Iron Acquisition in *Mycobacterium tuberculosis*

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ABSTRACT: The highly contagious disease tuberculosis (TB) is caused by the bacterium *Mycobacterium tuberculosis* (Mtb), which has been evolving drug resistance at an alarming rate. Like all human pathogens, Mtb requires iron for growth and virulence. Consequently, Mtb iron transport is an emerging drug target. However, the development of anti-TB drugs aimed at these metabolic pathways has been restricted by the dearth of information on Mtb iron acquisition. In this Review, we describe the multiple strategies utilized by Mtb to acquire ferric iron and heme iron. Mtb iron uptake is a complex process, requiring biosynthesis and subsequent export of Mtb siderophores, followed by ferric iron scavenging and ferric-siderophore import into Mtb. Additionally, Mtb possesses two possible heme uptake pathways and an Mtb-specific mechanism of heme degradation that yields iron and novel heme-degradation products. We conclude with perspectives for potential therapeutics that could directly target Mtb heme and iron uptake machineries. We also highlight how hijacking Mtb heme and iron acquisition pathways for drug import may facilitate drug transport through the notoriously impregnable Mtb cell wall.
5.3. IsdG-Type Heme Oxygenases
5.3.1. Overview
5.3.2. Structure and Mechanism of IsdG-Type Proteins
5.3.3. Heme Ruffling That Is Required for Heme Activation in IsdG-Type Heme Degraders
5.3.4. IsdG-Type Proteins Require an Electron Donor
5.4. Mtb MhuD Heme Degradation
5.4.1. Overview
5.4.2. Structure of MhuD
5.4.3. Mechanism of MhuD Heme Degradation
5.4.4. Unique Diheme Form of MhuD
5.4.5. Potential Mtb MhuD Electron Donors
6. Iron-Acquisition Pathways As Anti-TB Drug Targets
6.1. Targeting the Mycobactin Biosynthetic Pathway
6.2. Targeting Iron Uptake Pathways
6.3. Targeting Heme Uptake Pathways
7. Conclusions
Author Information
Corresponding Authors
ORCID
Notes
Biographies
Acknowledgments
Abbreviations
References

1. INTRODUCTION
1.1. Tuberculosis
Tuberculosis (TB) is one of the deadliest infectious diseases known to man. TB has plagued the world since ancient times and has even been discovered in ancient Egyptian mummies.1 Prior to the advent of anti-TB drugs in the 1940s, a TB diagnosis was considered a death sentence.2 Mycobacterium tuberculosis (Mtb), the etiological agent of TB, is a slow-growing, acid-fast bacterium.3 Mtb has evolved multiple mechanisms to evade host immune defenses, and upon host assault and antibiotic treatment, Mtb often moves into a latent state within the human host.3 Notably, latent TB is asymptomatic but can be activated by malnutrition or in immunocompromised individuals.4 Before the discovery of anti-TB drugs, TB patients were sent to isolated resorts to be quarantined and treated, which at the time consisted of little more than bed rest.5 Following the discovery of penicillin, the first anti-TB drug streptomycin was discovered in 1943 from Streptomyces griseus.6 Other anti-TB drugs soon followed including isoniazid, rifampicin, pyrazinamide, and ethambutol, and by the end of the 1970s, it was thought that TB could be eradicated globally.5 Unfortunately, this was not to be the case. Today, TB remains one of the most deadly infectious diseases worldwide.

1.2. Need for Novel Anti-TB Drugs
With the onset of AIDS in the 1980s, TB cases started to reemerge.7 HIV and TB are synergistic. AIDS weakens the immune system, allowing for latent TB to be reactivated into a full-blown active infection. An estimated one-third of the world’s population is infected with latent TB, and those living with HIV are 20–30 times more likely to develop active TB than those without HIV.6 In 2016, 10.4 million people were diagnosed and treated for TB, and there were 1.7 million TB-related deaths. Notably 95% of these deaths were in low- or medium-income countries.6

Following the discovery of streptomycin in 1943, an array of anti-TB drugs was developed over the next several decades. First-line drugs such as isoniazid, rifampicin, pyrazinamide, ethambutol, and streptomycin are used in the standard cocktail to treat TB patients. These drugs are administered together as a drug regimen cocktail in accordance with guidelines set forth by the World Health Organization (WHO).5 The usual TB treatment regimen lasts more than 6 months and up to 18 months in the case of drug-resistant isolates.5 Unfortunately, many anti-TB drugs cause harsh side effects or result in medical complications such as liver damage. Moreover, the prolonged period of treatment can result in therapeutic noncompliance by patients that leads to the development of drug-resistant strains.6

The TB epidemic is compounded by the rise of multidrug-resistant TB (MDR-TB).7 Over the last two decades, the number of cases of MDR-TB resistant to anti-TB drugs isoniazid and rifampicin has been steadily increasing, with WHO currently estimating 160 000 new MDR-TB cases per year. Even more concerning, however, is the emergence of extensively drug-resistant TB (XDR-TB), which, like MDR-TB, is resistant to isoniazid and rifampin but has additional resistance to fluoroquinolone and second-generation anti-TB drugs (such as amikacin, kanamycin, or capreomycin). To date, only 30% of XDR-TB cases are successfully treated. The MDR-TB and XDR-TB burden largely falls in India, China, and the Russian Federation.8 With the emergence and proliferation of MDR-TB and XDR-TB, the development of new classes of anti-TB therapeutics is critical. Drug discovery efforts are ongoing; however, only one new anti-TB drug, the highly toxic bedaquiline, has been approved in the last several decades. Due to its particularly harsh side effects, bedaquiline is only used to treat the most severe cases of MDR-TB and XDR-TB.8

1.3. Mtb Iron Assimilation
Because of its role in many essential biological processes, iron is essential for the survival of all living organisms including bacterial pathogens.9–11 To acquire iron, bacteria have evolved pathways to take up both host heme and nonheme iron (simply referred to as iron henceforth). As iron and/or heme acquisition is essential, the mechanisms of pathogen iron and heme uptake are thought to represent potential drug targets.

In this Review, we will focus on the Mtb siderophore-dependent iron and heme uptake pathways. We will start by describing sources of host iron that Mtb is likely to encounter during active infection. As Mtb is neither a Gram-negative nor Gram-positive bacterium, we will briefly discuss briefly bacterial siderophore-dependent iron and heme uptake mechanisms. Then, we will review the Mtb iron and heme uptake pathways in detail. We will include structural and functional analyses and discuss the relationship between Mtb iron and heme uptake pathways with those of Gram-negative and Gram-positive organisms. The topics of siderophore biosynthesis and regulation of siderophore-dependent iron and heme uptake pathways will be reviewed, albeit only briefly. While it has been suggested that Mtb may internalize holo-iron-binding proteins, transferrin (Tf) and lactoferrin (Lf),12,13
the molecular mechanism of internalization of holo-Tf and holo-Lf proteins is unknown and will not be discussed further. Finally, we will present ideas on how to exploit Mtb heme and iron uptake systems as a target for novel anti-Mtb drugs.

2. Overview of Iron in the Host

2.1. Role of Iron in Life

Metals are an integral part of life. Within the human body, the most abundant metal ions are Na+, K+, Mg2+, and Ca2+, while the most abundant transition metal is iron. Iron performs a multitude of extremely diverse roles in biology, with well-known roles in biological catalysis,14,15 electron transport,16 and oxygen transport and storage.17

During pathogenesis, the limited availability of free iron and heme iron in the host represents a serious challenge to the pathogen. Although iron is a useful catalyst when coordinated to a protein, free iron has toxic effects. In an aqueous environment, uncomplexed iron, predominately ferrous iron (Fe2+), is a potent activator of free-radical chemistry through the Fenton reaction, catalyzing the formation of highly reactive hydroperoxy and hydroxy radicals.18 Further, Fe2+ readily oxidizes to its ferric form (Fe3+), which has extremely low solubility (10^{-18} M) and results in ferric iron precipitate.19 As a consequence of iron’s reactivity and potential insolubility, mammals maintain very low concentrations of free, unbound iron. Indeed, the majority of iron is tightly coordinated to high-affinity iron transport, storage, and scavenging proteins.20 Furthermore, iron sequestration is a key component of the mammalian innate immune system, and humans have dedicated proteins whose sole role is to scavenge extracellular iron to prevent its uptake by pathogens.21,22 The battle for iron between the human host and bacteria has evolved into a sophisticated cat and mouse game, where bacteria have developed complex pathways to steal iron from host proteins, and to circumvent this, humans have evolved proteins to neutralize bacterial iron scavenging pathways.23,24

2.2. Sources of Host Iron

The human body contains ~5 g of iron, which is divided into ferric (Fe^{3+}) and ferrous (Fe^{2+}) iron ions, and heme iron, where iron is coordinated to protoporphyrin IX (PPIX). The

Table 1. Heme-Binding Properties of Host Heme Sources

<table>
<thead>
<tr>
<th>protein</th>
<th>coordination</th>
<th>(k_{on}) ((\mu M^{-1} s^{-1}))</th>
<th>(k_{off}) (s^{-1})</th>
<th>(K_a) (M^{-1})</th>
<th>ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>metHb α-chain (pH 7)</td>
<td>6c His/H2O</td>
<td>100</td>
<td>1.6 \times 10^{-4}</td>
<td>6.0 \times 10^{11}</td>
<td>27, 28</td>
</tr>
<tr>
<td>metHb β-chain (pH 7)</td>
<td>6c His/H2O</td>
<td>100</td>
<td>4.2 \times 10^{-3}</td>
<td>2.4 \times 10^{10}</td>
<td>27, 28</td>
</tr>
<tr>
<td>serum albumin</td>
<td>5c Tyr</td>
<td>~50</td>
<td>1.1 \times 10^{-2}</td>
<td>4 \times 10^{5}</td>
<td>28</td>
</tr>
<tr>
<td>hemopexin</td>
<td>6c His/His</td>
<td>ND</td>
<td>ND</td>
<td>10^6 to 1.9 \times 10^{14}</td>
<td>29, 30</td>
</tr>
</tbody>
</table>

The rate \(k_{on}\) refers to the bimolecular rate constant of heme binding to apo-protein in dilute conditions. The rate \(k_{off}\) represents the unimolecular rate of heme dissociation. The relationship between \(k_{on}\), \(k_{off}\), and the association constants \(K_a\) is \(K_a = k_{on}/k_{off}\). This is the same for Tables 2 and 3. ND = not determined, 5c = 5-coordinate, and 6c = 6-coordinate.

