# Attack on the Scourge of Tuberculosis: Patented Drug Targets

Reena Vohra, Meetu Gupta, Rekha Chaturvedi and Yogendra Singh\*

Institute of Genomics and Integrative Biology, Mall Road, Delhi 110007, India

Received: July 22, 2005; Accepted: September 13, 2005; Revised: November 06, 2005

**Abstract:** Tuberculosis is one of the most devastating bacterial diseases, with increasing rates of morbidity and mortality, despite the presence of effective chemotherapy and Bacillus-Calmette-Guerin (BCG) vaccine. The success of *Mycobacterium tuberculosis* lies in its ability to spread by aerosol droplets, evade the host immune system and to persist in pulmonary granulomas. The advancement in the field of molecular and cellular microbiology and the availability of transcriptome and proteome data of *M. tuberculosis* have aided in understanding the pathogenesis of this organism for developing more effective drugs. The current strategy of drug design is to identify gene products, which are essential for survival and virulence. To date, several gene products of mycobacteria, ranging from proteins involved in cell wall synthesis to energy generation and from entry into host to persistence, have been shown to be essential for the survival or virulence of *M. tuberculosis*. These proteins and their associated pathways are considered as promising drug targets against *M. tuberculosis* and several of these have been patent protected. Herein, we enlist drug targets against *M. tuberculosis* for which patents have been filed and issued during the last ten years. The significance of these drug targets in the development of drug is also discussed. This review presents a comprehensive account of the pivotal information for drug discovery and drug design to all researchers involved in tuberculosis research.

**Keywords:** *Mycobacterium tuberculosis*, tuberculosis, drugs, cell envelope, MabA, sigma factor, RNA polymerase, signal transduction, serine/threonine kinases, metabolism, virulence, isocitrate lyase, dormancy.

#### INTRODUCTION

Tuberculosis, caused by Mycobacterium tuberculosis, remains one of the biggest killers amongst the infectious diseases despite the availability of effective drugs and an attenuated Bacillus-Calmette Guerin (BCG) vaccine. Streptomycin was the first drug introduced in 1944 for the treatment of tuberculosis but almost immediately after its introduction many patients started showing resistance to this antibiotic [1-3]. Para-aminosalicylate (PAS) was introduced in 1946 that largely overcame the emergence of resistant strains [4]. A few years later, isoniazide (INH) was developed and initial treatment with both INH and streptomycin was even more effective. To date, many drugs are available, which are classified into two categories. First line therapy includes five medications: isoniazide (isonicotinic acid hydrazide), pyrazinamide (analog of nicotinamide), ethambutol [(S,S´)-2,2´(ethylenediimino)di-1-butanol], rifampicin (lipophilic ansamycine) and streptomycin (aminocyclitol glycoside) [5, 6]. Second line therapy, which is used exceptionally in the cases of drug resistance, includes cycloserine, capreomycin, fluoroquinolones, ethionamide, PAS, thioacetazone, rifabutin, clofazimine and some macrolides [7].

The major setback in controlling tuberculosis was the emergence of multidrug resistant tuberculosis (MDR-TB) during 1990-92. Currently, at least 50 million people are estimated to be affected with MDR-TB. A few MDR strains of *M. tuberculosis* were found to be resistant to many first line agents as well as some of the second line drugs [8]. Moreover, the high rate of coinfection with human

immunodeficiency virus (HIV) presented a challenge to the existing chemotherapies. One of the limitations of currently available drugs is their inability to act on latent bacilli. People carrying latent infection are at a risk of reactivation and this is one of the major barriers in controlling tuberculosis. Therefore, there is an urgent need to develop novel drugs that can act against both actively growing and dormant bacteria. The efforts for drug development are being coordinated by Global Alliance for TB Drug Development (www.tballiance.org), an organization that has been involved in developing public-private partnerships to bring out new, faster-acting and affordable drugs against tuberculosis.

Rational development of a new antitubercular agent requires the exploration of new means to understand the genetics and physiology of M. tuberculosis. In this regard, availability of the genome sequence of M. tuberculosis [9] and powerful genetic tools for manipulating mycobacteria have provided valuable information about the potential targets. Also, the information available from X-ray crystallographic studies of many of these targets has helped in designing novel chemotherapeutic agents. This review exclusively focuses on mycobacterial targets that have been patent protected from all over the world in the last ten years, Fig. (1). Herein, we present the information vis-à-vis proteins that are important for the survival of *M. tuberculosis* during infection and persistence in the host environment. The potential drug targets compiled in this review are likely to lead to new medication that should facilitate in controlling the spread of tuberculosis.

