Recent Insights into the Structure and Function of Mycobacterial Membrane Proteins Facilitated by Cryo-EM

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Abstract

Mycobacterium tuberculosis (Mtb) is one of the deadliest pathogens encountered by humanity. Over the decades, its characteristic membrane organization and composition have been understood. However, there is still limited structural information and mechanistic understanding of the constituent membrane proteins critical for drug discovery pipelines. Recent advances in single-particle cryo-electron microscopy and cryo-electron tomography have provided the much-needed impetus towards structure determination of several vital Mtb membrane proteins whose structures were inaccessible via X-ray crystallography and NMR. Important insights into membrane composition and organization have been gained via a combination of electron tomography and biochemical and biophysical assays. In addition, till the time of writing this review, 75 new structures of various Mtb proteins have been reported via single-particle cryo-EM. The information obtained from these structures has improved our understanding of the mechanisms of action of these proteins and the physiological pathways they are associated with. These structures have opened avenues for structure-based drug design and vaccine discovery programs that might help achieve global-TB control. This review describes the structural features of selected membrane proteins (type VII secretion systems, Rv1819c, Arabinosyltransferase, Fatty Acid Synthase, F-type ATP synthase, respiratory supercomplex, ClpP1P2 protease, ClpB disaggregate and SAM riboswitch), their involvement in physiological pathways, and possible use as a drug target.

Graphic Abstract

Tuberculosis is a deadly disease caused by Mycobacterium tuberculosis. The Cryo-EM and tomography have simplified the understanding of the mycobacterial membrane organization. Some proteins are located in the plasma membrane; some...
span the entire envelope, while some, like MspA, are located in the mycomembrane. Cryo-EM has made the study of such membrane proteins feasible.

Keywords  Cryo-EM · Drug discovery · Membrane protein · Mycobacterium · Protein structure

| Abbreviations | Description |
|---------------|-------------|
| AAA           | ATPases associated with diverse cellular activities |
| ABC           | ATP-binding cassette |
| Acyl         | Acyl-carrier protein M, Acr1: alpha-crystallin-related protein 1 |
| Adn           | ATP-dependent nuclease |
| AftD          | Arabinofuranosyltransferase D |
| AMPPNP        | Adenylly imidodiphosphate |
| BBH           | Benzoxylbenzylidene-hydrazine |
| BDQ           | Bedaquiline |
| Bpa           | Bacterial proteasome activator |
| CBM           | Carbohydrate-binding module |
| CDL           | Cardiolipin |
| CIII2/CIV2SOD2 | C_{III} and C_{IV} subunits of respiratory supercomplex bound to superoxide dismutase in C2, symmetry |
| Clp           | Caseinolytic proteases |
| Cryo-EM       | Cryo-Electron Microscopy |
| Cryo-ET       | Cryo-Electron Tomography |

| Subunits of aa_{3}-type cytochrome c oxidase |
| C-terminal domain |
| Decaprenyl-phosphate-arabinose |
| ESC conserved component |
| Ethambutol |
| 6 KDa early secretory antigenic target |
| ESCX-1 secretion-associated protein |
| ESAT-6 |
| Electron transport chain |
| Fatty Acid Synthase |
| Galactofuranosyl |
| Z-Gly-Leu-Phe-chloromethyl ketone |
| Glycine-glutamine-tyrosine-leucine |
| Human immunodeficiency virus |
| Heat shock protein |
| Mycobacterial-specific protein Y |
| Membrane spanning domain |
| Mycobacterium tuberculosis |
| Multi/extremely drug resistant |
| Nucleotide-binding domain |
| Proteasome accessory factor E |
Introduction

*Mycobacterium tuberculosis* (Mtb) is the causative agent of tuberculosis (TB)—a disease that costs as many as 200,000 deaths per year worldwide, and almost a quarter of the world population is infected with it (Gaensbauer and Broadhurst 2019). Even after decades of research, there still is a lack of an effective short treatment course or a potent vaccine and better diagnostics. Due to non-compliance with the drug regimen, a high percentage of TB patients drop-out from the treatment program that lasts at least 6 months in general (Awofeso 2008; Churchyard 2018). This high-dropout rate has led to increased cases of TB relapse and drug resistance, making TB even more challenging to treat. Although new drugs have been introduced, none of them have proven to be completely effective. The only notable exception is the use of Pretomanid in combination with Bedaquiline (BDQ) and Linezolid for the treatment of a specific type of highly treatment-resistant TB of the lungs that has shown promise (Bigelow et al. 2020).

Given the current situation, the World Health Organization’s (WHO) ‘End the TB’ strategic program that aims to reduce TB incidences by 80% and deaths by 90% by the year 2030 seems rather ambitious (World Health Organization 2020). The ongoing COVID-19 pandemic has made the situation even more challenging (Grüber 2020) by severely disrupting the TB, Human Immunodeficiency Virus (HIV), and Malaria programs worldwide. In addition, according to recent estimates, COVID-19 has set back worldwide TB treatment efforts by almost a decade and reducing gains to control the disease (The Global Fund 2020). Under such circumstances, we are in urgent need of effective treatment for TB.

A deep understanding of the mechanism of action and atomic details of critical elements contributing to any pathogenic micro-organism’s virulence is crucial for successful and effective drug development. The mycobacterial membrane enables the bacilli to survive in the hostile environment inside the host (Lin et al. 2002; Hahn et al. 2005; McClean 2012). The characteristic cell envelope, slow growth rate and unique metabolic features (e.g. glycolate pathway, diauxic growth, etc.) of Mtb make it an exciting but challenging system to study (Faller et al. 2004; Målen et al. 2011; Koul et al. 2014). The Mtb cell wall has a distinctive composition, and despite close resemblance with Gram-positive bacteria, its outer membrane contains a complex mixture of lipids such as mycolic acids that distinguishes it from other bacteria (Marrakchi et al. 2014).

An extensive physicochemical and structural characterization of Mtb membrane samples could help us understand its composition. Further, various ‘omics’ studies have revealed information about constituent membrane proteins and the secretome of the pathogen. However, there is a large gap in the understanding of the Mtb membrane proteins from their structural point of view. This has been a significant bottleneck in deciphering the roles and mechanism of action of these proteins, which in turn has thwarted the discovery and development of novel therapeutics.

Cryo-electron microscopy (cryo-EM) is proving instrumental in breaking this bottleneck (Munir et al. 2021). Recent advances in cryo-EM have been the silver lining and have proven to be extremely helpful in unraveling the structures of key drug targets of various pathogens, including Mtb. A relatable example has been structure elucidation of various membrane proteins and drug targets of Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) in record time. This has advanced drug discovery and vaccine development efforts against SARS-CoV-2 worldwide (Jeong et al. 2020; Wrapp et al. 2020). Not surprisingly, cryo-EM has also helped determine structures of key Mtb proteins that are important for virulence and survival of the pathogen and are important targets for drug discovery and vaccine development programs. Furthermore, developments in cryo-electron tomographic (cryo-ET) techniques and cryo-EM have contributed a tremendous amount of structural knowledge about mycobacterial membrane organization and its constituent membrane proteins. This has proven instrumental in modern drug discovery and a better understanding of host–pathogen interactions (Sani et al. 2010b).