Figure 1. Iron availability in macrophages. Macrophages uptake iron by sequestering Tf/Lf bound iron and storing released iron in ferritin nanocages. Heme is uptaken either by Hx and Hp-Hb complex receptors where the proteins are degraded in the endosome or by erythrophagocytosis of senescent red blood cells (RBCs). In both cases, heme is released into the macrophage cytosol. There, heme is degraded by heme oxygenase (HO-1) to release iron, carbon monoxide (CO), and biliverdin—discussed in section 5.1. Mtb residing in the macrophage phagosome will encounter iron and likely also heme and Hb.
majority of human nonheme iron is circulating in the bloodstream, where most of circulating Fe\(^{3+}\) ions are complexed by transferrin (Tf) and lactoferrin (Lf). Tf coordinates two Fe\(^{3+}\) ions tightly (\(K_a = 1 \times 10^{-23}\) M\(^{-1}\)) and shuttles iron between tissues.\(^{25}\) Lf is a glycosylated Fe\(^{3+}\) ion transporter that is part of the innate immune system and is predominantly found in secretory fluids.\(^{21}\) Importantly, Tf serves a key antimicrobial role by establishing iron-replete conditions.\(^{21}\) Noncirculating iron is found within tissues, where iron is coordinated to proteins such as iron-containing enzymes and regulatory proteins, and ferrous iron is stored in large ferritin nanocages.\(^{26}\)

### 2.3. Sources of Host Heme

Heme iron represents the most abundant form of circulating iron in the human body. Similar to free iron, heme is highly reactive. Therefore, the concentration of free heme is kept extremely low and heme is tightly bound to proteins (\(K_a = 10^8-10^{13}\) M\(^{-1}\)).\(^{28,31}\) Approximately 80% of all human iron is incorporated into the heme-containing oxygen carrier hemoglobin (Hb) in erythrocytes,\(^{32,33}\) making Hb the greatest human reservoir of any type of iron.\(^{34}\) Hb exists in two oxidation states with very different biophysical properties. Reduced Hb is an \(\alpha_2/\beta_2\) tetramer that transports O\(_2\) and binds ferrous heme very tightly. In contrast, oxidized Hb (metHb) breaks down into \(\alpha/\beta\) heterodimers and no longer binds O\(_2\).\(^{27,28}\) The binding affinity of metHb for ferric heme is considerably diminished, where the half-life of heme dissociation is \(\sim 2-3\) min for the \(\beta\) subunit of metHb (Table 1).\(^{27}\) Thus, for heme bound to metHb, heme dissociates passively on a time scale that is rapid enough to create a pool of labile heme that can be utilized by bacterial heme uptake systems. In a healthy human, a mere 1–2% of Hb is in the metHb form. However, the availability of metHb may rise during bacterial infection as many pathogens secrete hemolysins that lyse erythrocyte cells, resulting in Hb release into the blood serum.\(^{35}\) Hb released under hemolytic conditions is subject to rapid oxidation, and thus heme is more susceptible to dissociation and acquisition by a pathogen.\(^{27}\)

To prevent heme loss to pathogens, mammals have evolved mechanisms to recover heme dissociated from metHb. There are three primary proteins involved in metHb and free heme recovery: haptoglobin (Hp),\(^{22,36}\) hemopexin (Hx),\(^{37,38}\) and serum albumin.\(^{39}\) Hp is a tetrameric serum glycoprotein that forms a tight interaction with metHb to prevent heme dissociation.\(^{22,37}\) Hx is a monomeric glycoprotein that binds free heme with an extremely high affinity (Table 1). While serum albumin binds heme with relatively low affinity (\(K_a \approx 10 \times 10^6\) M\(^{-1}\)),\(^{39,28}\) due to its overall high abundance, it also plays an important role in reducing the free heme concentration in serum by binding free heme and transferring it to Hx.

### 2.4. Potential Sources of Host Iron for Mtb

TB is contracted via inhalation of Mtb into the lungs. Mtb is primarily an intracellular organism, residing in the phagosome of lung macrophages during initial infection (Figure 1). Additionally, TB has an asymptomatic, latent stage, where Mtb can remain dormant in granulomas for decades.\(^{4}\) During active infection, Mtb will extracellularly disseminate via the bloodstream from its primary infection site to secondary ones, such as the central nervous and lymphatic systems. In the macrophage phagosome, Mtb will encounter extremely low free iron levels,\(^{40}\) even though the macrophage cytosol contains relatively high iron levels (Figure 1).\(^{33}\) To overcome this, it has been suggested that Mtb siderophores promote iron diffusion across the phagosomal membrane.\(^{41}\) This hypothesis
is supported by $^{59}$Fe labeling experiments that demonstrate iron accumulation in Mtb residing in phagosomes.\textsuperscript{42,43} Moreover, macrophages degrade senescent red blood cells in the phagolysosome,\textsuperscript{44} making Hb a promising source of heme iron for macrophage-residing Mtb (Figure 1). Indeed, a recent study has established that the cytosolic concentration of the bioavailable heme pool within human lung cells is $400-600$ nM.\textsuperscript{45}

When Mtb is extracellular, such as in the bloodstream, Mtb may sequester iron from Tf, Lf, and ferritin. Extracellular Mtb will also encounter heme. Mtb possesses hemolysins that lyse erythrocyte cells,\textsuperscript{35} thereby releasing HB into the bloodstream as metHb and creating a pool of labile heme as well as heme that is weakly bound to serum albumin. Notably, in severe Mtb infection blood accumulates in the lung cavities, possibly resulting in elevated heme levels in the lungs.

3. BACTERIAL SIDEROPHORE-DEPENDENT IRON UPTAKE PATHWAYS

3.1. Siderophore-Dependent Iron Uptake

Siderophore-mediated iron uptake is part of the iron uptake repertoire in both Gram-negative and Gram-positive bacteria. Siderophores are iron-chelating small molecules ($M_w$ ranging from 100 to 2000 Da) secreted by microorganisms to facilitate iron scavenging, transport, and uptake.\textsuperscript{46–48} The defining feature of siderophores is their extremely high affinity for Fe$^{3+}$ ions with $K_a$ values up to $10^{52}$ M$^{-1}$.\textsuperscript{49} Siderophores are
opportunistic iron scavengers that are not known to display a preference regarding their iron source, as they remove iron from Tf, Lf, ferritin, and other human iron binding proteins. Once bound to ferric iron, ferric-siderophores are imported into bacteria by a variety of mechanisms (see section 3.4).

3.2. Siderophore Structure

The majority of siderophores form hexadentate, octahedral complexes with ferric iron. Siderophores can be classified into four major categories based on the ferric iron-binding moiety: carboxylates, catecholates, hydroxamates, and phenolates (Figure 2A), all of which are hard Lewis bases. Moreover, some siderophores contain multiple binding moieties.48

Pathogenic Mtb produces two siderophores termed “mycobactins”, mycobactin and carboxymycobactin (Figures 2B and 3), which are mixed-type siderophores with both phenolate and hydroxamate moieties (Figure 2).46 While the two siderophores share an identical core, consisting of five amino acids with a characteristic oxazolidine ring derived from salicylate (Figure 3),46,50 they differ in hydrophobicity and cellular localization. Owing to its long aliphatic tail, mycobactin is hydrophobic and cell-wall- and membrane-associated. In contrast, the secreted carboxymycobactin features a shorter tail that terminates with a carboxylate group, rendering it hydrophilic (Figure 3).51,52

Other mycobacterial species produce a variety of mycobactins, which all retain the same structural core (Figure 3) but are derivatized differently. While Mtb has been shown to produce mycobactin T (Figure 3A), Mycobacterium smegmatis and Mycobacterium paratuberculosis produce their own specific siderophores, mycobactin S and mycobactin J, respectively.46,47 Nonpathogenic, saprophytic mycobacteria also produce a different extracellular siderophore that does not have the mycobactin structural core, a formylated pentapeptide called exochelin;53 however, in this Review, we shall only discuss Mtb mycobactins (see Figure 3).

3.3. Siderophore Biosynthesis

3.3.1. Overview. Biosynthetic pathways of siderophore production can be classified into two types: nonribosomal peptide synthetase (NRPS)-dependent54 and NRPS-independent55 siderophore biosynthetic pathways.36,48,56 NRPSs are large multi-enzyme complexes that synthesize small cyclic or linear polypeptide products without requiring an mRNA template and usually utilize a thiotemplate mechanism for biosynthesis. NRPSs have been proposed to be responsible for the synthesis of many aryl-capped (or aromatic-ring-capped) siderophores such as Mtb mycobactin (Figure 3).48

3.3.2. Mtb Mycobactin Biosynthesis. Biosynthesis of mycobactin occurs in an NRPS-dependent manner. As the biochemistry and structures within this pathway have been recently reviewed,57,58 we will only present a brief summary of the pathway. In Mtb, mycobactin biosynthetic proteins are encoded by two gene clusters, mbt-1 and mbt-2, where each locus includes genes mbtA to mbtJ and mbtK to mbtN, respectively (Figure 4A).59,60 The proteins encoded by mbt-1 assemble the core of the mycobactin molecule, whereas those encoded by mbt-2 assemble the aliphatic hydrophobic side chain of mycobactin T.

In the mycobactin biosynthetic pathway (Figure 4B), several enzymes produce and activate precursor components. MbtI catalyzes the conversion of chorismate (1) to salicylic acid (3). Then MbtA activates salicylic acid and catalyzes its transfer to the thiolate domain of MbtB through an acyl adenylate intermediate (Figure 4B (4)). MbtB is part of the megasynthase composed of three NRPSs (MbtB, MbtE, and MbtF) and two polyketide synthases (MbtC and MbtD), which then assembles the core mycobactin structure (Figure 3, light blue).51 Not part of the megasynthase are MbtG, which
catalyzes the hydroxylation of L-lysine to N6-hydroxy-L-lysine, and MbtH, which chaperones the correct folding of the NRPSs (MbtB, MbtE, and MbtF). MbtJ is the only protein with an unknown function encoded by the mbt-1 locus. The proteins encoded by mbt-2 produce the functional mycobactin T (Figure 4B). First, the aliphatic, hydrophobic acyl chain is assembled by the enzymes MtbL, MbtM, and MbtN. Second, MbtK catalyzes the linkage of the acyl chain to the mycobactin core.

3.3.3. Regulation of Mycobactin Biosynthesis. Mycobactin synthesis is regulated by the iron-dependent regulator (IdeR) (Figure 4A), which has sequence similarity to the iron-dependent diphtheria toxin repressor from Corynebacterium diphtheria. IdeR represses genes associated with mycobactin synthesis in the presence of iron, while acting as a positive regulator for the Mtb iron-storage ferritin proteins BfrA and BfrB. IdeR also regulates many genes that encode proteins involved in Mtb mycobactin export and import (see sections 3.5 and 3.6). Importantly, an Mtb ideR deletion mutant does not survive in macrophages or a mouse model, suggesting that ideR is an essential gene for Mtb virulence.

3.4. Bacterial Siderophore Export and Import

Siderophore-mediated iron uptake has been extensively studied in E. coli and therefore warrants discussion to illustrate how bacteria may acquire iron and shuttle it across the outer and inner membranes to the cytoplasm. E. coli enterobactin (Figure 2B) is synthesized in the cytoplasm and is then translocated into the periplasm by a transmembrane protein, which is a member of the ATPase major facilitator superfamily (MFS). The multifunctional efflux channel TolC then exports enterobactin from the periplasm into the extracellular space. TolC-mediated export of enterobactin also requires a periplasmic membrane fusion protein and resistance nodulation cell division (RND) transporters (Figure 5).