### PATENT PROTECTED DRUG TARGETS

A patent is a set of exclusive territorial rights granted by the government of a country to the assignee for their creative ideas and expressions of the human mind that have

<sup>\*</sup>Address correspondence to this author at the Institute of Genomics and Integrative Biology, Mall Road, Near Jubilee Hall, Delhi 110007, India; Tel: 011-2766 6157; Fax: 011-2766 7471; E-mail: ysingh@igib.res.in

Fig. (1). An outline of Patent Protected drug targets of Mycobacterium tuberculosis.

The patented targets against which antitubercular drugs can be developed are shown in boxes. These targets are known to control various cellular processes required for survival or virulence of *M. tuberculosis*. The site of action of first line drugs (Rifampicin, Isoniazid, Pyrazinamide, Ethambutol and Streptomycin) are shown by arrows.

commercial value. At present, several gene products, which are part of many important pathways such as cellular metabolism, transcription, translation, cell envelope synthesis, signal transduction, persistence and virulence have been patented owing to their importance as promising drug targets (Table 1).

# PROTEINS INVOLVED IN CELL ENVELOPE METABOLISM

Mycobacteria belong to a suprageneric taxon, the mycolata, which also includes Corynebacteria, Nocardiae and Rhodococci, by virtue of similarity in their cell wall architecture [10]. The mycobacterial cell envelope is a complex structure comprising mycolic acids covalently bound to peptidoglycan via arabinogalactan [11]. The cell wall of *M. tuberculosis* confers resistance against many antibacterial agents known to inhibit cell wall synthesis such as isoniazide, ethionamide, isoxyl, and the thiosemicarbazones. An enriched knowledge of the biosynthesis and assembly of the cell envelope components has provided vital clues for the development of novel antimycobacterial agents.

After phagocytosis by macrophages, M. tuberculosis overcomes several environmental stresses, such as those caused by toxic reactive oxygen and nitrogen species. The detrimental effect of the reactive radicals is circumvented by modifications such as cyclopropanation of mycolic acids, which renders mycobacterium resistant to lipid peroxidation [11]. Mycolic acid cyclopropanating enzyme (MACE, US5573915) [12], also known as cyclopropane mycolic acid synthase (cma) is responsible for cyclopropanation at distal position of the mero chain in the -mycolic acid series [13]. Non-pathogenic mycobacteria, M. smegmatis do not cyclopropanate their mycolic acids. Heterologous expression of cma in M. smegmatis not only resulted in cyclopropana-tion of -mycolates but also significantly protected recombinant M. smegmatis from oxidative stress [13]. A homolog of cma is also present in M. leprae, suggesting that cyclopropanation of mycolic acids is specific to pathogenic forms of mycobacteria [14]. Therefore, therapeutic agents, which affect the activity of MACE, hold the key to successful elimination of pathogenic mycobacteria. PCT patent numbers WO9638581 and US5573915 issued