So far, 75 cryo-EM structures of mycobacterial proteins have been deposited in the Protein data bank (PDB) (Table 1) (Burley et al. 2019). The majority of these structures are membrane proteins. However, very few of these have been tested as targets for drug discovery, while others are yet to be explored. In this review, we discuss the structures of some of the key Mtb proteins and protein complexes elucidated by EM. The list includes cryo-EM structures of type VII secretion systems, Rv1819c, Arabinosyltransferase, Fatty Acid
| Sr. no | PDB code | Title of the PDB Entry | References |
|--------|----------|------------------------|------------|
| 1      | 5V93     | Cryo-EM structure of the 70S ribosome from *M. tuberculosis* bound with Capreomycin | Yang et al. (2017) |
| 2      | 5ZEP     | *M. smegmatis* hibernating state 70S ribosome structure | Mishra et al. (2018) |
| 3      | 6DZI     | Cryo-EM Structure of *M. smegmatis* 70S C(minus) ribosome 70S-MPY complex | Li et al. (2018a) |
| 4      | 5ZEB     | *M. smegmatis* P/P state 70S ribosome structure | Mishra et al. (2018) |
| 5      | 5V7Q     | Cryo-EM structure of the large ribosomal subunit from *M. tuberculosis* bound with a potent linezolid analog | Yang et al. (2017) |
| 6      | 5ZET     | *M. smegmatis* P/P state 50S ribosomal subunit | Mishra et al. (2018) |
| 7      | 6DZP     | Cryo-EM Structure of *M. smegmatis* C(minus) 50S ribosomal subunit | Li et al. (2018a) |
| 8      | 6DZK     | Cryo-EM Structure of *M. smegmatis* C(minus) 30S ribosomal subunit with MPY | Li et al. (2018a) |
| 9      | 5XYM     | Large subunit of *M. smegmatis* ribosome | Li et al. (2018b) |
| 10     | 5XYU     | Small subunit of *M. smegmatis* ribosome | Li et al. (2018b) |
| 11     | 5ZEU     | *M. smegmatis* P/P state 30S ribosomal subunit | Mishra et al. (2018) |
| 12     | 5061     | The complete structure of the *M. smegmatis* 70S ribosome | Hentschel et al. (2017) |
| 13     | 5060     | 50S large ribosomal subunit from *M. smegmatis* | Hentschel et al. (2017) |
| 14     | 505J     | 30S small ribosomal subunit from *M. smegmatis* | Hentschel et al. (2017) |
| 15     | 6EDT     | *M. tuberculosis* RNAP open promoter complex with RbpA/CarD and AP3 promoter | Boyaci et al. (2019) |
| 16     | 6EE8     | *M. tuberculosis* RNAP promoter unwinding intermediate complex with RbpA/CarD and AP3 promoter | Boyaci et al. (2019) |
| 17     | 6EEC     | *M. tuberculosis* RNAP promoter unwinding intermediate complex with RbpA/CarD and AP3 promoter captured by Corallopyronin | Boyaci et al. (2019) |
| 18     | 6C04     | *M. tuberculosis* RNAP Holo/RbpA/double fork DNA-closed clamp | Boyaci et al. (2018) |
| 19     | 6BZO     | *M. tuberculosis* RNAP Holo/RbpA/Fidaxomicin/upstream fork DNA | Boyaci et al. (2018) |
| 20     | 6FBV     | *M. tuberculosis* RNA polymerase in complex with Fidaxomicin | Lin et al. (2018) |
| 21     | 6HWH     | Functional obligate respiratory supercomplex from *M. smegmatis* | Wiseman et al. (2018) |
| 22     | 6C06     | *M. tuberculosis* RNAP Holo/RbpA/Fidaxomicin | Boyaci et al. (2018) |
| 23     | 6C05     | *M. tuberculosis* RNAP Holo/RbpA in relaxed state | Boyaci et al. (2018) |
| 24     | 6M7J     | *M. tuberculosis* RNAP with RbpA/Us fork and Corallopyronin | Boyaci et al. (2019) |
| 25     | 5LZP     | Binding of the C-terminal GQYL motif of the bacterial proteasome activator Bpa to the 20S proteasome | Bolten et al. (2016) |
| 26     | 7BVF     | Cryo-EM structure of *M. tuberculosis* arabinosyltransferase EmbA-EmbB-AcpM2 in complex with ethambutol | Zhang et al. (2020b) |
| 27     | 6SGZ     | *M. smegmatis* Structure of protomer 2 of the ESX-3 core complex | Famelis et al. (2019) |
| 28     | 6SGX     | *M. smegmatis* Structure of protomer 1 of the ESX-3 core complex | Famelis et al. (2019) |
| 29     | 6SGW     | *M. smegmatis* Structure of the ESX-3 core complex | Famelis et al. (2019) |
| 30     | 6BGL     | Doubly PaF-E-capped 20S core particle in *M. tuberculosis* | Hu et al. (2018) |
| 31     | 6BGO     | Singly PaF-E-capped 20S CP in *M. tuberculosis* | Hu et al. (2018) |
| 32     | 6EYD     | *M. smegmatis* RNA polymerase Sigma-A holoenzyme | Kouba et al. (2019) |
| 33     | 6UGM     | *M. smegmatis* ESX-3 translocon complex | Poweleit et al. (2019) |
| 34     | 6P6W     | Structure of *M. smegmatis* RNA polymerase core | Kouba et al. (2019) |
| 35     | 7BVC     | *M. smegmatis* arabinosyltransferase EmbA-EmB-AcpM2 in complex with ethambutol | Zhang et al. (2020b) |
| 36     | 7BVG     | *M. smegmatis* arabinosyltransferase EmbA-EmB-AcpM2 in complex with di-arabinose | Zhang et al. (2020b) |
| 37     | 7BX8     | *M. smegmatis* arabinosyltransferase complex EmbB2-AcpM2 in symmetric "resting state" | Zhang et al. (2020a) |
| 38     | 7BWR     | *M. smegmatis* arabinosyltransferase complex EmbB2-AcpM2 in substrate DPA bound asymmetric "active state" | Zhang et al. (2020a) |
| 39     | 6VGQ     | ClpP1P2 complex from *M. tuberculosis* with GLF-CMK bound to ClpP1 | Vahidi et al. (2020) |
| 40     | 6VGK     | ClpP1P2 complex from *M. tuberculosis* | Vahidi et al. (2020) |
| 41     | 6VGN     | ClpP1P2 complex from *M. tuberculosis* bound to ADEP | Vahidi et al. (2020) |
| 42     | 7BVE     | *M. smegmatis* arabinosyltransferase EmbC2-AcpM2 in complex with ethambutol | Zhang et al. (2020b) |
| 43     | 3I83     | *M. tuberculosis* Heptameric EspB Rosetta model | Solomonson et al. (2015) |
| 44     | 5ZEY     | *M. smegmatis* Trans-translation state 70S ribosome | Mishra et al. (2018) |
Recent Insights into the Structure and Function of Mycobacterial Membrane Proteins Facilitated…