Upon scavenging extracellular ferric iron, E. coli ferric-enterobactin (FeEnt) is imported into the periplasm utilizing a specific β-barrel outer membrane receptor (OMR) through a TonB-dependent transport mechanism (Figure 5). In the periplasm, a periplasmic binding protein (PBP) binds and delivers FeEnt to the inner membrane ATP-binding cassette (ABC) transporter complex, so it can be translocated into the cytoplasm where the siderophore is modified by FeEnt esterase to promote the release of iron (Figure 5). Moreover, E. coli has also evolved to import ferric siderophores from other bacteria using specific siderophore-binding TonB-dependent receptors, such as ferrichrome from Aspergillus. Once in the cytoplasm, an E. coli ferric-siderophore reductase induces iron release by reducing ferric iron to ferrous iron, resulting in iron dissociation.

In Gram-positive bacteria, siderophore export and import is simplified because only one membrane needs to be traversed. However, these export and import pathways have been relatively understudied. While it is known that ferric-siderophores are recognized by siderophore-binding proteins (homologous to PBPs) located in the cell membrane and that a permase assists in the translocation of the ferric-siderophore across the membrane, the molecular mecha-
nisms of siderophore transport through the Gram-positive cell-wall are poorly understood.

3.5. Mtb Siderophore Export

3.5.1. Overview. The two Mtb siderophores, carboxymycobactin and mycobactin (Figure 3), are utilized differently. Carboxymycobactins are relatively soluble and are exported into the extracellular environment, where they scavenge iron from human iron-containing proteins such as Tf and Lf. Mycobactins, on the other hand, due to their insolubility, are anchored to the Mtb outer membrane and cell-wall environments. Export of mycobactin and carboxymycobactin across the inner membrane is dependent on MmpL (mycobacterial membrane protein large) proteins, MmpL4 and MmpL5, along with their small membrane-associated accessory proteins, MmpS (mycobacterial membrane protein small) proteins, MmpS4 and MmpS5 (see Figure 8). More recently, it was proposed that ESX-3, one of the five type VII secretion systems in Mtb, also contributes to iron acquisition as ESX-3 mutants display defects in ferric-carboxymycobactin uptake and accumulation of cell-wall-associated mycobactins (see Figure 8). However, the precise role ESX-3 plays in iron import/export is unknown and thus will not be further discussed in this Review.

3.5.2. Mtb MmpL Proteins. MmpL proteins appear to act in both siderophore export and heme import pathways. We will therefore present a broad description of this important class of Mtb proteins. MmpL proteins belong to the RND permease superfamily of transmembrane transporters, which associate with outer membrane factors. This assembly is stabilized by periplasmic membrane fusion proteins to form a three-component efflux pump (extensively reviewed in ref 87), which is driven by proton motive force. Mtb MmpL proteins are functionally important and have been shown to mediate the transport of a variety of substrates across the mycobacterial inner membrane, including Mtb lipids and siderophores. Mutational analyses revealed that one of the MmpL proteins, MmpL3, is essential for Mtb viability, and several other MmpL proteins are necessary for Mtb virulence in mice infections including MmpL4, MmpL5, MmpL7, MmpL8, MmpL10, and MmpL11.

The structures of canonical Gram-negative RND transporters are homotrimeric, where each monomer harbors 12 transmembrane helices (TM) with N-terminal and C-terminal periplasmic domains inserted between TM1 and TM2 and between TM7 and TM8, respectively. Each periplasmic domain comprises two structurally similar porter subdomains (N-terminal porter subdomains, PN1 and PN2, and C-terminal porter subdomains, PC1 and PC2, containing a βαββαβ motif) and a “middle” docking subdomain (DN or DC) (Figure 6B and C).

Mtb has 13 MmpL members of the RND family, all of which are actinobacteria-specific inner membrane proteins of ~1 000 residues. Mtb MmpL proteins structurally cluster into two distinct subfamilies of RND transporters (Figure 6A). MmpL cluster I proteins have an N-terminal domain similar to an RND porter subdomain and a C-terminal domain that resembles that of RND proteins, comprising two porter subdomains and a docking subdomain (Figure 6B and C). In contrast, MmpL cluster II proteins have minimal periplasmic architecture, whereby the N- and C-terminal domains each contain a single porter subdomain (Figure 6B and C). Unlike other RND or MmpL cluster I proteins, each MmpL cluster II member has a C-terminal cytoplasmic domain (Figure 6B and C). In addition, some of the cluster I MmpL members are encoded along with a small membrane-associated accessory protein, MmpS.

3.5.3. Structures of MmpL4/5 and MmpS4/5. Both proteins involved in siderophore export, MmpL4 and MmpL5, are members of MmpL cluster I (see section 3.5.2, Figure 6A). Thus, they are predicted to contain two periplasmic domains, a D1 porter subdomain and a larger D2 domain composed of two porter and one docking subdomains (Figure 6B and C). MmpL4 and MmpL5 associate with MmpS4 and MmpS5, respectively. The structure of the mature form of MmpS4 (without its N-terminal signal peptide or single transmembrane tether) was solved by NMR and revealed an immunoglobulin-like fold comprising a 3- and 4-stranded β-sheet along with a disordered C-terminal (Figure 7). The sequence identity between MmpS4 and MmpS5 is >50%, suggesting similar structural folds. It was demonstrated that MmpS4 interacts only with the D1 domain of MmpL4.

3.5.4. Mtb Siderophore Export Mechanism. MmpL 4/5 are required for mycobactin and carboxymycobactin export (see Figure 8). In addition, several experiments suggest the accessory proteins MmpS4 and MmpS5 are also necessary for mycobactin and carboxymycobactin secretion. First, it was demonstrated that an MtbΔmmpS4/5 double mutant is not viable under low-iron conditions, suggesting both proteins take part in iron acquisition. Interestingly, the growth of the single MtbΔmmpS4 and MtbΔmmpS5 mutants are not attenuated in low-iron conditions, suggesting MmpS4 and MmpS5 have redundant roles. While MmpS4 and MmpS5 are likely to be involved in ferric iron acquisition, they do not participate in heme uptake because the MtbΔmmpS4/5 double mutant grows in media supplemented with heme. Next, it was shown that MtbΔmmpS4/5 can take up exogenous iron-loaded mycobactin, indicating that the MmpL4/S4 and MmpL5/S5 systems are involved specifically in mycobactin export but are not required for mycobactin import. Biosynthesis of mycobactin and its secretion may occur in a coordinated manner because MmpS4/5 localizes to the inner membrane along with the mycobactin biosynthetic enzyme, MtbG. This led to speculation that the MmpL4/S4 and MmpL5/S5 systems form a multienzyme complex with MtbG to couple mycobactin biosynthesis and its export. Similar coordinated
3.6. Mtb Ferric-Siderophore Import

3.6.1. Overview. Iron assimilation is essential for Mtb growth and virulence. However, import mechanisms for ferric mycobactins are poorly defined, as several necessary steps required for the import of mycobactins are unknown. One aspect of importing mycobactin is that has been recently well-characterized is the role of IrtA/IrtB. IrtA/IrtB is a heterodimeric protein involved in both the import of ferric-carboxymycobactin into the Mtb cytoplasm and the reduction of iron in the ferric-carboxymycobactin complex to release iron into the cytoplasm.

An assessment of Mtb MmpL4 and MmpL5 deletion mutants within mice revealed that MmpL4 is required for full virulence, while MmpL5 is essential for Mtb survival in the lung. With respect to MmpS4 and MmpS5, the growth of the single MtbΔmmpS4 and MtbΔmmpS5 mutants were attenuated in mice lungs. More strikingly, for the MtbΔmmpS4/mmpS5 double mutant there was 100% survival over a 180-day period compared to those mice infected with wild-type Mtb that had a 0% survival rate in the same time period. This is the strongest attenuation observed in Mtb mutants lacking iron utilization genes, suggesting that MmpL4/S4 and MmpS4/S5 mycobactin export systems are required for Mtb virulence and represent promising drug targets.

3.6.2. Potential Protein Players in the Periplasm. The current consensus is that hydrophobic mycobactins are cell-surface receptors or cell-wall associated and are involved in transporting ferric iron from the extracellular environment through the cell-wall environment. In support of this hypothesis, experiments show that ferric-carboxymycobactin can transfer its ferric iron cargo to mycobactin. Next, two predicted ferric-siderophore PBPs, FecB and FecB2, may shuttle ferric-mycobactin or ferric-carboxymycobactin from the Mtb cell-wall environment to the periplasmic side of the IrtA/IrtB transporter complex in the inner membrane (Figure 8).

FecB is predicted to be part of the Mtb iron-regulated IdeR regulon. A study in Mycobacterium avium shows that M. avium FecB is upregulated under low-iron conditions, suggesting this protein plays a role in iron acquisition.

Mtb FecB and FecB2 are predicted to belong to the Type III PBP superfamily. Mtb FecB has moderate sequence identity to other ferric-siderophore PBPs such as E. coli FhuD (Fe\(^{3+}\) enterobactin) and Pseudomonas aeruginosa FepB (Fe\(^{3+}\)-enterobactin catechol). The structure of Mtb FecB has been solved (PDB: 4PM4, Figure 9A) and is composed of N- and C-terminal lobes consisting of mixed α/β structures, where the two lobes are linked by a rigid “backbone” helix, characteristic of PBPs. Indeed, Mtb FecB2 has high structural homology to other ferric-siderophore and heme PBPs (such as IsdE and PhuT). Mtb FecB2 has the highest structural similarity with Staphylococcus aureus HtsA, which binds a hydroxamate-type siderophore, staphyloferrin A. Within the HtsA-ferric-staphyloferrin A complex, iron is not

![Figure 8: Mtb mycobactin-dependent iron uptake. Green arrows are carboxymycobactins, and the ones with acyl tails are mycobactins. MmpL4/S4 and MmpL5/S5 systems export mycobactin. Carboxymycobactins scavenge iron from host iron-binding proteins and then putatively transfer iron to cell-wall-associated mycobactins. PBPs proteins, FecB and FecB2, are proposed to shuttle iron to the periplasmic side of the inner membrane, where the IrtA/IrtB complex shuttles ferric-carboxymycobactin across the inner membrane. Iron is released into the cytoplasm by ferric reductases, IrtA-NTD and ViuB, upon which carboxymycobactin is recycled. ESX-3 is involved in mycobactin import—export by an unknown mechanism.](Image)
directly coordinated by HtsA residues (Figure 9C), and like the E. coli FhuD-gallichrome complex (Figure 9B), HtsA utilizes a combination of tyrosine and arginine residues to bind its siderophore noncovalently. The binding cleft of FecB2 contains two arginines (Arg184 and Arg221) and a tyrosine residue (Tyr39) that could be involved in stabilizing the mycobactin backbone and/or coordinating ferric iron. However, the structure of Mtb FecB2 featured an empty binding site that does not shed light on the protein’s potential cargo or mode of binding.