 Table 1.
 Patent Protected Drug Targets of M. tuberculosis

S. No.	DRUG TARGET	CLASSIFICATION	PATENT NUMBER <sup>a</sup>	PDB ID
1	Arylamine N-acetyl tranferases	Cellular Metabolism	EP1082441, EP2002516109T, WO9961625, GB9811407	1GX3
2	Proteasome associated genes and DNA repair genes	Cellular Metabolism	US2004213776	-
3	Acetolactate synthase and ketol acid reducto isomerase	Cellular Metabolism	AU2451397, US5998420, WO9737660	-
4	Thymidylate synthase	Cellular Metabolism	WO02072805, GB0105763D	-
5	SigA and SigB	Transcription	AU706367, AU4550896, AU5018496, CA2171600, CN1201465, EP0832116, GB2298862, IE960132, JP2000503525, NZ286081, WO9638478, SE9501976, SE9502596, SE9503246,	-
6	SigF, OrfX, and OrfY	Transcription	AU732858, AU2580297, CA2249208, EP0910403, JP2000508525T, US5700925, US5824546, US6004764, WO9735611	-
7	AlgU	Transcription	CA2278314, EP0970192, JP200150903, US6355469, WO9831789	-
8	RNA Polymerase alpha-subunit	Transcription	AU7625298, CA2274098, EP0956347, EP2001505438, US6355464, WO9824891	-
9	WhiB1, WhiB2, WhiB3 and WhiB4	Transcription	EP1245683, US6645505, US2002164573	-
10	Adenylyl cyclase	Signal Transduction	CA2447110, EP1258528, EP1390487, US2005048603, WO02092805	Crystallization done, No PDB ID available <sup>b</sup>
11	Protein kinase B, G, H and J	Signal Transduction	AU2003206594, WO03074728	1MRU
12	Tyrosine phosphatases	Signal Transduction	WO2005005639	1U2Q
13	DevR	Signal Transduction	EP1472339, GB2398302, WO03066838	-
14	Isoleucyl tRNA synthase	Translation	US5756327	-
15	Methionyl tRNA synthase	Translation	US5798240	-
16	Seryl tRNA synthase	Translation	US5656470	-
17	MabA (FabG1)	Cell Envelope	AU2003258723, EP1490491, FR2837836, WO03082911	1UZN
18	EmbC, EmbA, and EmbB	Cell Envelope	AU6467598, US6015890, WO9841533	-
19	MACE	Cell Envelope	AU5881496, US5573915, WO9638581	1L1E
20	D-alanine racemase	Cell Envelope	US2003133952	-

(Table 1) Contd....

S. No.	DRUG TARGET	CLASSIFICATION	PATENT NUMBER <sup>a</sup>	PDB ID
21	Rv3133c, Rv2623, Rv2626c	Dormancy Related Genes	AU1610002, GB0030368, GB2386420, US2004242471, WO0248391	-
22	Resuscitation Factor (RpfA-E)	Dormancy Related Genes	AU749414, AU7779798, CA2292898, EP0983361, GB9711389, GB9811221, JP2002503106, NZ501568, WO9855624, ZA9804838	1XSF
23	Rv1174c and phospholipids	Dormancy Related Genes	WO0245736	-
24	Isocitrate lyase	Dormancy Related Genes	AU3391602, US2003018166, WO0233118	1F8M, 1F8I
25	Alkyl hydroperoxide reductase (AhpC)	Virulence	WO9954479	2BMX
26	AhpD, dihydrolipoamide dehydrogenase, and dihydrolipoamide succinyltransferase	Virulence	US2003190325	1KNC
27	SecA	Virulence	AU5170896, US5885828, WO9626276	1NL3, 1NKT

<sup>&</sup>lt;sup>a</sup> The first two letters in the patent number corresponds to PCT (Patent Corporation Treaty) contracting states: AU, Australia; AT, Austria; CA, Canada; CH, Switzerland; CN, China; DE, Germany; FR, France; EP, Europe; GB, Great Britain; IE, Ireland; JP, Japan; NZ, New Zealand; SE, Sweden; TW, Taiwan; US, United States of America; WO, World Intellectual Property Organization; ZA, South Africa.

bwww.doe-mbi.ucla.edu/TB/

1996, relate to MACE and its DNA or protein compositions useful for both diagnosis and designing of therapeutics for treatment of tuberculosis and other mycobacterial infections. The patent discloses the invention of a method for determining the ability of a compound to inhibit the cyclopropanation of mycolic acids in pathogenic mycobacteria.

MabA, a NADPH-dependent -ketoacyl ACP reductase, is involved in the biosynthesis of long chain fatty acids, the precursors of mycolic acids (WO03082911) [15]. Unlike mabA of M. smegmatis, mabA of M. tuberculosis is in an operon with inhA, an NADH-dependent 2-trans-enoyl ACP reductase [16]. Both genes together with KasA/B ( -ketoacyl synthase) form type II fatty acid elongation system (FAS-II) [17-19]. INH is known to inhibit InhA and MabA by formation of a covalent adduct between Mn<sup>III</sup>-activated isoniazid and the MabA/InhA cofactor [20]. Crystal structure analysis revealed that MabA has specific functional and structural properties when compared to other homologous bacterial -ketoacyl reductases, such as large hydrophobic substrate binding pocket and preference for long chain substrates [21]. These distinct properties of MabA make it a potential drug target specific for mycobacteria. Also, the mechanism of inhibition by a metabolite of INH could serve as a model for rational drug designing. WO03082911 published in 2004 relates to the recombinant native and mutant protein MabA. The invention provides crystallographic co-ordinates for designing and screening ligands inhibiting the enzymatic activity of MabA.