Synthase (FAS), F-type ATP synthase, respiratory supercomplex, Caseinolytic proteases (Clp) P1P2 protease, ClpB disaggregate and S-adenosylmethionine (SAM) riboswitch proteins-protein complexes. These proteins-protein complexes are involved in distinct and vital cellular processes like transport, virulence, electron transport, etc., in Mtb and are promising candidates for drug development. These cryo-EM structures have opened new avenues for novel drug discovery programs. In addition, how recent advents in cryo-EM have helped us in gaining insights into the Mtb membrane organization is briefly discussed. Compiled structural information about mycobacterial membrane and membrane proteins deduced using cryo-EM would provide the researchers with a visual picture of reaction mechanisms associated with them and help design novel antituberculars.

Cryo-Electron Tomography and Mycobacterial Membrane Organization

The mycobacterial cell wall is a unique structure that acts as a permeability barrier, and it is essential for survival, virulence, and pathogenesis (Barry Iii et al. 1998; Etienne et al. 2002). Over the decades, researchers have been
trying to elucidate the mycobacterial membrane structure using transmission electron microscopy. However, sample preparation methods that involved fixing with various solvents and staining with heavy metal salts in different labs led to imaging of dehydrated membranes (Daffé and Marrakchi 2019). The information obtained from these studies thus suffered from sample preparation artifacts along with lack of details (Paul and Beveridge 1992; Zuber et al. 2008). Advances in sample preparation methods, specially CEMOVIS (Al-Amoudi et al. 2004; Hoffmann et al. 2008) and cryo-EM of the mycobacterial capsular layer (Sani et al. 2010a), have allowed the visualization of fully hydrated and unstained mycobacterial membrane providing new structural insights into membrane organization. This unique combination of CEMOVIS and cryo-ET of plunge frozen membranes has provided an improved understanding of the structure of lipid bilayer and the outer membrane (Hoffmann et al. 2008).

Summing up the information available, the four-layered envelope of Mtb is about 75–80-nm-thick and can be described from the exterior to the interior of the cell (Daffé and Marrakchi 2019). A capsule layer of ~35 nm thickness is associated with ~8 nm outer membrane made up of mycolipids. The outer membrane is, in turn, linked to a ~35 nm thick cell wall primarily made up of the arabinogalactan layer tethered to the peptidoglycan layer (Alsteens et al. 2008). This is followed by a periplasmic bilogalactan layer tethered to the peptidoglycan layer to a ~35 nm thick cell wall primarily made up of the ara-up of mycolipids. The outer membrane is, in turn, linked to a ~35 nm thick cell wall primarily made up of the arabinogalactan layer tethered to the peptidoglycan layer (Alsteens et al. 2008). This is followed by a periplasmic space and ~7 nm thick granular layer tethered to the cell’s plasma membrane (Lee et al. 2005), where membrane proteins are embedded (Besra and Chatterjee 1994). Interestingly, about thirteen enzymes are involved in synthesizing the arabinogalactan-peptidoglycan complex, which are unique to Mycobacteria (Lee et al. 2005; Crick and Brennan 2008) and hence are putative drug targets against Mtb.

While the understanding of Mtb cell envelops has improved due to their imaging in cryo-conditions, the resolutions achieved are still low. Moreover, it has been shown that lab-grown bacteria may have compositional heterogeneity in the membrane due to the kind of media used (Sani et al. 2010a) or the presence of detergents that may lead to capsule shedding (Hoffmann et al. 2008; Sani et al. 2010a). These problems may be overcome by cryo-EM of vitrified Mtb engulfed by macrophages. However, these are usually too thick to be vitrified properly. The recent developments in sample vitrification methods, high-pressure freezing, and preparation of thin lamella suitable for cryo-EM via FIB-SEM may make it possible (Gorelick et al. 2019; Ravelli et al. 2020). Besides, advances in cryo-ET and the establishment of unique Biosafety Level–3 cryo-EM facilities have raised the hopes of carrying out an exhaustive and detailed structural characterization of native Mtb envelop and membranes (Parvate 2018).

Cryo-EM Structures of Mtb Membrane Proteins

The majority of membrane proteins are recalcitrant to crystallization. Thus, several key Mtb membrane protein structures could not be determined by X-ray crystallography, severely impacting the progress of the Mtb drug discovery field. One of the key issues had been the requirement of milligram quantities of highly pure, compositionally, and conformationally homogenous protein sample for crystallization (Acharya and Lloyd 2005). Moreover, most Mtb membrane proteins express poorly in recombinant expression systems and require extensive optimization of expression and purification conditions (Korepanova et al. 2007). Unlike cytosolic proteins, membrane proteins are highly unstable when extracted from the lipid bilayer (Haltia and Freire 1995). Therefore, to mimic such an environment, detergents/lipids are added to the sample during membrane protein extraction and purification. Due to the presence of lipids/detergents, crystallization becomes more difficult. Technological advancements in cryo-EM, such as the development of direct electron detectors and stable microscopes, have proven beneficial for protein structure determination, especially for membrane proteins (Frank 2017). It overcomes the above two limitations as it requires less amount of sample and does not require crystallization (Renaud et al. 2018). By overcoming these two significant barriers, cryo-EM has accelerated the membrane protein structure determination.

Cryo-EM single-particle analysis technique has led to the structure determination of several essential Mtb proteins providing novel structural and mechanistic insights into their functions. We discuss some of the protein–protein complexes that are therapeutically important targets and whose structures were recently solved in a near-native environment due to current advances in the cryo-EM technique.

Structural Insights into Mycobacterial Secretion Systems and Transporters

ESX/Type VII Secretion System

Bacterial cell membranes contain specific protein complexes used by the pathogenic bacteria to secrete virulence factors helping them to invade the host. Such secretion systems are classified into different types (e.g., Type I, Type II, Type IX) based on their structure, composition, and function (Green and Mecsas 2016; Lasica et al. 2017). Mycobacteria consist of five paralogous ESX (ESX: ESAT-6 Secretion System, ESAT-6: 6 kDa early secretory
antigenic target) secretion systems important for virulence, immune modulation, gene transfer, nutrient and metabolite uptake, and cell physiology of the bacterium (Bunduc et al. 2020). These systems release certain effectors, commonly known as virulence factors, that help the Mtb infect the host. It is proven that the mycobacterial cytosolic translocation is mediated via ESAT-6, a secretion product of the ESX-1 secretion system, and established a connection between the translocation of pathogenic mycobacterium and its virulence (Houben et al. 2012). Although the role of effector proteins from ESX-1, 3, and 5 systems have been traced, the lack of three-dimensional structure of the ESX system had limited the knowledge of its mechanism of operation until the structure of ESX-3 from \textit{M. smegmatis} was solved at ~4 Å via EM (Famelis et al. 2019). During the limited supply of iron, ESX-3 is expressed (Rodriguez et al. 2002; Macikag et al. 2009) and is reported to be associated with various functions such as metal ion homeostasis (Serafini et al. 2009; Siegrist et al. 2009; Tinaztepe et al. 2016; Tufariello et al. 2016), repair of phagosomal damage, and phagosome maturation (Mehra et al. 2013; Portal-Celhay et al. 2016; Mittal et al. 2018).

