sulfoxide (DMSO), copper sulfate, zinc sulfate, magnesium sulfate, and calcium chloride were purchased from Millipore Sigma. Biotechnology (BT) grade Chelex 100 resin, sodium form, was purchased from Bio-Rad. The synthetic diisonitrile chalkophore was synthesized and characterized as previously described (compound 4 in reference 27).
**General growth conditions, strains, and DNA manipulations.** *M. tuberculosis* Erdman and *M. marinum* M strain were grown and maintained in 7H9 medium (broad spectrum broth) or on 7H10 agar supplemented with 10% oleic acid-albumin-dextrose-catalase (OADC) and 0.05% glycerol (7H9 OADC/7H10 OADC). Broth cultures were additionally supplemented with 0.02% Tween 80. Genetic mutations were generated by specialized transduction utilizing the temperature-sensitive phage phAE87 (50). Mutant strains were confirmed by PCR (see Fig. S2 in the supplemental material) followed by sequencing. *M. marinum* transposon mutants were a generous gift from the Tobin Lab at Duke University. For a complete strain list with relevant features, see Table S1. Plasmids utilized in this study were generated using standard molecular techniques and are listed with their features in Table S2.

**Chelated copper- and zinc-depleted 7H9 media and buffers.** Ion-deficient buffers were generated by treatment with 10 g/L of biotechnology-grade Chelex resin for 24 h at room temperature, followed by 0.2-μm filtration.

7H9 medium was generated from primary components as follows: 0.5 g/L of ammonium sulfate, 1 g/L of monopotassium phosphate, 2.5 g/L of disodium phosphate, 0.1 g/L of sodium citrate, 0.5 g/L L-Glutamic acid, 0.001 g/L of pyridoxine hydrochloride, and 0.0005 g/L of biotin were dissolved in 850 mL of deionized water. Once dissolved, 10 mL of 50% glycerol, 2.5 mL of 20% Tween 80, and 100 mL of ADS (albumin, dextrose, and saline) supplement were added and the total volume was adjusted to 990 mL with deionized water. The resulting solution was then treated with 10 g/L of Chelex resin for 24 h at 4°C. Resin was removed by 0.2-μm filtration, and iron, magnesium, and calcium levels were restored to 0.04 g/L, 0.05 g/L, and 0.0005 g/L, respectively, by the addition of 4, 5, and 0.5 mL of 10-mg/mL sterile filtered solutions of ferric ammonium citrate, magnesium sulfate, and calcium chloride dissolved in UltraPure distilled water. Sterile filtered 10-mg/mL solutions of copper sulfate and/or zinc sulfate dissolved in UltraPure distilled water were used to restore Cu and/or Zn levels to 0.001 g/L, when indicated.

**Cation limitation and chelation sensitivity growth assays.** For liquid growth assays, the indicated strains were pregrown in nonchelated medium until reaching an optical density at 600 nm (OD600) of ~1.0. Cells were then collected by centrifugation (3,700 × g, 10 min) and washed twice with chelated phosphate-buffered saline plus 0.02% Tween 80 (PBS-Tween 80). Growth assays were initiated at a calculated OD600 of 0.005 by the addition of 1 mL of washed culture at an OD600 of 0.05 to 9 mL of chelated 7H9-ADS deficient in either copper or zinc ions or in chelated 7H9-ADS replete for all ions. Selective ion chelators and synthetic chelophore analogues were assayed in chelated 7H9-ADS replete for all ions. *M. tuberculosis* and *M. marinum* growth assays were conducted at 37°C and 30°C, respectively, with growth assays via daily OD600 measurements. To maintain logarithmic growth and ensure that all cells entered log phase to remove any confounding effects of extended lag phase of any mutant or condition, cultures were diluted 1:10 upon reaching an OD60 of 0.4, and thus, the y axes of many growth curve plots show “dilution corrected OD600” that incorporates this dilution factor. For *M. marinum* and *M. tuberculosis* TTM survival assays, cells were pregrown and washed as indicated above. Replicate 2-mL tubes containing chelated 7H9-ADS replete for all ions and the indicated concentrations of TTM, or a DMSO vehicle control, were inoculated at an OD600 of 0.01. After 7 days of incubation at 30°C (*M. marinum*) or 37°C (*M. tuberculosis*), aliquots were removed and plated on nonchelated 7H10-OADC plates to determine growth/survival. CFU per microliter of culture were determined after 12 days of incubation at 30°C (*M. marinum*) or 21 days at 37°C (*M. tuberculosis*).