Ethambutol (EMB) is a frontline antituberculosis drug, which targets the mycobacterial cell wall. Resistance to EMB was used as a tool to identify genes participating in cell wall biosynthesis, which led to the identification of

mycobacterial embCAB gene cluster (US6015890) comprising embC, embA and embB genes [22-24]. EmbA and are arabinosyl transferases, which utilize arabinofuranosyl phosphodecaprenol for arabinosylation of cell wall arabinogalactan, the major polysaccharide of the mycobacterial cell wall [25-27]. Over expression of Emb proteins [23] and mutations of conserved residue Met306 are associated with resistance to EMB in mycobacteria [22, 23, 28]. Identification and characterization of resistance to EMB has provided information for development of new chemotherapeutic agents against these mutated Emb proteins. WO9841533 published in 1998 and US6015890 issued in 2000 relate to the identification, cloning, sequencing and characterization of the embCAB operon. This patent also provides one or more single-stranded nucleic acid probes, which specifically hybridize to the wild type embCAB operon or the mutated embCAB operon that may be used in the diagnosis of drug-resistant mycobacterial strains.

D-Alanine is an essential component of dipeptide D-alanyl-D-alanine, involved in the cross-linking of peptidoglycan strands in bacteria [29]. D-alanine racemase (alrA) catalyzes the conversion of L-alanine into D-alanine [30], and D-alanine-D-alanine ligase catalyzes the subsequent dimerization of D-alanine into the key dipeptide, D-alanyl-D-alanine [31]. Inactivation of D-alanine racemase was shown to affect the survival of *M. smegmatis* in phagocytic cells (US2003133952) [32]. Thus, inhibitors can be designed against D-alanine racemase as novel antituberculosis drugs targeting peptidoglycan biosynthesis in mycobacteria. US2003133952 published in 2003 relates to methods of making live-attenuated vaccines against pathogenic mycobacteria using *alrA* mutants. Furthermore, *alrA* mutants

can be used for screening antimycobacterial agents that are synergistic with peptidoglycan inhibitors.

### PROTEINS EMPLOYED IN TRANSCRIPTION AND TRANSLATION

Genes required for survival following uptake by macrophages can provide insight into mycobacterial pathogenesis and provide novel targets for developing antibacterial agents. The ability to adapt to intracellular stress requires regulation of complex gene expression mediated mainly by sigma factors. Sigma factors are involved in transcription initiation by interacting with RNA polymerase. DNA-dependent RNA polymerase is a basic unit of the bacterial transcription apparatus. The holoenzyme is a complex consisting of five protein subunits: two copies of subunit and one copy each of , 'and sigma subunit. ' subunits are invariant in a given bacterial species and together with subunit form core RNA subunit is involved in protein-protein polymerase. The interactions with transcription activators and protein-DNA interactions [33-35]. The amino terminal domain of subunit is also required for the assembly of core RNA polymerase. RNA polymerase is a well-documented drug target and rifampicin is a highly specific inhibitor of mycobacterium RNA polymerase [36]. Due to emergence of resistant strains, development of new compounds that interfere with the enzymatic activity of RNA polymerase or disrupt interaction of a subunit with the core enzyme is urgently required. Healy et al., have patented the method for high-throughput screening of RNA polymerase inhibitors and identified anti-tuberculosis compounds that specifically inhibit mycobacterial transcription (EP0956347) [37]. Patent applications WO9824891 published in 1998, EP0956347 and US6355464 issued in 2002 relate to novel nucleic acids encoding the RNA polymerase alpha subunit from M. tuberculosis. The invention also provides vectors comprising the nucleic acids, cells comprising the vectors, and methods for producing M. tuberculosis alpha subunit.

Sigma factors are interchangeable RNA polymerase subunits that are responsible for promoter recognition. Prokaryotes usually have a constitutively expressed principal sigma factor, which is responsible for the transcription of essential housekeeping genes, and a number of alternative sigma factors that are transcriptionally and/or posttranslationally activated in response to specific environmental signals [38]. Mycobacterial genome encodes at least 14 sigma factors [9]. Sigma factor A (sigA; rpoV), (WO9638478) [39], a principal sigma factor, is an essential gene and has been shown to be involved in virulence of M. tuberculosis [40]. A mutation in sigA from arginine to histidine at amino acid residue 522 resulted in the loss of virulence of M. bovis and complementation of the attenuated M. bovis mutant with the M. tuberculosis wild type sigA restored virulence [40]. The transcription of sigB (WO9638478) [39] is regulated differentially from that of sigA. Transcription of the sigB gene increases significantly when *M. tuberculosis* enters the stationary phase at 10 days of microaerophilic incubation and under various stress conditions [41]. These findings suggest that SigB may be an alternative or secondary sigma factor, which controls a large stationary-phase regulon.