The ESX-3 secretion system consists of core components called ‘ESC conserved component,’ (Ecc) i.e., Ecc viz; EccB, EccC, EccD, EccE, and fifth component MycP. These components are responsible for releasing virulence factors EsxA and EsxB (Abdallah et al. 2006; Brodin et al. 2006; Ohol et al. 2010; Sloan Siegrist et al. 2014; Ates et al. 2015; Van Winden et al. 2016). The Ecc components together form a hexameric pore in its cell envelope. This pore releases virulence factors that help them infect the host and evade the actions of the host immune system. The membrane-anchored element ‘MycP’ possesses protease activity, while the fork

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**Fig. 1** a1 Structure of EccB dimer from the ESX-3 core complex showing fork architecture at the periplasm (PDB: 6SGY). a2 The ESX-3 core complex structure formed by EccC, EccD, and EccE subunits is arranged laterally. The pale green box is the trans-membrane region. The translocon pore is formed by the EccD subunit (PDB: 6SGW). b The structure of ABC transporter Rv1819c. In the inset, the binding site for the nucleotide (PDB: 6TQE) is shown. The substrate enters from the cap end of the transporter. The conversion of ATP to ADP leads to the import of the incoming substrate from the NBD end (Color figure online)
domain of EccB is towards the periplasmic face (Fig. 1a1). The ESX-3 complex is made up of two protomers. Each protomer constitutes EccB3, EccC3, and EccE3 as a single copy and EccD3 in two copies of homo-dimers. The EccB3 from each protomer is attached to a transmembrane helix (TMH) that anchors it firmly within two protomers and holds the two protomers together with the help of the fork domain (Fig. 1a2). Together, these two protomers form the secretion pore with dimensions large enough to transport dimeric substrate proteins in folded conformation across its cell envelope. Another structure resolved at 3.7 Å, has elaborated the presence of ESX-1-secreted proteins in folded conformation across its cell envelope (Fig. 1a2). Together, these two protomers form the secretion pore with dimensions large enough to transport dimeric substrate proteins in folded conformation across its cell envelope.

The structure of ESX-1 secretion associated protein B (EspB), a secretion protein was also predicted from a cryo-ET, with a resolution of about 30 Å from Mtb (Solomonson et al. 2015). It confers sevenfold symmetry with approximate dimensions of 100 Å×80 Å cylinders with about 50 Å pore. It has been reported that EspB C-terminal domain (CTD) is processed by the membrane-bound protease MycP1P2 following secretion (McLaughlin et al. 2007; Xu et al. 2007; Ohol et al. 2010; Solomonson et al. 2013; Wagner et al. 2013). A visual snapshot of mycobacterial membrane organization using electron microscopy with proteomic confirmation of the presence of ESX-1-secreted proteins has been reported (Sani et al. 2010b). Moreover, an EM structure of ESX-1 secreted Mtb-EspB has been reconstructed at 3.37 Å resolution providing important structural and functional insights. The seven units of EspB form a heptameric cylinder with dimensions of 90 Å×90 Å. The central channel pore is about 45 Å wide. The overall cylindrical arrangement of EspB indicates that it could facilitate the transport of ESX-1 substrates such as EsxA-EsxB or B form DNA molecules (Korotkova et al. 2015). The ESX-3 is proven to be indispensable for Zinc and iron metabolism in Mtb. BBH7, a benzylxybenzylidene-hydrazine compound, has been identified as an inhibitor of the ESX system, which induces Zinc stress in Mtb, thereby severely inhibiting bacterial growth (Rybniker et al. 2014; Cole 2016). Combining the structural information with the knowledge of inhibitors like BBH7, highly specific inhibitors against ESX systems could be designed that may reduce the virulence of Mtb and thus could be an effective alternative antitubercular strategy.

Rv1819c, an ABC Transporter

The ATP-binding cassette transporters (ABC transporters) are probably one of the most studied superfamilies of the multi-subunit transporters, which transport a broad spectrum of molecules at the expense of ATP (Choudhury et al. 2002). The presence of membrane-spanning domains (MSD) and the nucleotide-binding domains (NBD) is typical of ABC transporters. The ATPase activity is associated with the NBD, which is positioned towards the cytoplasm. ATP hydrolysis provides the energy required to transport substrates across membranes (De Rossi et al. 2006). The gene rv1819c was shown to be associated with the uptake/transport of cobalamin in Mycobacteria (Gopinath et al. 2013). However, the protein Rv1819c was related to the transport of antimicrobial peptides like Bleomycin (Domenech et al. 2009) and efflux of antibiotic ciprofloxacin when expressed in C. glutamicum, indicating its potential role in providing resistance through drug efflux (Mazando et al. 2013).

Recently, the structure of the Rv1819c homodimer in the nucleotide bound state was determined at 3.5 Å resolution using cryo-EM (Rempel et al. 2020). The electron density for detergent micelles, observed in the cryo-EM map, helped approximate membrane position around the protein. The structure also revealed the presence of an ABC exporter fold in Rv1819c. The characteristic features in this fold that distinguish it from the similar fold found in previously reported ABC transporters are the unusually large hydrophilic cavity and an additional TMH-0 towards the N-terminal (Korkhov et al. 2012; Choudhury et al. 2014; Lin et al. 2015). The cavity volume measures about 7,700 Å³ spanning almost the entire width of the lipid bilayer membrane. It is made up of two lateral chambers, larger across the membrane, while a smaller one acts as a junction between the larger chamber and the NBD (Fig. 1b). The structures also revealed a 17 amino acid residues long loop in TMH3, constriciting between the two chambers near the membrane towards the cytoplasm. The cavity’s inner side is lined by the network of polar and negatively charged amino acid residues. Such a lining could facilitate the exchange of hydrophilic molecules across the cavity, thus in and out of the membrane. Considering the cavity volume, it could facilitate the transport of antibiotics, too, as seen in transporters of Microcin-J25 export ATP-binding/permease protein McjD and the peptidase-containing ATP-binding cassette transporters (PCAT) (Choudhury et al. 2014; Lin et al. 2015). Further, the structure shows that the exporter fold cavity of the Rv1819c is big enough to accommodate about six cobalamin moieties. This indicates that there is no specific, stable binding site for cobalamin in the cavity. This hypothesis is strengthened with the observation that no substrate was eluted during the co-purification experiments of the Rc1819c. However, the presence of several antimicrobial peptides, including Bleomycin, was confirmed in the cavity. Since Mtb is a slow-growing pathogen, such a large cavity in a non-specific exporter could explain the uptake and accumulation of hydrophilic antibiotics inside the bacterial cell (Taber et al. 1987; Aller et al. 2009). In another study, it was observed that Rv1819c helped the Mtb cells in excreting out the antibiotics, serving as a salvage pathway for Mtb (Balganesh et al. 2010). It would be interesting to see if the Rc1819c could rescue...
the Mtb from antibiotic stress when treated with a specific inhibitor designed using current structural information. If it could do so, specific inhibitors against Rc1819c would prove to be effective antituberculars.