**RNA extraction and RT-qPCR.** For TTM-stimulated RT-qPCR, the indicated strains were pregrown and washed as described above. Triplicate 30-µL cultures of chelated 7H9-ADS replete for all ions were then inoculated at an OD600 of 0.02. Upon reaching an OD600 of 0.6 to 0.7 cultures were treated with the indicated concentration of TTM, or a DMSO vehicle control, for 5 h. Cells were then collected by centrifugation (3,700 × g, 10 min), suspended, and stored in 1 mL of TRIzol reagent for downstream RNA extraction and processing described below. For ion-deficient RT-qPCR assays, 100-µL cultures of *M. tuberculosis* (Erdman) were pregrown in nonchelated 7H9 OADC medium to an OD600 of ~2.0. Cells were collected by centrifugation (3,700 × g, 10 min), washed in chelated PBS-Tween 80, and concentrated to an OD600 of 6.0. Triplicate cultures were then inoculated at a starting OD600 of 0.2 by the addition of 1 mL of washed culture to 29 mL of chelated 7H9-ADS containing the indicated complement of ions. At 48 h after inoculation, cells were collected by centrifugation (3,700 × g, 10 min), suspended, and stored in 1 mL of TRIzol reagent for downstream RNA extraction and processing described below.

Cells in 1 mL of TRIzol were mechanically disrupted with zirconia bead via three 30-s pulses in a BioSpec Mini24 BeadBeater. Following lysis, beads were removed by centrifugation at 20,000 × g for 5 min and total RNA was isolated using the Direct-zol RNA miniprep kit (Zymo Research) as directed by the manufacturer. Contaminating genomic DNA was removed using the Turbo DNA-free kit (Invitrogen). A total of 500 ng of resulting total RNA was used to synthesize cDNA via random priming utilizing the Maxima H-Minus cDNA synthesis kit (Thermo Fisher). Real-time qPCR was performed on a 7500 real-time PCR system (Applied Biosystems). The amplification product was detected by SYBR green using the Dynamo Flash qPCR kit (Thermo Fisher). For each gene of interest (GOI), normalized cycle threshold (C<sub>T</sub>) was determined relative to the housekeeping gene *sigA*. Relative expression level was calculated using the formula 2<sup>-ΔΔCT</sup> where CT = C<sub>TMax</sub> - C<sub>TGOI</sub> and Primer sets used to amplify individual GOI are listed in Table S3.

**Immunoblotting.** Lysates for immunoblotting were prepared from duplicate cultures of the indicated strains and washed and treated as described for RNA extraction above, except that postcentrifugation pellets were washed once with 1 mL of lysis buffer (350 mM sodium chloride, 20 mM Tris (pH 8.0), 1 mM 2-mercaptoethanol) prior to suspension in 0.8 mL of lysis buffer plus ~100 µL of zirconia beads. Lysis was performed by three 45-s pulses in a BioSpec Mini24 BeadBeater with 5-min intervening rest periods on ice. Beads and debris were removed by centrifugation at 20,000 × g for 15 min at 4°C, and the resulting supernatant was mixed 1:1 with 2× Laemmli sample buffer supplemented with 0.1 M...
dithiothreitol (DTT). Twenty microliters of each sample, heated for 10 min at 100°C, was then separated on 4 to 12% NuPAGE bis-Tris polyacrylamide gels. Separated proteins were transferred to nitrocellulose and probed with the appropriate antibodies. Antibodies used in this study were monoclonal anti-E. coli RNA polymerase β (BioLegend), rabbit polyclonal anti-GFP (Rockland), and sdAb anti-ALFA tag-horseradish peroxidase (HRP) (NanoTag Biotechnologies). PPE1-GFP blots were quantitated using ImageJ software. Relative GFP signal was normalized to corresponding Rpoβ loading control levels and DMSO or replete-ion complement controls.