WO9638478 published in 1996 and GB2298862 issued in 1999 relate to the provision of novel nucleic acid molecules coding for SigA and SigB subunits of M. tuberculosis RNA polymerase. The invention further provides screening assays for compounds, which inhibit the interaction between a sigma subunit and a core RNA polymerase.

sigH, (AlgU, US6355469) [42] was shown to be induced after heat shock and after exposure to the thiol-specific oxidizing agent, diamide [43]. M. tuberculosis mutants lacking sigH showed reduced immunopathology in infected animals [44]. M. tuberculosis sigF mutant strain was shown to be attenuated in immunocompetent mice [45, 46] and moreover, the expression level of sigF is upregulated in stationary phase and in stress conditions [47]. SigF is regulated by M. tuberculosis OrfX (anti-sigma factor) and OrfY (anti-anti-sigma factor) proteins. The M. tuberculosis SigF, OrfX and OrfY (WO9735611) [48] can be used to screen for dormancy activators, which trigger growth arrest during active tuberculosis infection and can also be used to screen for antagonists that induce reactivation in patients with latent tuberculosis. Reactivation will render antimycobacterial drugs more effective, because the available drugs are typically more potent toward actively growing bacilli. Thus, sigma factors are logical targets for the development of the transcriptional inhibitors. US6355469 patent issued in 2002 and WO9831789 published in 1998 relate to DNA encoding M. tuberculosis RNA polymerase AlgU sigma subunit protein and methods of detecting inhibitors of the RNA polymerase. The invention also encompasses sequence-conservative and function-conservative variants of this sequence. US5700925 patent issued in 1997 is directed to a gene involved in latency and a diagnostic method for detecting latent M. tuberculosis. The invention also relates to M. tuberculosis vaccines expressing mutant sigF genes. Patent application WO9735611 published in 1997 and AU732858 patent issued in 2001 disclose a method for identifying a gene or a protein, which is regulated by a sigma factor of M. tuberculosis.

WhiB proteins were originally described in Streptomyces coelicolor and were shown to be involved in sporulation and cell septation [49]. WhiB family proteins are also present in M. tuberculosis and M. bovis BCG. WhiB proteins (WhiB1, WhiB2, WhiB3 and WhiB4, US6645505) [50] of M. tuberculosis have been patented and disruption of these genes in M. bovis BCG revealed that whiB1 is an essential gene and whiB2, whiB3 and whiB4 are involved in growth and septum formation. Furthermore, WhiB proteins were characterized by the yeast two-hybrid system that demonstrated DNA binding and transcriptional activation properties of these proteins [50]. WhiB3 has also been shown to be involved in virulence of *M. tuberculosis* as gene inactivation led to reduction in the survival in immunocompetent mice [51]. Patent number US6645505 issued in 2003 discloses an in vivo drug screening method taking advantage of the yeast two-hybrid and provides a method of using whiB genes. The drugs against WhiB2 and the WhiB4 of M. tuberculosis H<sub>37</sub>Rv and M. bovis BCG will be particularly useful where drug resistance has developed against the WhiB1 and WhiB3 or where the anti-WhiB1 and anti-WhiB3 drugs are allergic or toxic.

The components of the translational apparatus are prominent targets for antibiotics. Aminoacyl-tRNA synthetases catalyze the transfer of specific amino acid to its corresponding tRNA to form aminoacyl-tRNAs, which are used during protein synthesis. Eukaryotic-like isoleucyltRNA synthetase (IleRS, US5756327) of M. tuberculosis [52, 53] has been found to be resistant to the prokaryotic IleRS-targeted antibiotic, pseudomonic acid [54, 55]. Moreover, M. tuberculosis methionyl-tRNA synthetase (MetRS, US5798240) [56] has characteristic class I signature sequences (HVGH and KMSKS) but lacks the Zn<sup>2+</sup> binding motif and the C-terminal dimerization appendix [57]. Because the amino acid sequences of the tRNA synthetases have diverged with time, significant differences exist between the structures of the enzymes from mammals and mammalian pathogens [58]. These differences can be exploited by identifying inhibitors, which have specific activity against these mycobacterial tRNA synthetases. Moreover, the seryl-tRNA synthetase (US5656470) [59] of M. tuberculosis has also been characterized by biochemical assays and antisense strategy. US5756327 patent issued in 1998 relates to IleRS [52]. Recombinant DNA constructs encoding IleRS can be used for the construction of tester strains to identify inhibitors of the essential tRNA synthetase. US5798240 patent issued in 1998 relates to the isolated and/or recombinant nucleic acids, which encode MetRS [56]. The invention also provides antisense nucleic acid that can hybridize to the mRNA of MetRS of mycobacteria. US5656470 patent issued in 1997 relates to isolated and/or recombinant nucleic acids, which encode seryl-tRNA synthetases of mycobacterial origin [59].