**Proteins Involved in Mycobacterial Cell Wall Synthesis and Maintenance**

**Arabinosyltransferases: Members of Mycobacterial Cell Wall Synthesis Pathway**

As described earlier, the unique cell wall of Mtb has a distinctive structure called mycolyl-arabinogalactan-peptidoglycan. It consists of three essential components: mycolic acids, arabinogalactan, and a cross-linking network of peptidoglycans (Alderwick et al. 2015) synthesized by various transferases and synthases unique to mycobacteria. Recently, cryo-EM enabled the structure determination of some of the key enzymes involved in cell wall biosynthesis. These structures hold particular importance in anti-TB drug development since most of the constituents associated with these enzymes are exclusive to Mtb and thus could act as excellent drug targets. One example is membrane-associated arabinosyltransferases EmbA, EmbB, EmbC enzymes that are members of the mycobacterial cell wall synthesis pathway against which the drug ethambutol (EMB) is considered to be active. EMB is a front-line antitubercular used to treat multi-drug resistant TB (MDR-TB) (Sreevatsan et al. 1997). Interestingly, mutations in these arabinosyltransferases result in drug resistance to EMB (Safi et al. 2008; Sun et al. 2018).

Structures of EmbA-EmbB and EmbC-EmbC complexes in different states, i.e., bound to their glycosyl donor and acceptor, EMB, were determined (Zhang et al. 2020b). These structures help in elucidating mechanisms of EmbA-EmbB and EmbC-EmbC complexes and mode of inhibition by EMB. EmbA and EmbB together form a heterodimer, while EmbC forms a homodimer for its activity. EmbB, a fifteen TMH spanning protein, catalyzes the transfer of arabinose from the donor decaprenyl-phosphate-arabinose (DPA) to its arabinosyl acceptor. On the cytoplasmic face of EmbB is an acyl-carrier protein (AcpM), which is suggested to be associated with the transferase activity. The DPA and di-arabinose bound structures help us understand the overall mechanism of catalysis. The EmbA-EmbB complex catalyzes the branching reaction, while the elongation reaction is catalyzed with the help of the EmbC dimer. In another report, the complex EmbB-EmbB'-AcpM2 in resting-state on binding to DPA gets activated, as shown in Fig. 2a (Zhang et al. 2020a). It leads to the transfer of arabinose from the donor DPA to its arabinosyl acceptor. EMB bound to EmbB structure explains the interaction of the drug with the active site in EmbB (Fig. 2a). The drug EMB competes for the binding sites for substrates in EmbB and EmbC (Goude et al. 2009). The 3.3 Å EM structures of EmbB (substrate and drug bound) provided crucial insights into the mode of action of substrate binding and presented a map of how resistance against ethambutol could have occurred due to mutations accumulated around the active site (Tan et al. 2020a).

Similarly, arabinofuranosyltransferase D (AftD) is another class of transferase. The structure from *M. abscessus* at 2.9 Å resolution (Tan et al. 2020b) revealed similarities in overall architecture with EmbBA, EmbB, and EmbC (Zhang et al. 2020b). The AftD structure is α-helix rich with sixteen TMHs and a cytoplasmic portion consisting of both α and β components. The cytoplasmic α/β folds are homologous to EmbC, three carbohydrate-binding modules (CBMs), and one structural β-sandwich domain. Besides, AftD has a conserved glycosyltransferase (type C) fold, which binds to complex arabinose glycans. The high-resolution structure of AftD enabled the identification of the active site for the transferase activity and highlighted the importance of Arg1389 residue.

**Mycobacterial Fatty Acid Synthase**

Mycolic acid is a significant and unique fatty acid found in the Mtb cell wall playing a vital role in pathogenicity (Marakchi et al. 2014). The enzyme system that catalyzes the synthesis of fatty acid precursors of mycolic acids is known as the FAS complex. In Mtb, there are two types of FAS systems FAS1 and FAS2. FAS1 is found in other mycolic acid producers like certain *Bifidobacteriales* and *Actinomycetales* species and in Mtb, it is known to produce fatty acid precursors with a carbon chain length of C16-18 and C26. However, in *M. smegmatis*, FAS produces C24 fatty acids instead of C26 (Trivedi et al. 2004). FAS2 is involved in the elongation of products of FAS1 in Mycobacteria (Takayama et al. 2005). With the help of other enzymes, FAS2 converts C16-18 long products of FAS1 to C45-64 long meroacids, which are further converted to mycolic acids (Bhatt et al. 2007).

The first mycobacterial FAS1 structure was reported at a global resolution of 7.5 Å from *M. smegmatis*, succeeded by a 3.3 Å structure from Mtb, determined using single-particle cryo-EM (Boehringer et al. 2013; Elad et al. 2018). The FAS1 complex structure appears like a barrel of ~ 25 nm diameter. Even though the resolution is not high, almost all the helices were resolved in the density map. Although separate β-strands were not visible, their presence could be confidently marked due to flattened density clouds corresponding to β-sheets in the MsFAS model. At the same time, these features could be characterized in Mtb FAS1. However, a crucial difference in the reactive pocket of fungal FAS1 and MsFAS is visible. MsFAS at the opening of the reactive pocket is broader than that observed in fungal FAS1 (Gipson et al. 2010). These
The multifunctional, mycobacterial FAS1 is a 2MDa complex consisting of α6 homo-hexamers (Fig. 2b). Each protomer is a product of six subunits named acetyltransferase (AT), enoyl reductase (ER), dehydratase (DH), malonyl transacylase (MPT), ketoacyl reductase (KR), and ketoacyl synthase (KS), assuming respective function as indicated by its name (Fig. 2b). Although considered homologous, *M. smegmatis* FAS1 (MsFAS) is 900 amino acid residues shorter than the fungal FAS1. This difference could be attributed to the presence of the C-terminal phosphopantetheinyl transferase domain in the fungal FAS. This structural data, particularly regions that are different from fungal FAS, can be used for virtual screening and identification of drug candidates that could target mycolate synthesis.

**Electron Transport Chain Proteins are Important Therapeutic Targets**

**F-type ATP Synthase**

The mycobacterial F-type ATP synthase is an essential enzyme among its respiratory chain protein members, which provides ATPs crucial for survival. Encoded by *unc* operon, the ATP synthase consists of six subunits. F1 subunit crystal structure had been reported (Zhang et al. 2019a). However, the complete enzyme structure was lacking, which was recently reported from *M. smegmatis* using cryo-EM (Fig. 3a) (Guo et al. 2020a).