Mtb FecB and FecB2 have 24% sequence identity and may be functionally redundant. However, FecB2 has a predicted 35-residue signal peptide, whereas FecB is predicted to have a 68-residue N-terminal extension containing both a signal peptide and a membrane lipid attachment site, suggesting that FecB and FecB2 localize to different cellular locations and may work in tandem to shuttle ferric-carboxymycobactin and/or ferric-mycobactin through the lipid-rich cell-wall environment to the periplasmic side of the inner membrane. There are two potential mechanisms of PBP-dependent siderophore transport through the periplasm and/or the cell wall. They may use an affinity gradient manner to ensure directional transport similar to how S. aureus imports heme. Alternatively, a more complicated mechanism is possible where iron dissociates from ferric-mycobactin or ferric-carboxymycobactin within the periplasm as observed in P. aeruginosa siderophore iron transport.

3.6.3. Ferric-Carboxymycobactin Inner Membrane Permease—IrtA/IrtB. It has been shown that IrtA/IrtB is an inner membrane ABC heterodimeric transporter, which is critical for the import of Fe\(^{3+}\)-carboxymycobactin from the Mtb periplasm to the cytoplasm. We propose that ferric-carboxymycobactin bound to Mtb FecB or Mtb FecB2 directly transfers its ferric-carboxymycobactin cargo to the IrtA/IrtB complex for translocation to the cytoplasm. The genes that encode for IrtA/IrtB are co-ordinated and are downstream of a predicted IdeR-binding site. IrtA and IrtB are predicted to have six and five TMs, respectively, and both have C-terminal ATPase domains. IrtA and IrtB likely form a heterodimeric ABC transporter based on their similarity to the Y. pestis iron transporter, YbtPO. IrtA has an additional domain at its N-terminus, which is predicted to be a cytoplasmic ferric-siderophore reductase domain (IrtA-NTD) (discussed further in section 3.6.4).

Mutational analysis demonstrates that both IrtA and IrtB are required for fully functional Mtb iron import; however, the exact roles of IrtA and IrtB are still not fully resolved. Under iron-deplete conditions, inactivation of IrtB alone or both IrtA/IrtB led to significant growth defects. The effects were more severe with inactivated IrtA/IrtB than inactivated IrtB alone, suggesting that IrtA can form a homodimer that retains partial iron-import activity. As the carboxymycobactin concentration in the culture filtrate was measured to be equivalent in the MtbΔirtA/irtB mutant and wild-type strains, IrtA/IrtB does not act as a carboxymycobactin exporter. Further, as addition of ferric-carboxymycobactin as the sole iron source did not restore growth in the MtbΔirtA/irtB double mutant, it is clear that the IrtA/IrtB complex is involved in carboxymycobactin-dependent iron uptake. Notably, the MtbΔirtA/irtB mutant resulted in impaired replication in both macrophages and mouse lungs, suggesting that IrtA/IrtB is required for virulence.

In contrast, a study by Farhana et al. also suggests that M. smegmatis IrtA is involved in mycobactin export; however, compared to the study in Mtb, M. smegmatis IrtB alone is involved in ferric-mycobactin import. To resolve this apparent contradiction between studies, the precise function of IrtA and IrtB requires further investigation to determine whether these proteins form functional homodimers or a heterodimer.

3.6.4. Cytoplasmic Ferric-Carboxymycobactin Reduction. Once ferric-carboxymycobactins reach the Mtb cytoplasm, iron is released for utilization or storage. In some Gram-negative bacteria, the release of iron from ferric-siderophores is achieved through the flavin-dependent reduction of siderophore-bound iron by a ferric-siderophore reductase. As discussed earlier, the affinity of ferrous iron to the siderophore is substantially lower compared to that of ferric iron. Thus, iron reduction in the ferric-siderophore complex results in the release of ferrous iron. IrtA-NTD is predicted to be a ferric-siderophore reductase. Additionally, IrtA-NTD possesses a RXYS(T) motif, which has been associated with flavin adenine dinucleotide (FAD) binding in ferric oxidoreductases. Notably, unlike IrtA, IrtB lacks an analogous domain.

Results from in vitro binding experiments show that IrtA-NTD binds FAD in a 1:1 ratio. To show that FAD-dependent IrtA-NTD is required for fully functional IrtA/IrtB carboxymycobactin-dependent iron uptake, a mutational analysis of the reversed FAD-binding motif (Arg70, Tyr72, and Thr73) was carried out. The results demonstrated that mutating IrtA-NTD Tyr72 and Thr73 to Ala resulted in diminished growth under low-iron conditions, while the IrtA-NTD Arg70Ala mutation did not affect growth, suggesting that Tyr72 and Thr73 are critical for IrtA-NTD FAD binding and activity. Taken together the results suggest that FAD binding to IrtA-NTD is required for fully functional IrtA/IrtB, and furthermore, because IrtA-NTD is located on the cytoplasmic side of the inner membrane, IrtA-NTD is a cytoplasmic ferric-siderophore reductase.
In addition to IrtA-NTD, the Mtb proteome is predicted to encode another ferric-siderophore reductase, ViuB, (Figure 8), which is homologous to IrtA-NTD98 and a ferric-siderophore reductase ViuB from *Vibrio cholerea*.113 Mtb ViuB is induced under low-iron conditions110 and in bacillus Calmette-Guérin (BCG, an attenuated strain of *Mycobacterium bovis*).114 However, the Mtb ViuB knockout mutant showed no phenotype when grown in low-iron conditions or macrophages.115 Thus, further studies of Mtb ViuB are still required to determine if it plays a functional role in siderophore-mediated iron uptake.

4. BACTERIAL HEME UPTAKE PATHWAYS

4.1. Heme Uptake in Gram-Negative Bacteria

Heme uptake pathways have been discovered in a large number of Gram-negative bacteria and have been extensively studied in *P. aeruginosa*, *Serratia marcescens*, *Yersinia pestis*, and *Haemophilus influenzae*.116−122 Two general mechanisms of heme uptake are found in Gram-negative bacteria. Heme is acquired either through outer membrane receptors that bind host heme and/or heme proteins such as Hb or Hx (Figure 10A and B)118,120−122 or through the action of secreted heme-scavenging proteins called hemophores (Figure 10C).116,117,119 After heme has been acquired from the host, it is transported across the bacterial outer membrane using a TonB-dependent outer membrane receptor (Figure 10).118 Once in the periplasm, heme is transferred to PBPs, which shuttle heme to an inner membrane ABC transporter.118,123 In some species, multiple separate heme uptake pathways are present. In such cases, the individual heme uptake pathways are proposed to have complementary roles to enable heme scavenging from alternate heme sources or are differentially utilized depending on host heme availability.125,124

4.2. Heme Uptake in Gram-Positive Bacteria

Heme uptake pathways have also been discovered in several Gram-positive species, where those of *S. aureus*,125,126 *Streptococcus pyogenes*,127 and *Bacillus anthracis* have been studied extensively.128 These organisms use a sophisticated cascade of heme transport proteins to chaperone heme through the bacterium’s thick peptidoglycan cell wall and an ABC transporter to translocate heme across the inner membrane (Figure 11).

A characteristic feature in all Gram-positive heme uptake pathways studied to date is the presence of heme uptake proteins containing structurally conserved NEAT domains (Figure 12).130,131 While NEAT domains are not conserved at the amino acid level, they all share a common, superimposable immunoglobulin-like fold.132,133 NEAT domains serve many functional roles and can act as hemophores (e.g., *B. anthracis* IsdX1 and IsdX2),128,133,134 cell-surface receptors (e.g., *S. aureus* IsdB and IsdH),126,135,136 and cell-wall-anchored heme transporters (e.g., *S. aureus* IsdC and IsdA).104,137 Some NEAT domain proteins contain a single domain (e.g., IsdX1),128 while others are modular, with some featuring multiple NEAT domains in a single polypeptide chain, such as IsdX2.133 The functional properties of a NEAT domain are determined by the immediate heme coordination environment, secondary coordination sphere, and a 310-helix (Figure 12) that is located above the NEAT heme binding site and mediates protein−protein interactions.128 *S. aureus* heme extraction from metHb is achieved via direct protein−protein interactions between NEAT domains and metHb.151 For a comprehensive review on NEAT domains, we direct the reader to ref130.

4.3. Mtb Heme Uptake

4.3.1. Overview. Mtb heme uptake is distinct from other bacteria, lacking close homology to proteins associated with either Gram-negative or Gram-positive bacterial heme uptake systems, apart from homologues of PBPs that might play roles

Figure 10. Heme uptake mechanisms in Gram-negative bacteria. Heme is acquired using an outer membrane receptor specific for either (A) heme or a heme protein such as Hb, (B) hemopexin, or (C) a secreted hemophore. Heme is transferred across the outer membrane in a TonB-dependent manner, before being shuttled to the inner membrane by a PBP, where an ABC permease translocates heme into the cytoplasm. Heme is then degraded by a heme-degrading protein to release iron.
predicted small molecule PBP.146 pathway involves two surface-exposed proteins and one shown that the heme-analogue gallium-protoporphyrin IX region between genes Rv0202c and Rv0207c.90 The second mycobacterial proteins found in the Mtb nonoperonic genomic section 5.3.

Fi finally translocated across the membrane using an ABC transporter and degraded via the IsdG-type heme degraders, IsdG and IsdI; see section 5.3.

Figure 11. Heme uptake in Gram-positive S. aureus. Heme is acquired using cell-surface receptors that directly interact with and scavenge heme from Hb. Heme is then shuttled across the cell wall through a series of heme transporters via protein–protein interactions. Heme is finally translocated across the membrane using an ABC transporter and degraded via the IsdG-type heme degraders, IsdG and IsdI; see section 5.3.

Figure 12. Structure of the third NEAT domain of the S. aureus heme scavenger IsdH in its heme-bound form (PDB: 2Z6F).

In both siderophore-dependent iron and heme uptake pathways in Mtb; see Figure 13. The first comprises mycobacterial proteins found in the Mtb nonoperonic genomic region between genes Rv0202c and Rv0207c.90 The second pathway involves two surface-exposed proteins and one predicted small molecule PBPs.146

4.3.2. Evidence of Mycobacterial Heme Uptake. 4.3.2.1. Initial Evidence. Indirect evidence of an Mtb heme uptake pathway was first found in research unrelated to iron acquisition. It was observed that a recombinant BCG strain harboring a defective mycobactin biosynthetic pathway was able to slowly replicate in SCID mice, suggesting it uses an alternative iron source, likely heme iron.147 It has also been shown that the heme-analogue gallium-protoporphyrin IX (GaPPIX) is lethal to M. smegmatis. This implies that mycobacteria can acquire GaPPIX and process it in the cytoplasm to release toxic Ga metal, or that it is directly incorporated into heme proteins, whereby it disrupts essential pathways.148 This result further demonstrated that mycobacteria can take up metalloporphyrins (MPPs), such as heme.