# PROTEINS OF SIGNAL TRANSDUCTION PATHWAYS

After entering macrophages, cross-talk modalities exist between the mammalian host and the pathogen, with the result that the host's defense system is bypassed and the pathogen survives and proliferates. M. tuberculosis genome encodes 15 putative adenylyl cyclases (ACs), indicating the presence of signal transduction pathways mediated through cAMP as a second messenger. It has also been reported that cAMP levels increase in macrophages infected with mycobacteria leading to an inhibition of phagosomelysosome fusion [60]. Amongst the 15 putative ACs, Rv2435c and Rv1625c are grouped with mammalian ACs based on their sequence homology (WO02092805 and US2005048603) [61, 62]. As the closest progenitor of mammalian AC, Rv1625c gene product comprises a protein with six transmembrane helices and a single cytosolic catalytic domain, which dimerizes to form a 12transmembrane, homodimeric enzyme, with two substratebinding sites at the dimer interface in contrast to the heterodimeric mammalian enzyme [63]. Structural and biochemical analysis revealed that despite the high (60% similarity to guanylyl cyclases) sequence similarity to the mammalian enzymes, Rv1625c has a unique substrate binding pocket that has not been reported in any other cyclase so far [63, 64]. These differences could in principle be exploited in designing novel drugs. US2005048603 published in 2005 and WO02092805 published in 2002 relate to a method for expression of membrane proteins wherein a portion of a nucleotide sequence coding for the membrane protein is fused to a portion of a nucleotide sequence coding for an adenylyl cyclase of *M. tuberculosis* [61, 62]. The main advantage of this inventive method is that the recombinant membrane protein can be effectively expressed in prokaryotic as well as in eukaryotic systems. The invention further provides a kit for expression of membrane proteins.

Protein kinases and phosphatases are essential for virulence in a number of bacterial species that modulate the host-signaling network. M. tuberculosis has two functional tyrosine phosphatases, MptpA and MptpB (WO2005005639) [65], which are secreted into the culture filtrate by actively growing mycobacterial cells [66]. mptpA and mptpB knock out strains of M. tuberculosis were attenuated in the lungs and spleen of infected animals. The ability of mutant strains to survive in macrophages activated with IFN- was highly impaired [67]. This suggests that tyrosine phosphatases of M. tuberculosis help in its survival in the host cells by dephosphorylating proteins that are involved in IFNmediated signaling pathways. WO2005005639 published in 2005 relates to the role of mptpA and mptpB in the virulence and pathogenesis of mycobacteria. The invention provides mutant mycobacterium strains harboring a modified tyrosine phosphatase gene (mptpA or mptpB) wherein the mutant mycobacterium strain is incapable of expressing the active tyrosine phosphatase.

Eukaryotic type serine/threonine kinases of M. tuberculosis [68] are also attractive therapeutic targets. Because of their significance in signal transduction and their role in circumventing the hostile environment, many of them have been patent protected. Mutant strains of protein kinase G (PknG) [69, 70] and protein kinase H (PknH) were unable to survive under in vivo conditions (WO03074728) [71]. PknG was shown to be secreted in host cells, causing phagosomal maturation block [70]. The inhibition of phagosome-lysosome fusion may be mediated by phosphorylation of host proteins by PknG. PknH can phosphorylate EmbR, a protein hypothesized to modulate the levels of arabinosyltransferases involved in arabinan biosynthesis of arabinogalactan, a key molecule of the mycobacterial cell wall [72, 73]. pknB (protein kinase B, WO03074728) [71], one of the four *M. tuberculosis* kinases, conserved in the downsized genome of Mycobacterium leprae, is presumed to play an important role in the processes that regulate the complex life cycle of mycobacteria. Transposon-mutagenesis has shown that PknB and PknG are required for growth of mycobacterium under in vitro conditions [74]. PknB inhibitor 1-(5-isoquinolinesulfonyl)-2-methylpiperazine (WO03074728) [71, 75] was found to inhibit the growth of M. bovis (BCG) and M. smegmatis. WO03074728 published in 2003 relates to a method for identifying compounds capable of affecting the activity of serine/threonine kinases (PknB, PknG, PknH and PknJ) of M. tuberculosis. Knockout mutants of PknG and PknH display slower growth and viability both in vitro and in vivo. The compounds of this invention can be incorporated into a variety of formulations for therapeutic administration.