The structure of mycobacterial F-type ATP synthase holds particular importance since the recently approved antitubercular drug BDQ specifically inhibits mycobacterial F-type ATP synthase (Haagsma et al. 2009). It

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*Fig. 2 a Mycobacterial Arabinosyltransferase complex. The complex is composed of two chains, each of EmbB and AcpM. On binding of DPA (red-green spheres), resting-state complex (PDB: 7BX8) is activated (PDB: 7BWR). The arabinosyltransferase is inhibited by ethambutol (Etmb: red) binding in the active site in EmbB (7BVC). b The barrel-shaped MsFAS is a hexamer with one monomer chain shown in cyan (PDB: 6GJC). Each chain is made up of 6 subunits, namely—acetyltransferase (AT), enoyl reductase (ER), dehydratase (DH), malonyl transacylase (MPT), ketoacyl reductase (KR), and ketoacyl synthase (KS) (Color figure online)*
was speculated that it binds either to the c subunit of the enzyme or targets ε subunit (Preiss et al. 2015; Kundu et al. 2016). However, the recent cryo-EM structure describes the full-length enzyme in different drug bound states while resolving all of its subunits. The EM structure remarkably reveals that there are in all 7 BDQ binding sites at F₀ region (Fig. 3b) in the ATP synthase: 5 BDQ molecules exclusively bind to the c chains of the ring called the ‘lagging site’ (Fig. 3c) while the remaining 2 interact with the subunit a and subunit c, called the ‘leading site’ (Fig. 3c). These findings have ruled out the proposed interaction of the BDQ with the ε subunit and have elucidated the overall mechanism and biochemistry of the drug action (Guo et al. 2020b). A recent report also indicates that the drug BDQ can target human mitochondrial ATPase (Luo et al. 2020). This has posed serious concerns regarding the treatment of Multi/Extremely drug resistant (M/XDR) TB. Thus, the availability of the complete structure of mycobacterial ATPase is a significant advancement in designing highly specific inhibitors.

Mycobacterial Respiratory Supercomplex

Several membrane-bound multi-protein enzyme complexes together form the respiratory chain in Mtb (Gong et al. 2018). Popularly termed as respiratory supercomplexes, which help transfer electrons from reduced nicotinamide adenine dinucleotide (NADH) to molecular oxygen (O₂), thus linking this electron transfer to transmembrane proton translocation (Roy and Wani 2020). The net result is the generation of a proton gradient that drives the ATP production and transport across the membrane (Rao et al. 2008).

The M. smegmatis respiratory supercomplex’s (Fig. 4a1) structure was determined at 3.3–3.5 Å resolutions with its crucial cofactors like heme clearly visible in the electron density map (Fig. 4a2) (Wiseman et al. 2018). The complex is primarily made up of two sub-complexes, complex III and complex IV. The dimer of complex III lies in the middle and is flanked by individual complex IV subunits from both sides. The center for oxygen reduction lies in complex IV, while the quinol site is present in complex III. The

![Image](https://example.com/image.png)
electron transfer between these two sub-complexes is facilitated via a cytochrome subunit bridged between complex III and IV. The complex of dimeric $C_{III}-C_{IV}$ arrangement is C2 symmetric with dimensions of $200 \times 70 \times 120$ Å. Such symmetry is the first of its kind observed among respiratory supercomplexes so far. At a global resolution of 3.5 Å, all the prosthetic groups of respiratory supercomplex, menaquinone, and heme molecules could be resolved and built. The two menaquinone molecules could be observed in the conserved quinone binding site located in the $C_{III}$ complex. The complex $C_{III}$ comprises of QcrA, QcrB, and QcrC subunits, while the complex $C_{IV}$ consists of CtaC, CtaD, CtaE, and CtaF subunits. When the supercomplex is in an open state, the cc head of the QcrC is bound to only complex $C_{IV}$, while in the closed state, the head domain is associated with both $C_{III}$ and $C_{IV}$ complexes (Gong et al. 2018).

Mycobacterial $bc$-type $C_{III}$ supercomplex appears similar to the dimeric association, as seen in bacterial $bc_{1}$-type and mitochondrial $C_{III}$; however, certain subtle structural differences are present (Xia et al. 1997). Q203, an imidazopyridine amide, is a candidate reported to be an effective inhibitor of cytochrome $bc_{1}$-aa$_3$ complex, binds explicitly to QcrB of the supercomplex (Lee et al. 2020). Since complex $C_{III}$ is an essential ROS production site, it is a crucial target.
to produce lethal stress in mycobacteria (Zhao and Drlica 2014). The superoxide dismutase (SodC) enzyme is known to protect the electron transport chain (ETC) of mycobacteria from this ROS stress when phagocytized by human macrophages, thus helping in the survival of the bacterium inside the host lungs (D’Orazio et al. 2001). It has been found that the SodC is integral to the respiratory supercomplex situated at the periplasmic face. It is probably involved in scavenging the ROS produced by the host, providing a robust defense mechanism to Mtb (Bhattacharya 2015). In summary, the cryo-EM structure of a C_{III}–C_{IV} respiratory supercomplex from M. smegmatis illustrates how an electron transfer pathway occurs in complex C_{III} and reaches the final oxygen reduction step in the complex C_{IV}, a new branch of electron transfer for energy transduction in association with superoxide dismutase domain acting as the detoxing agent of ROS.

**Structural Insights into Key Mycobacterial Proteases**

**ClpP1P2 Protease**

Proteostasis is a necessary process for maintaining overall protein turnover in the cell using various synthesis and lysis pathways. Specific proteolytic machines in assistance with AAA + unfoldase (AAA: ATPases Associated with diverse cellular Activities) partner recognize certain substrate proteins and mediate their unfolding and further carry out the proteolytic activity (Sauer and Baker 2011). Among such proteases, the ClpP system (Caseinolytic proteases P) is crucial as it helps maintain homeostasis and assist virulence in several bacteria. Particularly in Mtb, the ClpP1P2 protease system is of major importance and indispensable for bacterial growth and virulence (Frees et al. 2014; Bhandari et al. 2018), making it an attractive drug target. Unlike other bacteria, Mtb contains two genes, clpP1 and clpP2, encoding for MtClpP1 and MtClpP2 proteins, respectively (Personne et al. 2013; Alhuwaider and Dougan 2017).