Data availability. All strains used in this work are available from the corresponding author upon request. There is no external data set associated with the paper. The uncropped blots for Fig. 2B and C and Fig. 4E are available for review upon request and were included in the submission materials for the manuscript.

SUPPLEMENTAL MATERIAL
Supplemental material is available online only.

FIG S1, TIF file, 0.1 MB.
FIG S2, TIF file, 2.8 MB.
FIG S3, TIF file, 0.8 MB.
FIG S4, TIF file, 2.6 MB.
FIG S5, TIF file, 2.4 MB.
TABLE S1, XLSX file, 0.01 MB.
TABLE S2, XLSX file, 0.01 MB.
TABLE S3, XLSX file, 0.01 MB.

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REFERENCES
1. Murdoch CC, Skaar EP. 31 May 2022. Nutritional immunity: the battle for nutrient metals at the host–pathogen interface. Nat Rev Microbiol https://doi.org/10.1038/s41579-022-00745-6.
2. Mey AR, Gómez-Garzón C, Payne SM, Slauch JM. 2021. Iron transport and metabolism in Escherichia, Shigella, and Salmonella. EcoSal Plus 9:eESP-0034-2020. https://doi.org/10.1128/ecosalplus.ESP-0034-2020.
3. Reddy PV, Puri RV, Chaupan P, Kar R, Rohilla A, Khera A, Tyagi AK. 2013. Disruption of mycobactin biosynthesis leads to attenuation of Mycobacterium tuberculosis for growth and virulence. J Infect Dis 208:1255–1265. https://doi.org/10.1093/infdis/jit250.
4. Wells RM, Jones CM, Xi Z, Speer A, Danilchanka O, Doornbos KS, Sun P, Wu F, Tian C, Niederweis M. 2013. Discovery of a siderophore export system essential for virulence of Mycobacterium tuberculosis. PLoS Pathog 9:e1003120. https://doi.org/10.1371/journal.ppat.1003120.
5. Lui S, Guo H, Adamson J, Clas JD, Davis TD, Chavadi SS, Warren JD, Quadri LEN, Tan DS, Bishai WR. 2013. Pharmacokinetic and in vivo efficacy studies of the mycobactin biosynthesis inhibitor salicyl-AMS in mice. Antimicrob Agents Chemother 57:5138–5140. https://doi.org/10.1128/AAC.00918-13.
6. Lamb AL. 2015. Breaking a pathogen’s iron will: inhibiting siderophore production as an antimicrobial strategy. Biochim Biophys Acta 1854:1054–1070. https://doi.org/10.1016/j.bbabap.2015.05.001.
7. Koh E-I, Henderson JP. 2015. Microbial copper-binding siderophores at the host-pathogen interface. J Biol Chem 290:18967–18974. https://doi.org/10.1074/jbc.R115.644329.
8. Calmettes C, Ing C, Buckwalter CM, El Bakkouri M, Chieh-Lin Lai C, Pogoutse A, Gray-Owen SD, Pomes R, Moraes TF. 2015. The molecular mechanism of zinc acquisition by the neisserial outer-membrane transporter ZnuD. Nat Commun 6:9796. https://doi.org/10.1038/ncomms9796.
9. Jean S, Juneau RA, Criss AK, Cornellson CN. 2016. Neisseria gonorrhoeae evade calprotectin-mediated nutritional immunity and survives neutrophil extracellular traps by production of TdfH. Infect Immun 84:2982–2994. https://doi.org/10.1128/IAI.00319-16.
10. Maurakis S, Keller K, Maxwell CN, Pereira K, Chazin WJ, Criss AK, Cornellson CN. 2019. The novel interaction between Neisseria gonorrhoeae TdfJ and human S100A7 allows gonococci to subvert host zinc restriction. PLoS Pathog 15:e1007937. https://doi.org/10.1371/journal.ppat.1007937.