4.3.2.2. Demonstration of a Heme Uptake Pathway. Heme uptake was directly examined using an Mtb strain (MtbΔmbtB) that has a disrupted mycobactin biosynthetic pathway. This mutant strain replicates poorly in media containing ferric iron as the sole iron source, indicating that mycobactin-mediated iron uptake is nonfunctional.90 In contrast, addition of heme or Hb to the media results in a restoration of growth, comparable to growth of MtbΔmbtB supplemented with exogenous mycobactin. These results indicate that Mtb has a heme uptake system.

4.4. First Heme Uptake Pathway

4.4.1. Identification of Protein Players. A single 10.6 kDa protein, Rv0203, was identified from pulldown experiments on Mtb culture filtrate using heme-agarose beads followed by mass spectrometry analysis, suggesting Rv0203 may represent an extracellular heme binding protein.90 The search for additional Mtb heme uptake proteins focused on the genomic vicinity surrounding Rv0203. Two other proteins, MmpL3 (Rv0206c) and MmpL11 (Rv0202c), were hypothesized to have a role in heme transport across the inner membrane as they are predicted to be large RND-like transmembrane transporters (Figure 13).

To directly probe the roles of Rv0203, MmpL3, and MmpL11 in heme uptake, knockout mutants were constructed in the mycobactin-deficient strain, MtbΔmbtB. The MtbΔmbtBΔmmpL3 mutant could not be made, as mmpL3 is an Mtb essential gene.149 Both MtbΔmbtBΔmmpL3 and MtbΔmbtBΔmmpL11 displayed normal growth kinetics when grown in media containing ferric iron supplemented with exogenous mycobactin. However, both mutants displayed a significant growth defect compared to MtbΔmbtB when grown in media with both iron and heme present but without exogenous mycobactin. This strongly suggests that both Rv0203 and MmpL11 are required for Mtb to utilize heme as an iron source. MmpL3 shares ~30% sequence homology with MmpL11 and, therefore, may also play a role in Mtb heme uptake.90

4.4.2. Structural Biology of Mtb Heme Uptake Proteins. 4.4.2.1. Rv0203 Structure. The crystal structure of apo-Rv0203 reveals an entirely α-helical protein that crystallizes as a dimer of dimers, where each individual Rv0203 protomer consists of five α-helices (Figure 14A).90 Each promoter in the tetramer structure is L-shaped and arranged around a central core composed of 4 α-helices. The central helices (α5) represent the Rv0203 C-termini and are critical in stabilizing dimer-dimer interactions. Tetrameric Rv0203 is also observed in solution, where Rv0203 exists in a concentration-dependent equilibrium with the dimeric form.

A structural homology search revealed that Rv0203 features a unique fold because there were no other proteins with an overall similar fold. Interestingly, structural comparison with the S. marcescens hemophore, HasA, which has no structural or sequence homology to Rv0203 (Figure 14A and B), reveals that Rv0203 surface residues Tyr59, His63, and His89 are arranged in a similar motif as the HasA heme binding residues His32, Tyr75, and His83 (Figure 14).116 The three Rv0203
residues are located in α-helices 1 and 3 within a hinge region on the protein surface that may be able to accommodate a heme molecule, leading to the hypothesis that Rv0203 residues Tyr59, His63, and His89 represent the Rv0203 heme binding site (Figure 14C).

4.4.2. Role of Rv0203. Mutagenesis studies demonstrated that the proposed heme binding residue Tyr59 is required for heme binding. Rv0203 variants Tyr59Ala and Tyr59Phe both bind heme poorly, suggesting this residue is essential in heme binding. Further evidence for Tyr-heme iron ligation is found in the spectroscopic signatures of Rv0203, which have the greatest similarity with a high-spin phenolate-ligated heme molecule. While the heme binding properties of Rv0203 are known, the holo-Rv0203 crystal structure has not yet been determined and several aspects of the Rv0203 heme binding mechanism remain unclear. The cleft between α-helices 1 and 3 in apo-Rv0203 is too small to fully accommodate a heme molecule. It remains to be seen if heme binding induces a conformational change to fully accommodate heme within the cleft. Kinetic studies of heme binding to Rv0203 suggest that the interaction is dominated by the Tyr-heme iron ligation, with little added contribution from noncovalent interactions. Compared to heme transport proteins in S. aureus and S. marcescens, Rv0203 binds heme relatively weakly as the rate of heme dissociation, $k_{off}$, is 0.08 s$^{-1}$ (Tables 2 and 3), precluding affinity-driven heme acquisition from metHb. Further experiments showed that Rv0203 does not steal heme from metHb through protein-protein interactions, suggesting Rv0203 may represent a hemophore that binds free heme, akin to the hemophore-like protein HmuY from P. gingivalis. Alternatively, Rv0203 may function as a heme

Figure 13. Mtb heme uptake. In the first pathway, the proposed proteins are colored in orange, and in the second they are light orange. In the first pathway, heme uptake involves a secreted protein, Rv0203. The heme source for Rv0203 is still not known. Rv0203 may bind free heme or work in combination with a hitherto unidentified partner that facilitates heme release from Hb. Heme transport across the mycobacterial outer membrane occurs via an unknown mechanism. Heme uptake across the inner membrane involves heme transfer between Rv0203 and two RND transmembrane transporters, MmpL3 and MmpL11, that accept heme from Rv0203 through protein-protein interactions. Finally, heme in the cytosol is degraded by MhuD to release iron. In the second pathway, PPE36 and PPE62 are cell-surface associated and predicted to bind free extracellular heme. FecB2 is a PBP protein, also predicted to bind heme, which shuttles heme through the Mtb periplasm. Asterisks indicate proposed steps.

Figure 14. Structure of Rv0203 (PDB: 3MAY) in comparison with HasA (PDB: 1DKH). (A) Tetrameric structure of Rv0203. The proposed heme binding site is boxed for one monomer. (B) Structure of HasA. (C) Model for holo-Rv0203. (D) Heme binding site of HasA.
transport protein that binds free heme and shuttles its cargo to the outer membrane and/or through the cell wall and periplasm. On the basis of the low-resolution CmpL1 structure, a model of MmpL3 was constructed, revealing the protein’s main structural elements (Figure 15A).158 The MmpL3 model consists of 12 TMs and 3 soluble domains, consistent with the predicted topology of the MmpL Cluster II subfamily (see Figure 6). The D1 and D2 porter subdomains interact on the periplasmic side, while the large D3 domain is located on the cytoplasmic side.158 On the basis of the CmpL1 structure, the MmpL3 trimer model is arranged around the central channel, presumed to be involved in substrate transport, and also features a periplasmic “head” domain located above the channel (Figure 15A). Notably, no density was observed for the cytoplasmic D3 domain. Because MmpL11 has a similar topological prediction as MmpL3, one expects MmpL11 to feature a similar overall architecture as MmpL3. MmpL3 and MmpL11 have minimal periplasmic domain oligomerization represents a functionally relevant behavior that occurs during inner membrane heme transfer.158

4.4.3. Structures of MmpL3 and MmpL11. MmpL3 and MmpL11 are both members of the RND superfamily of transmembrane transporters,90,92 as described in section 3.5.2. MmpL3 and MmpL11 have at least two roles in Mtb. Both are proposed to be involved in heme uptake and are also required for the export of important Mtb cell-wall lipids, including mycolic acids.90,153

Both MmpL3 and MmpL11 D1 domains bind heme. Their heme binding properties are similar, and both are characteristic of a mixed spin system with a combination of His, Tyr, and His/Tyr-heme iron ligation. Heme coordination behavior that occurs during inner membrane heme transfer.158

Table 2. Heme Binding Properties of Bacterial Heme Uptake Proteins

<table>
<thead>
<tr>
<th>protein</th>
<th>coordination</th>
<th>$k_{in}$ (µM$^{-1}$ s$^{-1}$)</th>
<th>$k_{off}$ (s$^{-1}$)</th>
<th>$K_s$ (M$^{-1}$)</th>
<th>ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rv0203</td>
<td>5c O/Tyr</td>
<td>133</td>
<td>0.0082</td>
<td>1.6 × 10$^7$</td>
<td>150</td>
</tr>
<tr>
<td>Mmp3-D1</td>
<td>5c/6c unknown ligands</td>
<td>29</td>
<td>$k_f = 0.0036$ (44%); $k_c = 0.0004$ (56%)</td>
<td>8.1 × 10$^7$</td>
<td>152</td>
</tr>
<tr>
<td>MmpL1-D1</td>
<td>5c/6c unknown ligands</td>
<td>53</td>
<td>0.34</td>
<td>1.6 × 10$^7$</td>
<td>152</td>
</tr>
<tr>
<td>PE22-PPE36</td>
<td>unknown</td>
<td>ND</td>
<td>ND</td>
<td>250</td>
<td>146</td>
</tr>
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<td>ND</td>
<td>ND</td>
<td>2500</td>
<td>144</td>
</tr>
<tr>
<td>Rv0265c (FecB2)</td>
<td>unknown</td>
<td>ND</td>
<td>ND</td>
<td>3333</td>
<td>146</td>
</tr>
</tbody>
</table>

**ND** = not determined, 5c = 5-coordinate, and 6c = 6-coordinate.

On rate refers to ferrous CO-heme binding. $k_f$ and $k_c$ represent the fast and slow rates of heme dissociation, respectively.

4.4.3.1. Heme Binding Properties of MmpL3 and MmpL11. Both MmpL3 and MmpL11 D1 domains bind heme. Their heme binding properties are similar, and both are characteristic of a mixed spin system with a combination of His, Tyr, and His/Tyr-heme iron ligation. Heme coordination to both D1 domains induces domain oligimerization, possibly along the heme interface. Given that MmpL3 and MmpL11 assemble into trimers, it is feasible that heme-induced D1 domain oligimerization represents a functionally relevant behavior that occurs during inner membrane heme transfer.152

Figure 15. (A) Predicted structural model of MmpL3 and (B) structure of MmpL11-D2 porter domain (PDB: 4Y0L).
The heme binding kinetics of both MmpL3 and MmpL11 D1 domains are complex and are dependent on both the heme redox and D1 oligomeric states. Ferrous-CO heme binding to both MmpL3 and MmpL11 D1 domains displays the typical linear relationship between heme concentration and binding rate, yielding association rates similar to those of other heme transport proteins. In contrast, ferric heme binding to D1 domains is multiphasic. Ferric heme binding reaction likely represents a composite of multiple events, including heme binding to D1 domains in various oligomeric states and heme-D1 oligomerization.152 The $K_a$ values of MmpL3 and MmpL11 D1 domains are higher than that of Rv0203, suggesting that directional transfer from Rv0203 to D1 domains is possible (Table 3) based on a heme affinity gradient.