The structures of MtClpP1P2 were recently solved in two conformations; an apo form, which is a compact conformation, and cyclic acyldepsipeptides (ADEPs) bound conformation, which is relatively more relaxed and ordered at the N-terminus. The overall ClpP complex (single ClpP protein 200 residues long) appears like a barrel of about 83 Å height with a central channel formed by two rings stacked one upon another (Fig. 4b). The products of gene clp1 and clp2, i.e., proteins Clp1 and Clp2, form a heptameric ring. The ClpP2 heptamer ring is stacked above that of the ClpP1 ring coaxially to produce a barrel-like structure. In the bacterial ClpP complex, a ‘handle’ region made up of a helix and a β-sheet in each protomer, mediates the interactions within the rings (Liu et al. 2014). The handle region helps the catalytic triad Asp-His-Ser maintain the active site’s geometry for proper binding of the substrate polypeptide (Szyk and Maurizi 2006). The 3.0 Å resolution structure of the apo form of mycobacterial ClpP complex suggests that the β-strands in the handle region are disordered, leading to the distorted catalytic triad. This explains the inactivity of the enzyme system in the apo form. On binding with the ADEP, there is no change in the overall architecture of the ‘handle’ domain (Vahidi et al. 2020).

According to one report, the chloromethyl ketone derivative of compound Bortezomib specifically inhibits the MtClpP1P2 system, imparting cytotoxic effects on Mtb (Moreira et al. 2017). In the same report, it was shown that the inhibitor does not act on human proteases. This structural information of the MtClpP1P2 system could be exploited to design inhibitors highly specific to the protease active site and could be further developed as novel antitubercular candidates like Lassomycin, Rufomycin bind to the ClpC1- an ATP unfoldase, thereby affecting the ATP-dependent protein unfolding from proteolysis, ultimately showing a bacterial effect on active and latent TB infections (Dougan et al. 2020).

**ClpB—an ATP-Driven Disaggregase**

Like the ClpP system described earlier, another protease system, known as ClpB disaggregate, functions in association with heat-shock proteins. In bacteria, ClpB is a ring-shaped ATP-driven disaggregate belonging to the family of Hsp100 chaperones (Doyle et al. 2013). Under stress conditions, the ClpB, in association with other chaperons and accessory proteins, helps specific proteins refold from an aggregated state, thus helping the cell survive and overcome the stress (Mogk 1999; Doyle and Wickner 2009; Lupoli et al. 2016). Like Hsp104, ClpB utilizes ATP hydrolysis energy, and thus, it is known as ‘protein disaggregate,’ where it carries polypeptides through its central pore (Weibezahn et al. 2004). The cryo-EM structure of Mtb ClpB was determined at ~ 3.8–3.9 Å, in different conformations (AMPPNP bound conformer, open and closed state, AMPPNP- Adenylyl imidodiphosphate) (Yu et al. 2018). The structure indicates that ClpB is a hexamer arranged in a ring (Fig. 4c). Each protomer consists of three domains: an N-terminal domain (NTD), and NBD1 and NBD2 as two AAA+ type nucleotide domains that carry out ATP assisted peptide translocation (Gates et al. 2017). Thus, each hexamer accommodates 12 NBDs. Eukaryotic Hsp104 and mycobacterial ClpB belong to the same family (Doyle et al. 2013). Still, there are key differences, especially in the NBD regions of these two proteins. The Mtb ClpB has both NBD1 and NBD2 moving in an upward direction. However, in Hsp104, the two NBDs move in opposite directions, i.e., if NBD1 moves upwards, then the NBD2 orients itself downwards. This movement creates a temporary gap in the Hsp104 ring, which is absent
in ClpB. This structural difference could probably reflect a different mode of ATP hydrolysis in ClpB and Hsp104. Further experiments, however, are warranted to confirm this observation. ClpB is crucial for Mtb survival inside the host (Lupoli et al. 2018). The insights gained from the ClpB structure could help in developing it into a novel anti-TB drug target.

**Cryo-EM Structures that Elucidate Basic Cellular Processes in Mtb**

Apart from the proteins mentioned above, a few other crucial structures were reported recently. These structures are essential considering their druggability and have provided us with a more in-depth understanding of mycobacteria’s underlying cellular processes. The list includes Mtb and *M. smegmatis* ribosomes, small heat-shock protein Acr1, proteasome assembly and its bacterial proteasome activator (Bpa), nuclease AdnAB motor, RNA polymerase, and few others (Table 1). The structure of α-crystallin related protein 1 is among the first few mycobacterial protein structures determined using cryo-EM (Kennaway et al. 2005).

The cryo-EM structures of ribosomes describe how Mtb remodels its ribosomes in Zinc-depleted conditions (Li et al. 2018a). The 70S structure has underlined the differences between Mtb and other bacterial ribosomes, especially the variations in lengths of specific rRNA helices (Li et al. 2018b). The 70S structure now enables us to understand the movement between specific ribosomal proteins, handle, small subunit, and other regions/domains of the complete ribosome assembly (Yang et al. 2017). These structures offer detailed information about the transcription occurring at the ribosome. Additionally, two non-ribosomal proteins were detected in these structures. These two proteins could be some transcription regulators; however, these are yet to be identified. The EM reconstructions of the *M. smegmatis* 70S ribosomes in the hibernating state, trans- translating state (Fig. 5a), and the peptidyl tRNA bound state at the resolutions 4.1 Å, 12.5 Å, and 3.4 Å, respectively, highlight the new role of bS1 protein in 70S protection during hibernation in mycobacteria. With these structures’ help, we now have a complete snapshot of H54a-bS1 interactions that prevent subunit dissociation and degradation (Mishra et al. 2018).

Since, using cryo-EM, protein structures could be captured in various intermediate steps of transcription, they improve the understanding of the underlying conformational transitions. The available EM structures of mycobacterial RNA polymerase in core and holoenzyme containing σA—a principal sigma factor form have filled this gap (Kouba et al. 2019). These structures depict the steps where the promoter partially melts within the RNA polymerase cleft. This leads to the restriction of DNA at the RNA polymerase active site. The structures of RNA polymerase-promoter DNA complex intermediates elaborate on how and why clamp opening is essential to open a passage for the entry of single-stranded DNA into the active pocket, including fork-loop 2 and switch 2 structures in promoter melting (Boyaci et al. 2019). This RNA polymerase is arrested by antibiotic Fidaxomicin, in an open state, i.e., Fidaxomicin on binding in the cleft of RNA polymerase when in an open state (Fig. 5b) halts the transcription by stopping the promoter from interacting with the RNA polymerase (Boyaci et al. 2018; Lin et al. 2018). Alongside transcription regulation, cryo-EM also sheds light on other molecular machines such as those involved in homologous recombination. AdnAB is a helicase-nuclease present in mycobacteria. It is a dimeric complex with a C-terminal nuclease domain that creates cleavage of 5’ strand DNA. The structures of AdnB in the absence of DNA and complex with forked duplex DNAs before and after cleavage of 5’ ssDNA describe the process of DNA double-strand break resection in *M. smegmatis* (Fig. 5c) (Jia et al. 2019).