11. Stork M, Grijpstra J, Bos MP, Manas Torres C, Devos N, Poolman JT, Chazin WJ, Tommassen J. 2013. Zinc piracy as a mechanism of Neisseria meningitidis for evasion of nutritional immunity. PLoS Pathog 9:e1003733. https://doi.org/10.1371/journal.ppat.1003733.
12. Chaturvedi KS, Hung CS, Crowley JR, Stapleton AE, Henderson JP. 2012. The siderophore yersiniabactin binds copper to protect pathogens during infection. Nat Chem Biol 8:731–736. https://doi.org/10.1038/nchembio.1020.
13. Koh E-I, Hung CS, Parker KS, Crowley JR, Giblin DE, Henderson JP. 2015. Metal selectivity by the virulence-associated yersiniabactin metallophore system. Metallomics 7:1011–1022. https://doi.org/10.1039/c4mt00341a.
14. Behnlsen J, Zhi H, Aron AT, Subramanian V, Santus W, Lee MH, Gerner RR, Petras D, Liu JZ, Green RD, Price SL, Camacho J, Hillman H, Tjokrosurjo J, Montaldo NP, Hoover EM, Treacy-Abarca S, Gliston BA, Skaar EP, Criss WJ, Gameau-Tsodikova S, Lawrenz MB, Perry RD, Nuccio SP, Dorrestein PC, Raffatellu M. 2021. Siderophore-mediated zinc acquisition enhances enterobacterial colonization of the inflamed gut. Nat Commun 12:7016. https://doi.org/10.1038/s41467-021-27297-2.
15. Katumba GL, Tran H, Henderson JP. 2002. The Yersinia high-pathogenicity island encodes a siderophore-dependent copper response system in uropathogenic Escherichia coli. mBio 3:e00291-21. https://doi.org/10.1128/mBio.00291-21.
16. Bobrov AG, Kirillina O, Fetherston JD, Miller MC, Burlinson JA, Perry RD. 2014. The Yersinia pestis siderophore, yersiniabactin, and the ZnuABC system both contribute to zinc acquisition and the development of lethal septicaemic plague in mice. Mol Microbiol 93:759–775. https://doi.org/10.1111/mmi.12693.
17. Price SL, Vadyvaloo V, DeMarco JK, Brady A, Gray PA, Kehl-Fie TE, Gameau-Tsodikova S, Perry RD, Lawrenz MB. 2021. Yersiniabactin contributes to overcoming zinc restriction during Yersinia pestis infection of mammalian and insect hosts. Proc Natl Acad Sci U S A 118:2104073118. https://doi.org/10.1073/pnas.2104073118.

Botella H, Peyron P, Levillain F, Poincloux R, Poquet Y, Brandli I, Wang C, Tailleux L, Tilleul S, Garneau-Tsodikova S, Perry RD, Lawrenz MB, Khera A, Gao Q, Maridonneau-Parini I, Butler PD, Castagnoli PR, Gicquel B, de
32. Xiao Z, Neyroles O. 2011. Mycobacterial pI(-)-type ATPases mediate resistance to zinc poisoning in human macrophages. Cell Host Microbe 10:248–259. https://doi.org/10.1016/j.chom.2011.08.006.

35. Liu Z, Loegner TR, Wang F, Sacchettini JC. 2013. Structures of Mycobacterium tuberculosis FadD10 protein reveal a new type of adenylate-forming enzyme. J Biol Chem 288:18473–18483. https://doi.org/10.1074/jbc.M113.466912.

36. Gotzke H, Kilisch M, Martinez-Carranza M, Sogne-Idrissi S, Rajavel A, Schlichtzaehle T, Engels J, Jungmann R, Stenmark P, Opazo F, Frey S. 2019. The ALFA-tag is a highly versatile tool for nanobody-based bioscience applications. https://doi.org/10.1038/s41467-019-12301-7.