4.4.3.2. Heme Transfer between Rv0203 and MmpL3 and MmpL11 D1 Domains. Rv0203 transfers heme to both MmpL3 and MmpL11 D1 domains. The transfer is both rapid and unidirectional. Stopped-flow experiments demonstrate that the rate of heme transfer between Rv0203 and both D1 domains is faster than passive heme dissociation from Rv0203, indicative that heme transfer is accelerated by protein–protein interactions.152 In contrast, the reverse transfer proceeds at the same slow rate of passive release by holo-D1, indicating that D1 domains do not transfer heme to Rv0203. To determine the efficiency of heme transfer, biotinylated heme-Rv0203 and MmpL3 and MmpL11 D1 domains were incubated together, and after separation of biotinylated heme-Rv0203 and MmpL3 and MmpL11 D1 domains, heme loss was measured. The $K_a$ values of biotinylated heme-Rv0203 and MmpL3 and MmpL11 D1 domains are higher than that of Rv0203, suggesting that Rv0203 transfers heme to both MmpL3 and MmpL11 D1 domains.

4.5. Second Mtb Heme Uptake Pathway

4.5.1. Identification of Protein Players. More recently, the Niederweiss group utilized an innovative methodology to identify potential cell-surface and periplasmic proteins thought to be involved in heme uptake. An Mtb transposon library screen was used to identify mutants with greater resistance to toxic heme analogue GaPPIX than wild-type Mtb.156 These experiments identified two mycobacteria-specific proteins (PPE36 and PPE62) and one previously described (see section 3.6.2) predicted periplasmic protein (FecB2) whose attenuated expression resulted in Mtb mutants with resistance to GaPPIX.147 This suggests that these three proteins may play a role in GaPPIX transport to its protein target or could be direct targets of the resulting GaPPIX toxicity. Moreover, the Mtb transposon mutants exhibited reduced growth in the presence of heme, as compared to wild-type Mtb, further suggesting that these proteins play a role in heme uptake. In support of these results, the Mtb$\Delta$ppe62 and Mtb$\Delta$fecB2 mutants also displayed attenuated growth in heme alone, while the Mtb$\Delta$ppe36 mutant was nonviable with heme as the sole iron source. Interestingly, the Mtb$\Delta$fecB2 mutant also had reduced growth in the presence of ferric-carboxymyoglobin, suggesting that FecB2 might play a role in both ferriderophore and heme acquisition pathways.

4.5.2. Predicted Protein Functions. Further biochemical analysis of PPE36, PPE62, and FecB2 demonstrated that both PPE proteins are likely cell-surface-exposed, while FecB2 is a membrane-tethered PBP. Additionally, it was shown by surface plasmon resonance that these proteins bind heme, albeit only weakly (Table 3).

Interestingly, the PPE proteins are unique to mycobacteria and usually form a heterodimer with their operonic mycobacteria-specific PE partner (they are named for conserved PE and PPE N-terminal motifs). Members of the Mtb PE/PPE protein family have been shown to be important for secreted or cell-wall-associated virulence and immunogenicity factors.159,160 PPE proteins usually have a conserved N-terminal 180-residue domain that forms a 4-helical bundle with its operonic PE protein partner. The PPE protein C-terminal regions are polymorphic and have a variety of different functions, such as enzymatic activities, or contain large repeat domains or attachment domains that interact with other Mtb proteins.161 PPE36, which is required for growth in heme-supplemented media, is a 243-residue protein and has a predicted 180-residue N-terminal PPE helical domain and a short disordered C-terminal domain. However, the functional role of PPE36 in heme uptake remains elusive.

PPE62 is a much larger protein than PPE36, comprising 582 residues. The N-terminal domain is predicted to adopt the expected PPE helical structure. The initial half of the C-terminal domain shows a right-handed /β-helix composed of three parallel /β-sheets repeats, which is structurally reminiscent of HxuA,162 a cell-surface Hx receptor in H. influenza involved in removing heme from Hx (Figure 10C). Thus, it is possible that PPE62 has a similar Hx-binding function as HxuA. However, it should be noted that growth of the PPE62 deletion mutant was only tested in media with free heme as the only iron source, and thus, it seems unlikely that the sole function of PPE62 is to steal heme from Hx, like HxuA.

FecB2 is a putative heme binding PBP chaperone. The structure of Mtb FecB2 has been solved in its apo-form (PDB code: 4PM4; Figure 16A), as described in section 3.6.2. FecB2 has a similar fold to PBP IsdE from S. aureus126 and PhuT from P. aeruginosa,163 both of which chaperone heme through the periplasm. Heme iron in IsdE is six-coordinate with methionine and histidine ligands (Met78 and His229) (Figure 16B),126 whereas PhuT donates a tyrosine ligand (Tyr71) to heme.

Figure 16. Top panels are ribbon diagrams of the PBPs. The bottom panels are close-ups of the substrate-binding sites. Siderophores are shown in white stick, and active-site Tyr and Arg residues are indicated. (A) Mtb FecB2 (PDB: 4PM4; blue), (B) S. aureus heme:IsdE complex (PDB: 2Q8Q; pink), and (C) P. aeruginosa heme:PhuT complex (PDB: 2R79; orange).
heme iron, which is further stabilized by two arginines, Arg73 and Arg228 (Figure 16C). Within the FecB2 small molecule binding site, there are two arginines (Arg184 and Arg221) and a tyrosine Tyr39, which may be involved in heme iron coordination (Figure 16A). FecB2 contains a predicted N-terminal signal peptide followed by a cysteine residue still present in its mature form, leading to the speculation that it is a lipoprotein localized to the periplasmic side of the inner membrane. Thus, it has been proposed that FecB2 may be an inner membrane-localized PBP that chaperones heme through the periplasm to an inner membrane heme transporter.

4.6. Problems with the Potential Components of Mtb Heme Uptake

4.6.1. MmpL Proteins As Small Molecule Exporters. As mentioned above, Rv0203 binds heme with a lower affinity than the secreted hemophore HasA and does not acquire heme from Hb. Nevertheless, Rv0203 was shown to be critical in efficient Mtb heme uptake. We proposed that Rv0203 is a free heme scavenger; however, it is also possible that an unidentified protein assists Rv0203 in scavenging heme from Hb, similar to the HmuY system described in P. gingivalis, where heme release from metHb is catalyzed by Hb proteases that partially degrade metHb.

MmpL proteins have been shown to export small molecules including lipids and siderophores. MmpL3 has been shown to export trahalose monomycolate and also acts as a flippase for mycolic acids. Additionally, MmpL11 has been shown to export lipids in Mtb and M. smegmatis. These data raise the question of whether MmpL3 and MmpL11 are really bidirectional transporters. An alternative role for MmpL3 and MmpL11 in heme metabolism has been presented, whereby the proteins’ function is to prevent heme toxicity within the Mtb cytoplasm by enabling heme secretion. However, experimental investigation of this hypothesis is required.

4.6.2. Proteins Identified by GaPPIX Screen have a Low Affinity to Heme. All three proteins, PPE36, PEE62, and FecB2, which were identified through the aforementioned GaPPIX Mtb transposon library experiment, bind heme with millimolar affinity. The heme-binding affinity of these proteins is at least 1000-fold lower as compared to other known proteins involved in bacterial heme uptake systems (Tables 2 and 3), raising uncertainty about their proposed roles in GaPPIX or heme transport. On the basis of the low heme-binding affinities, it is possible that these proteins are not the direct target of GaPPIX but play an indirect role in GaPPIX import and Ga toxicity.

FecB2 is proposed to be a PBP heme transporter protein similar to S. aureus IsdE; however, heme binding to IsdE is at least 6 orders of magnitude tighter than binding to FecB2. FecB2 has a lone Tyr39 in its small molecule binding pocket that could potentially coordinate heme iron, analogous to P. aeruginosa PhuT. However, in the FecB2 apo-structure, this tyrosine appears to be shielded from coordinating ferric iron as it π-stacks with Trp58 (Figure 16A). Because FecB2 has an extremely low affinity for heme compared to S. aureus IsdE, one could speculate that FecB2 does not function as a periplasmic heme chaperone.

Figure 17. (A) Structures of chromophore products of bacterial heme degradation. (B) Proposed mechanisms of oxidative heme degradation. In the first monoxygenase step, production of the intermediate meso-hydroxyheme is common to HO, MhuD, and IsdG/I. The HO (blue), MhuD (red), and IsdG/I (green) products branch off after this step to produce different chromophores, side-products, and iron.

Chemical Reviews
Review
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As mentioned in section 4.5.2, the C-terminal region of PPE62 is predicted to have a β-propeller-like fold similar to HxuA, a cell-surface Hx receptor involved in Gram-negative heme uptake.167 The low heme-binding affinity of PPE62 is consistent with its proposed structure, as HxuA binds Hx tightly but does not bind heme itself and HxuA is not required in free heme uptake.162 It therefore remains to be seen if PPE62, which was proposed to be part of a free-heme acquisition pathway,168 is a bifunctional protein with dual Hx and heme-binding properties, or if PPE62 is in fact an Hx-binding protein.

5. CYTOPLASMIC HEME DEGRADATION TO RELEASE IRON

5.1. Overview

The first discovered and most extensively studied heme-degrading enzyme is human heme oxygenase 1 (hHO-1), which oxidatively degrades heme to release ferrous iron (Figure 17).168 Heme oxygenase (HO) is the canonical heme-degrading protein in eukaryotes and has also been found in several bacteria including *P. aeruginosa*, *Corynebacterium diphtheriae*, and *Neisseria meningitidis*.169–171 In the past decade, a novel class of non-HO heme degraders has been discovered, the IsdG-type branch.172 Notably, Mtbc also harbors a noncanonical IsdG-type heme-degrading protein, MhuD. However, the MhuD and IsdG heme-degradation mechanisms differ substantially, and efforts have been made to understand the unique steps in MhuD heme degradation.