Proteasomes are the assemblies of proteins that perform regulated proteolysis at times required by the cell (Becker and Darwin 2017). For example, the mycobacterial proteasome is activated by proteasome accessory factor E (PafE) (Fig. 5d). This activation promotes proteolysis through proteasome, helping in the virulence of Mtb in the human host. PafE binds to proteasome via glycine–glutamine–tyrosine–leucine (GQYL) motif at its C terminus, independent of ATP. This binding stimulates proteolysis (Hu et al. 2018). Like PafE, Bpa is another molecule that activates the 20S proteasome with the GQYL motif’s help (Bolten et al. 2016).

**S-Adenosylmethionine IV Riboswitch RNA**

In bacteria, specific RNAs can regulate downstream gene expression when bound to cellular metabolites, especially in response to a change in those particular metabolites’ concentration. Such RNAs are known as riboswitches. Even though riboswitch RNA does not fall into the membrane protein category, its structure holds particular importance. Despite its small size of 40 kDa, a relatively high-resolution EM structure was determined. These metabolite-binding riboswitches are highly specific but non-coding in nature. The metabolite S-adenosylmethionine (SAM), when acting as a ligand for riboswitches, regulates sulfur metabolism in bacteria (Roth and Breaker 2009).

SAM acts as a cofactor in several methylation reactions, wherein it particularly recognizes seven types of riboswitches like SAM-I, SAM-II... to SAM VI, and SAM-I/IV (Mirihana Arachchilage et al. 2018). SAM-I, SAM-I/IV, and SAM-IV are commonly found SAM riboswitches; crystal structures are already available for them except SAM-IV (Montange and Batey 2006; Gilbert et al. 2008;
Lu et al. 2008; Trausch et al. 2014; Huang and Lilley 2018). Recently, the SAM-IV riboswitch cryo-EM structures in both apo and SAM-bound forms were successfully determined at an overall resolution of 3.7 Å and 4.1 Å, respectively (Zhang et al. 2019b). SAM-IV riboswitch comprises of P1, P2a, P2b, P3, and P5 as five helices with two ‘pseudoknot’ (PK) folds, PK-1 and PK-2. The comparison of SAM-IV structure with that of SAM-I marks critical differences between these two structures. The SAM-IV contains a pseudoknot PK2, which is not seen in SAM-I but lacks the P4 helix present in SAM-I. The kink-turn motif in SAM-I and IV contains six and four nucleotides, respectively (Montange and Batey 2006). It can be noticed that the SAM-I/IV riboswitch contains the P4 helix but lacks PK1 pseudoknot. However, the P4 helix is present in the SAM-IV riboswitch. Despite all these significant structural differences among these three classes of riboswitches (SAM-I, SAM-I/IV, and SAM-IV), the overall core topology at the SAM binding site in these riboswitches remains the same.

**Additional Cryo-EM Structures that could be Novel Drug Targets**

Despite the recent progress in structure determination of Mtb proteins, there are still several proteins that remain to be explored not just for drug discovery purposes but also
to delineate the mechanism of action of crucial enzymes involved in different metabolic processes.

Phenolphthiocerol/phthiocerol polyketide synthase subunit E (PpsE), a type I polyketide synthase, is involved in the elongation of C22-24 fatty acids by adding malonyl-CoA and methylmalonyl-CoA extender units. This elongation step yields phthiocerol derivatives (Astarie-Dequeker et al. 2009). In Mtb, PpsE is proven to be associated with the ESX-1 secretion system (Augenstreich et al. 2017), thus making it an exciting target. The msl3 gene product, mycolic panoate synthase, is a homodimer. It is a part of the polyacylated trehalose synthesis pathway. Mutations in this gene lead to cell aggregation (Dubey et al., 2002).

Another candidate protein, Lnt, is 93 kDa in size and catalyzes the N-acetylation of the N-terminal cysteine of apolipoprotein (Tschumi et al. 2009). Specific inhibitors of Lnt could be designed based on the structural information to study the after-effects of tri-acetylation inhibition in Mtb cell morphology. Apart from such membrane proteins, other relatively smaller proteins like iron import ATP-binding protein, Serine/Threonine-protein kinase, Prothample IX farnesytransferase should be targeted for structure determination since these hold great potential as novel drug targets (Nalamolu et al. 2019).

**Cryo-EM and Vaccine Development**

‘Structural vaccinology’ is a rational approach for vaccine development. It is primarily based on determining the atomic structure of the antigen or antigen–antibody complex, mainly against viruses (Anasir and Poh 2019). The cryo-EM technique facilitated the structure determination of giant viruses like the herpes simplex virus for vaccine development. (Dai and Zhou 2018). Based on a similar approach, vaccines could be developed against different viruses. Recent EM structures of the African swine fever virus (ASFV)—one of the most complex DNA viruses could help in a more in-depth understanding of the virus’s structure–function relationship (Gallagher and Harris 2020). Such progress should aid vaccine development and could form the basis of a new era of ‘Structural Vaccinology.’ The recent SARS-CoV-2 pandemic, several structures have been deposited that have proven instrumental in research and development on drug discovery and vaccine development for SARS-CoV-2 (Walsh et al. 2020). Similarly, in TB’s case, structural information from the above discussed Mtb proteins could be utilized to develop an effective treatment/vaccine against all TB types towards worldwide TB control by 2030 as per WHO’s goals.

At present, 14 TB vaccines are at different stages of clinical trials, while the live, attenuated *Mycobacterium bovis* bacille Calmette-Guérin (BCG) vaccine is still the most widely used (Schrager et al. 2020). With the amalgamation of available cryo-EM information and additional experiments to understand mycobacterial membrane using EM and tomography techniques, improved and effective vaccine alternatives to the classical BCG vaccine may be an achievable goal.

**Concluding Remarks**

As mentioned earlier, membrane protein structure determination is a cumbersome process since membrane proteins are difficult to crystallize. However, in the past decade, cryo-EM technology has taken a giant leap forward, making these limitations almost invisible. Since 2005, about 75 EM structures have been deposited (as updated on July 2020) from mycobacteria in the PDB. We have briefly discussed key membrane proteins that are therapeutically important. These structures include protein candidates from various biological pathways. Some are associated with cell growth and metabolic pathways, while others are associated with virulence, defense, and survival strategies. The snapshots provided by cryo-EM structures depicting proteins in various stages of their activity help us establish underlying mechanisms with structural evidence. Several structures of ribosomes help us visualize different steps in transcription. Not only membrane protein structures, we now have snapshots of membrane organization, which would prove instrumental in the expansion of knowledge of material transport across the mycomembrane, especially when mechanistic insights into the functioning of secretion systems and transporters have started to emerge, thanks to cryo-EM.

In recent years, we have witnessed a tremendous increase in structure determination using cryo-EM. Compared to the past two decades, we see exponential growth in the number of membrane protein structures determined using EM in the past three years (Supplementary Table S1). These membrane proteins are associated with crucial pathways involved in bacterial metabolism and/or virulence; at the same time, they are non-native to human physiology. Hence, biosynthesis or metabolic pathways associated with these proteins could serve as a goldmine of novel and effective anti-TB drug targets.

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Declarations

Conflict of interest PJP is co-founder and shareholder of www.cryosol-world.com.

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