37. Wang Q, Boshoff HM, Harrison JR, Ray PC, Green SR, Wyatt PG, Barry CE. 2020. PE/PPE proteins mediate nutrient transport across the outer membrane of Mycobacterium tuberculosis. Science 367:1147–1151. https://doi.org/10.1126/science.aav5912.

38. Mitra A, Speer A, Lin K, Ehr S, Niederweis M, Stallings CL. 2017. PPE surface proteins are required for heme utilization by Mycobacterium tuberculosis. mBio 8:e01720-16. https://doi.org/10.1128/mBio.01720-16.

39. Dechow SJ, Baker JJ, Murro MR, Abramovitch RB. 2021. pep5I variants promote non-replicating Mycobacterium tuberculosis to grow at acidic pH by selectively promoting glycerol uptake. bioRxiv https://doi.org/10.1101/2020.05.19.044480.

44. Liu T, Ramesh A, Ma Z, Wang SK, Zhang L, Green GN, Talaat AM, Sacchettini JC, Giedroc DP. 2007. CsrR is a novel Mycobacterium tuberculosis copper-sensing transcriptional regulator. Nat Chem Biol 3:60–68. https://doi.org/10.1038/nchembio844.

45. Ward SK, Abomoeak B, Hoye EA, Steinberg H, Talaat AM. 2010. CtpP: a putative copper exporter required for full virulence of Mycobacterium tuberculosis. Mol Microbiol 77:1096–1110. https://doi.org/10.1111/j.1365-2958.2010.07273.x.

46. Ward SK, Hoye EA, Talaat AM. 2008. The global responses of Mycobacterium tuberculosis to physiological levels of copper. J Bacteriol 190:2939–2946. https://doi.org/10.1128/JB.000309.

47. Li CX, Gleason JE, Zhang SX, Bruno VM, Cormack BP, Culotta VC. 2015. Metalloprotease and other cytoplasmic metalloenzymes. Curr Opin Biotechnol 50:349–357. https://doi.org/10.1016/j.copbio.2016.07.016.

48. Hadley RC, Gu Y, Nolan EM. 2018. Initial biochemical and functional evaluation of Mycobacterium avium, Mycobacterium tuberculosis, and Mycobacterium smegmatis putative copper exporters required for full virulence of Mycobacterium tuberculosis. Mol Microbiol 100:1145–1151. https://doi.org/10.1111/mmi.13961.

49. Wagner D, Maser J, Lai B, Cai Z, Barry CE, Hoener zu Bentrup D, DG R, Bermudez LE. 2005. Elemental analysis of Mycobacterium avium, Mycobacterium tuberculosis-, and Mycobacterium smegmatis-containing phagosomes indicates pathogen-induced microenvironments within the host cell’s endosomal system. J Immunol 174:1491–1500. https://doi.org/10.4049/jimmunol.174.1.1491.

50. Grosse-Siestrup BT, Gupta T, Helms S, Tucker SL, Voskul MI, Quinn FD, Karls RK. 2020. A role for Mycobacterium tuberculosis sigma factor C in copper nutritional immunity. Int J Mol Sci 21:3007. https://doi.org/10.3390/ijms21103007.

51. Obisesan AO, Zygiel EM, Nolen EM. 2021. Bacterial responses to iron withholding by calprotectin. Biochemistry 50:3337–3346. https://doi.org/10.1021/acs.biochem.0c00309.

52. Bardarov S, Bardarov S, Pavelka MS, Sambandamurthy V, Larsen M, Tufarelli J, Chan J, Hartful G, Jacobs WR. 2002. Specialized transduction: an efficient method for generating marked and unmarked targeted gene disruptions in Mycobacterium tuberculosis, M. bovis BCG and M. smegmatis. Microbiology 148:3007–3017. https://doi.org/10.1099/00221287-148-10-3007.