5.2. Heme Oxygenase

HOS degrade heme into the chromophore biliverdin IXα, carbon monoxide (CO), and ferrous iron (Figure 17).168 Multiple HO structures have been solved across many different species.172–174 The structure of canonical HO is composed of an entirely α-helical fold (Figure 18).173 The heme-binding pocket is composed of proximal and distal helices, where the heme iron is coordinated by a histidine residue on the proximal side with a network of ordered waters on the distal side.175 Heme is anchored in the HO active site by Arg and Lys residues that interact with the heme propionates. Moreover, the heme molecule is slightly solvent-exposed and nearly planar.173

The mechanism for HO-mediated heme degradation occurs via three successive monooxygenation reactions, reviewed extensively in ref 175. The first monooxygenation step is activation of molecular oxygen by the heme iron center. The ferric heme is reduced to its ferrous form while coordinating dioxygen and is converted into the activated ferric hydroperoxy [Fe3+−O2H] intermediate that is stabilized by the network of water molecules surrounding the distal heme pocket.173,176–178 The Fe3+−O2H species is cleaved, resulting in the production of the α-meso-hydroxoheme intermediate.179,180 In the second monooxygenase step, the α-meso-hydroxoheme reacts with molecular oxygen to yield the second intermediate, α-verdoheme, and also results in the release of CO.181 Finally, a second round of molecular oxygen activation occurs during the third and final monooxygenation step, followed by hydroxylation of α-verdoheme to produce the final product, biliverdin, and to release ferrous iron.175

5.3. IsdG-Type Heme Oxygenases

5.3.1. Overview. The noncanonical IsdG-type heme-degrading protein family is the other major class of heme-degrading proteins, and the primary sequence and structure of IsdG-type enzymes are quite distinct from canonical HOs.172 IsdG-type heme-degrading proteins are primarily found in bacteria, although recently a branch has been discovered in lower eukaryotes.182 The family is named after IsdG, which was first characterized in *S. aureus*. IsdG and its homologue IsdI are part of the *S. aureus* isd heme uptake operon.183

5.3.2. Structure and Mechanism of IsdG-Type Proteins. IsdG-type heme-degrading enzymes are structurally distinct from HOx184,185 and consist of a homodimer exhibiting a ferrodoxin-like α+β-barrel fold, a similar fold to that in Figure 19A.184,185 There are two predominately hydrophobic heme-binding active sites per homodimer. Like HO, the heme iron is coordinated through a histidine residue (His75 in *S. aureus* IsdG numbering) on the proximal side; however, on the distal side, the water network is absent (Figure 19B). Instead, the heme iron on the distal side is coordinated to a chloride ion that in turn is coordinated by Asn6.185 IsdG and IsdI convert heme to novel chromophores, staphylobilin isomers,186 formaldehde, and iron (Figure 17).187 The staphylobilin isomers result from cleavage of the tetrapyrrole ring at the β/δ-meso-carbon and an oxygenation reaction at the δ/β-meso-carbon, respectively, followed by the release of the β/δ-meso-carbon as formaldehyde.

5.3.3. Heme Ruffling That Is Required for Heme Activation in IsdG-Type Heme Degraders. A defining feature in *S. aureus* IsdG-type enzymes is the presence of a highly ruffled heme in their active sites.185 The *S. aureus* IsdG-N7A and IsdI structures feature extraordinary out-of-plane heme distortions of 1.9 and 2.3 Å, respectively (Figure 19C).185 NMR experiments show that the observed heme ruffling appears to alter the electronic structure of the heme, possibly sensitizing the heme meso-carbons to nucleophilic attack.186 The requirement for heme ruffling in IsdG-type heme degradation was demonstrated by mutational analysis. Within the IsdG/I heme-binding pocket, Trp66 forms a hydrophobic interaction with the heme protoporphyrin ring to

![Figure 18. Structure of hHO-1 (PDB: 1N45) showing the water network surrounding the distal heme pocket.](image-url)
promote ruffling. An IsdG-Trp66Ala variant resulted in a reduction of heme ruffling, which correlated with a severe attenuation in heme degradation. Moreover, the heme electronic structure in IsdG-Trp66Ala is different from that of wild-type MhuD, such that the meso-carbons are probably not activated. Together, these results suggest that heme ruffling in IsdG-type heme-degrading proteins is required to increase the heme molecule’s reactivity in the hydrophobic active site.

Despite knowing the heme-bound IsdG and IsdI structures, our understanding of the intermediate steps in IsdG/I-catalyzed heme degradation is poor. However, several elegant studies partially elucidate the IsdG mechanism and suggest that the first oxidation step forms the β/δ-meso-hydroxyheme intermediates.

5.3.4. IsdG-Type Proteins Require an Electron Donor. IsdG and IsdI require reducing equivalents to degrade heme. It has been shown that S. aureus proteins, NtrA and IruO, donate electrons to IsdG and IsdI heme-degrading proteins. IruO is a FAD-containing NADPH-dependent reductase, while NtrA is also thought to be a novel nitroreductase capable of S-nitrosoglutathione activity.

5.4. Mtb MhuD Heme Degradation

5.4.1. Overview. An IsdG-type protein is also found in Mtb, MhuD (mycobacterial heme-utilizing degrader). MhuD shares high sequence and structural homology with IsdG-type heme-degrading proteins; however, MhuD degrades heme into unique chromophores, mycobilin isomers (Figure 17), and iron.

5.4.2. Structure of MhuD. Similar to IsdG and IsdI, MhuD forms a homodimeric ferredoxin-type α+β-barrel fold (Figures 19A and 20A). The structure of MhuD has been solved in both a diheme form (discussed later) and a cyanide-inhibited monoheme form. In the cyanide-inhibited MhuD-monoheme structure, heme iron is coordinated by proximal His75 and a distal cyanide. Most notably, the heme molecule in the MhuD active site is rotated ~90° about the axis normal to the tetrapyrole ring compared to the heme orientation observed in IsdG and IsdI. Examination of the MhuD-monoheme structure revealed that the heme molecule was distorted by 1.4 Å, a decrease from the 1.9 and 2.3 Å heme distortions observed in the IsdG-Asn7Ala and IsdI structures, respectively (Figure 19C). Heme distortion, like in IsdG and IsdI, appears to be induced by the neighboring Trp66 residue, and mutation of Trp into a less bulky hydrophobic residue results in reduced heme-degrading activity. Variable-temperature magnetic circular dichroism, NMR, and electronic spectroscopy experiments in MhuD indicate that ruffling of the heme delocalizes spin density from the central iron and pyrrole rings, as also seen in IsdI, potentially making the meso-carbons more susceptible to nucleophilic attack.

5.4.3. Mechanism of MhuD Heme Degradation. MhuD heme degradation can be divided into two steps. It was suggested that the first step in MhuD heme degradation is similar to the monooxygenation step of IsdI and HO (Figure 17). In this mechanism, MhuD is proposed to oxidize heme into β/δ-meso-hydroxyheme intermediates via formation of a [Fe3+–O2−] species. It was also proposed that there is no release of CO in MhuD-dependent heme degradation, and so it follows that there is no formation of the verdoheme intermediate. Once the β/δ-meso-hydroxyheme intermediate is formed (Figure 17), it has been demonstrated that meso-hydroxyheme is converted into mycobilin by a dioxygenase step. For the second step to occur, meso-hydroxyheme is thought to undergo radical localization to the α-meso-carbon before dioxygenation at the α-meso position into a dioxygenate intermediate, which is proposed to spontaneously decompose into Fe-mycobilin (Figure 17). Thus, MhuD degradation of heme is highly unusual as its active site performs two different reactions, a monooxygenase reaction followed by a dioxygenase reaction. The first monooxygenase reaction, which requires a proton-donating environment, is possibly the rate-limiting step in the hydrophobic MhuD heme-binding pocket. However, it is thought that distal Asn7, one of the only polar residues in the distal heme pocket, forms a hydrogen bond to the terminal hydroxyl group of Fe3+–O2H to stabilize the intermediate. The second dioxygenase reaction step, where the ruffled heme may allow radical localization on the α-meso-carbon, is likely enabled by the MhuD hydrophobic active site environment. Furthermore, heme ruffling may also promote the initial monooxygenase reaction by changing the electronic structure of heme iron. Thus, it appears that heme ruffling is required for the coupled monooxygenase and dioxygenase reactions within the MhuD active site.

5.4.4. Unique Diheme Form of MhuD. MhuD is capable of binding up to two hemes per active site (Figure 20A). The feature has not been observed in any other IsdG-type protein.
studied to date. In the MhuD-diheme structure, the protoporphyrin rings are stacked 3.45 Å apart and oriented such that the propionate groups from one heme are rotated ∼80° relative to those from the second heme (Figure 20B).195

The more solvent-exposed heme iron is coordinated by His75, while the solvent-protected heme iron is coordinated by a chloride ion and further stabilized by Asn7.195 The MhuD active site can accommodate two heme molecules through the straightening of α-helix 2 that is kinked in the MhuD-monomer structure, together with a conformational change in the flexible loop region following α-helix 2 (Figure 20B). Furthermore, both heme molecules in each MhuD-diheme active site are essentially planar.195 As one might anticipate, the MhuD-diheme form had no heme-degrading activity. Loss of activity could be due to three possible scenarios: (1) the planarity of the two heme molecules resulting in a diheme monoheme structure, together with a conformational change in the flexible loop region following α-helix 2 (Figure 20B). Furthermore, both heme molecules in each MhuD-diheme active site are essentially planar.195

5.4.5. Potential Mtb MhuD Electron Donors. An electron donor is essential for MhuD-dependent heme cleavage. There are numerous Mtb candidates that could be potential electron donors based on homology to S. aureus IsdG electron-donor proteins, IruO and NtrA.192,194 All candidates, however, remain to be tested to elucidate the electron donor for MhuD.

6. IRON-ACQUISITION PATHWAYS AS ANTI-TB DRUG TARGETS

6.1. Targeting the Mycobactin Biosynthetic Pathway

Over the last several decades, there have been tremendous efforts to design inhibitors to interrupt mycobactin-mediated iron acquisition and, in particular, to inhibit mycobactin biosynthesis (Figure 4).57,201 Although 14 proteins are responsible for the biosynthesis of mycobactin, only two have been studied extensively as potential drug targets, MbtA and MbtI. Both MbtA and MbtI catalyze initial steps in the biosynthetic pathway (Figure 4B). MbtA catalyzes the conversion of chorismate to salicylic acid, the substrate of the Mbt megasynthase complex. MbtA activates salicylic acid and catalyzes its transfer to the thiolate domain of MbtB through an acyl adenylate intermediate (Figure 4B (4)). Many of the designed inhibitors are based on chorismate (Figure 4B (1)) and isochorismate analogues (Figure 4B (2)) for MbtI and variations of the acyl-adenylate intermediate (Figure 4B (4)) for MbtA. Additionally, some progress has been made on developing inhibitors for MbtM, which is involved in fatty acid chain activation for linkage to the mycobactin core structure.59 For more in depth information on inhibitors for MbtA, MbtI, and MbtM, we direct the reader to a detailed review by Meneghetti, Villa, and co-workers.57

A potential new anti-TB drug target within the mycobactin biosynthetic pathway is MbtH. It is required for the correct folding of the three NRPSs, MbtB, MbtE, and MbtF, which are constituents of the megasynthase involved in mycobactin core synthesis.62 Thus, interruption of MbtH chaperone activity for the Mtb NRPSs would disrupt mycobactin synthesis and consequently inhibit iron acquisition and Mtb growth.

6.2. Targeting Iron Uptake Pathways

The proteins involved in Mtb mycobactin export and import represent both a new set of anti-TB drug targets and a potential gateway to introduce antibacterials via a “Trojan horse-like” mechanism.

As siderophore receptors and import/export machinery are highly selective, compounds with a similar structure to the mycobactin core (Figure 3) could have an inhibitory effect on the mycobactin-dependent iron uptake pathway. In fact, M. smegmatis mycobactin S (a stereoisomer of Mtb mycobactin T) is toxic to Mtb.202 A recent example further highlights the potential of applying a Trojan horse strategy to inhibit Mtb growth. In one study, mycobactin was conjugated with artemisinin (Figure 21A), a natural product isolated from the plant Artemisia annua and a known treatment for Plasmodium falciparum malaria.205 In P. falciparum, artemisinin is activated by heme and results in its cleavage and generation of free radicals that in turn damage many susceptible proteins.204 The mycobactin-artemisinin conjugate (Figure 21B) has strong and selective inhibitory effects against Mtb (MIC = 0.39 μg/mL) as well as MDR and XDR strains,205 thus demonstrating that mycobactin can be utilized in a Trojan horse-like manner.
Another mycobactin analogue of particular interest is a recently synthesized mycobactin-maleimide analogue that already has inhibitory effects against Mtb (MIC$_{90}$ = 0.88 μg/mL) and will allow the eventual synthesis of a series of novel mycobactin-drug conjugates. Finally, siderophores coordinated to toxic metals have been shown to inhibit bacterial growth. For example, when E. coli uptakes the scandium-enterobactin complex, bacterial growth is attenuated, suggesting that this might be an alternate avenue for exploiting Mtb mycobactins as therapeutics.

Another approach to attenuate Mtb growth is to design iron chelators that target the Mtb cytoplasm. A novel compound, pyrazolopyrimidinone (PZP) (Figure 22), was found to attenuate Mtb growth; however, mutations in the ESX-3 complex (see Figure 8) resulted in resistance to PZP. Because ESX-3 is thought to play a role in mycobactin transport, there was speculation that PZP acts by interfering with mycobactin-dependent iron uptake. However, further investigation revealed that PZP acts as an Mtb cytoplasmic iron chelator and does not interrupt mycobactin-dependent iron acquisition pathways.

Notably, in a comprehensive study identifying genes involved in Mtb antibiotic resistance, FecB was central to all antibiotics tested. Mtb FecB is a proposed PBP involved in siderophore-mediated iron uptake—see section 3.6.2. A MtbΔfecB mutant was hypersensitive to the uptake of a variety of antibiotics, including vancomycin and verapamil. Thus, FecB may be a good anti-TB drug target as its inhibition may prevent the development of new MDR and XDR Mtb strains.

6.3. Targeting Heme Uptake Pathways

As Mtb heme uptake was only discovered in 2011, these pathways have not been actively investigated as anti-TB drug targets; however, heme-dependent iron uptake pathways may be suited for Trojan horse strategies or be targeted directly. A successful drug candidate would specifically target Mtb heme transport proteins, while having low affinity to host heme-binding proteins.

Heme analogues including metal-substituted protoporphyrins (MPPs) have antibacterial activity. Nearly two decades ago, it was demonstrated that GaPPIX was toxic against various bacteria including M. smegmatis. Further, MPP analogues were shown to inhibit heme import ABC transporters in Trypanosoma cruzi as well as disrupt the ability of S. aureus NEAT domain proteins to shuttle heme to the inner membrane-associated transport protein complex. Within the last five years, efforts have refocused on MPPs as potential antibacterials. Notably, in P. aeruginosa it was shown that GaPPIX inhibits growth by targeting cytochromes and likely interferes with cellular respiration. It was also demonstrated that both Ga- and Sn-PPIXs inhibit growth of Mycobacterium abscessus. Additionally, in M. abscessus-infected macrophages, treatment with MPPs inhibited mycobacterial growth but did not exhibit toxicity toward host macrophages. Moreover, 1 μM GaPPIX inhibited intracellular M. abscessus growth to a level significantly greater than 25 μM Ga(NO$_3$)$_3$, demonstrating that GaPPIX has greater antimycobacterial power than the toxic metal, Ga$^{3-}$, alone.

In the Sher laboratory, MPPs have been tested as antibacterials against Mtb infection. Mtb-infected mice treated with SnPPIX in combination with conventional anti-TB drugs had a significantly reduced bacterial load compared to Mtb-infected mice treated with anti-TB drugs alone. The overall effect of this combination SnPPIX therapy was to increase the rate of Mtb clearance from 21 weeks for the conventional anti-TB therapy alone to 17 weeks. However, SnPPIX was shown to be a potent inhibitor of host hHO-1 in vitro. In contrast, the Mtb heme-degrading protein MhuD was not inhibited by SnPPIX. Furthermore, SnPPIX does not inhibit Mtb growth in vitro with an MIC$_{90}$ of <200 μM. Taken together, these results suggest that the in vivo SnPPIX effect against Mtb is due to SnPPIX acting as a host-directed therapy upon mammalian HO rather than an Mtb-directed therapy. Unlike SnPPIX, GaPPIX does inhibit Mtb growth in vitro with an MIC$_{90}$ of 1.5 μM. Thus, it remains to be seen if GaPPIX may act as both a host-directed and an Mtb-directed therapy.

In addition to their potential as anti-TB agents, heme analogues may provide insights into the mechanism of heme uptake and be utilized to identify new players in the Mtb heme acquisition pathways. This has, in part, already been demonstrated with the identification of transposon mutants that are resistant to GaPPIX—as discussed in section 4.5.1. Moreover, heme analogues could be used to assess the importance of heme acquisition in infectivity and virulence.

Significantly, over the last 7 years, MmpL3 has become the most successful anti-TB drug target. High-throughput whole-cell screens identified several potent antimycobacterial agents that target MmpL3, including BM212 and SQ109. However, these agents attenuate Mtb growth by acting upon MmpL3 mycolic acid flippase function or by dissipating the energy of proton motive force across the inner membrane. Thus, it is unlikely that the effect of the MmpL3 inhibitors is due to the interruption of heme uptake. Compounds that target MmpL3 have been extensively reviewed in refs 154, 214, 220, and 221.

7. CONCLUSIONS

Over the last two decades, bacterial heme and iron acquisition pathways have been discovered and mapped out in a number of organisms. Nevertheless, the relationship between heme uptake and virulence is not always straightforward. The contribution of heme acquisition to virulence has been firmly established in only a select number of bacteria. Most notably, elegant work by Skaar and others demonstrated that in S. aureus heme iron is the preferred iron source over transferrin iron in the early stage of S. aureus infection. Additionally, in uropathogenic E. coli it has been shown that heme uptake is necessary for urinary tract colonization, and in P. gingivalis, which lacks an endogenous heme biosynthesis pathway, heme uptake is a critical virulence factor. These observations suggest that heme acquisition pathways have excellent potential as antibacterial targets. However, except for these referenced cases, the interplay between heme and iron...
uptake during early, active, and secondary pathogen infections or dormancy remains largely unknown.

Two mechanisms of iron acquisition have been discovered in Mtb, iron and heme uptake. Iron acquisition is dependent on mycobactin siderophores. Mycobactins are required for initial stages of Mtb infection.40,46 Thus, targeting Mtb mycobactin metabolism represents a promising strategy in designing novel anti-TB drugs, and to date, three mycobactin biosynthesis enzymes, MbtA, MbtI, and MbtM are being investigated as drug targets. However, further research is needed to create a broader understanding of Mtb iron metabolism because there are still large knowledge gaps surrounding the proteins involved in siderophore export and import. Unlike siderophore-mediated iron uptake, a clear link between heme uptake and infection has not yet been established in Mtb. To effectively understand how Mtb heme uptake can be used as a drug target, it is imperative that future research focuses on understanding if heme uptake is predominately utilized in the initial stage of Mtb infection, in secondary infection, or even in the dormant state.

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Notes
The authors declare no competing financial interest.

Biographies

Alex Chao received his Bachelor’s degree in Chemistry from Arizona State University, where he performed undergraduate research examining the insulin signaling pathway as well as studying a bacterial surface adhesion protein. Alex is now a Ph.D. candidate at the UC Irvine in the Department of Molecular Biology and Biochemistry and is a recipient of the prestigious National Science Foundation Graduate Research Fellowship. Currently, he is working to biochemically and structurally characterize the Mtb heme-degradation enzyme. Additionally, he is working to determine the structures of proteins involved in Mtb iron uptake.

Paul Sieminski studied glycerol–water mixtures at various compositions, using physical and thermodynamic methods to elucidate phase transitions in the system during his Bachelor’s degree in Biology at Pennsylvania State University. Interested in applying these principles to biological systems, using X-ray crystallography and biochemical methods, he initiated characterization of an important protein in Mtb siderophore trafficking during his Master’s degree at UC Irvine. Currently, Paul is a doctoral student studying structural biology and biophysics in the Department of Chemistry & Biochemistry at UC Los Angeles.

Cedric Owens is an Assistant Professor of Biochemistry at Chapman University in Orange, CA. After growing up and completing his Abitur in Bayreuth, Germany, he moved to Waterville, ME, to obtain B.A. degrees in Chemistry and Mathematics Sciences from Colby College. For his Ph.D., he joined the lab of Dr. Celia Goulding at UC Irvine, where he characterized mycobacterial heme uptake. From there, he moved to UC San Diego to investigate electron transfer in nitrogenase in the lab of Dr. Akif Tezcan as a USDA NIFA postdoctoral fellow. Currently, Cedric’s research interests focus on characterizing and engineering regulation of nitrogen fixation in agriculturally relevant bacteria.

Celia Goulding is a Professor of Molecular Biology and Biochemistry and Pharmaceutical Sciences at the University of California, Irvine. She received a B.Sc. (Hons) in chemistry and mathematics and a Ph.D. in physical organic chemistry from King’s College, London. After leaving the United Kingdom for the United States, she was a postdoctoral fellow with Professor Rowena Matthews at the University of Michigan studying metalloprotein enzymology. Moving further west to UCLA, as research faculty under the supervision of Professor David Eisenberg, Celia became interested in the mechanisms by which Mtb is able to survive in human hostile environments, employing X-ray crystallography and other biophysical and biochemical techniques. During her time at UCLA, she also spearheaded the Tuberculosis Structural Genomics Consortium, which has resulted in ~350 Mtb protein structures being deposited in the Protein Databank since its inception in 2001. Celia eventually accepted an independent position at UC Irvine where she and her group investigate the function and structure of metalloproteins involved in heme and iron uptake pathways in Mtb.

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ABBREVIATIONS

ABC ATP-binding cassette
AIDS acquired immune deficiency syndrom
CO carbon monoxide
FAD flavin adenine dinucleotide
FeEnt ferric-enterobactin
GaPIX gallium-protoporphyrin IX
Hb hemoglobin
hHO-1 human heme oxygenase 1
HIV human immunodeficiency virus
HO heme oxygenase
Hp haptoglobin
Hx hemopexin
IdE ir-dependent regulator
ITC isothermal titration calorimetry
Ll lactoferrin
MDR-TB multidrug-resistant TB
MFS major facilitator superfamily
MhuD mycobacterial heme-utilizing degrader
MMP metalloprotoporphyrin IX
MmpL mycobacterial membrane protein large
MmpS mycobacterial membrane protein small
Mtb Mycobacterium tuberculosis
NEAT near iron transport
NRPS nonribosomal peptide synthetase
NTD N-terminal domain
Omp outer membrane porin
PBP periplasmic binding protein
PPIX protoporphyrin IX
PZP pyrazolopyrimidinone
RD resistance nodulation cell division
SAM S-adenosylmethionine
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