The two-component system plays a central role in the adaptation of pathogenic bacteria to the environmental signals prevailing within host tissues. The genes encoding

the response regulator DevR (Rv3133c/DosR, EP1472339) [76] and cytoplasmic region of the histidine kinase DevS (Rv3132c/DosS), are part of a well characterized twocomponent system of M. tuberculosis [77]. Expression of Rv3134c-devR-devS operon has been shown to be induced in hypoxic cultures of M. tuberculosis [78]. Guinea pigs infected with the mutant strain of DevR showed a significant decrease in gross lesions in lung, liver and spleen compared to guinea pigs infected with the parental strain [79]. The mutant strain also showed loss in viability in oxygen deprived cultures [76]. These observations suggest that DevR-DevS system is involved in the virulence of M. tuberculosis and acts as a key regulatory link between oxygen limitation and the initiation and maintenance of adaptive response to hypoxia. Therefore, this genetic system could serve as a vital target for the development of new drugs for elimination of dormant bacilli. WO03066838 published in 2003 relates to a process for identifying a novel target for the development of therapeutic modalities and drugs effective against tuberculosis. The patent relates to testing of M. tuberculosis devR mutant strain for virulence in guinea pigs.

## ENZYMES INVOLVED IN CELLULAR METABO-LISM

The proteins involved in metabolic pathways are also promising targets as they are essential for survival of the Arylamine N-acetyltransferases pathogen. WO9961625) [80] belong to a unique family of proteins found in a wide range of organisms, catalyzing the acetyl-CoA dependent N-acetylation of the arylamines, N and O acetylation of arylhydroxy amines and acetylation of aromatic hydrazines. Human NAT2 acetylates and inactivates the antituberculosis drug, isoniazid (INH) and is polymorphic. A homologue of human NAT2 in M. tuberculosis has also been shown to acetylate and inactivate INH in vitro [81]. Introduction of a substrate or an inhibitor, which may diminish the activity of NAT against INH can act as an anti-mycobacterial drug. WO9961625 published in 1999 relates to arylamine N-acetyltransferase proteins from M. tuberculosis and M. smegmatis together with antibodies raised against these proteins, and methods of detecting mycobacteria using such antibodies. WO9961625 has also described the methods for screening compounds that are ligands of arylamine N-acetyltransferase. Such ligands. designed using three-dimensional structures of NATs can be used for treatment of mycobacterial infections.

Microbes synthesize branched-chain amino acids such as isoleucine, valine and leucine. The three reactions of the isoleucine and valine biosynthesis catalyzed by enzymes are common to both pathways. One of the intermediates of the valine pathway is used for the synthesis of leucine. Therefore, inhibition of the isoleucine-valine pathway enzymes, acetolactate synthase (ALS) and ketol acid reductoisomerase (KARI) will result in elimination of all the three branched-chain amino acids (US5998420) [82]. Herbicides that inhibit plant branched-chain amino acid biosynthetic enzymes were tested for inhibition of M. tuberculosis growth in vitro [83, 84]. Sulphometuron methyl and metsulphuron methyl, inhibitors of ALS, were indeed able to affect the growth of M. tuberculosis. Furthermore, inhibitors of both ALS and KARI were effective against drug-resistant clinical isolates. Animal studies showed that sulphometuron methyl significantly prevented the growth of M. tuberculosis in lungs [85]. As mammals do not have branched-chain amino acid biosynthetic enzymes, treatment with these compounds should be specific to pathogenic organism. US patent no. 5998420 issued in 1999, patent applications AU2451397 and WO9737660 published in 1997 relate to a method for treating tuberculosis by administering therapeutically effective amount of a compound that inhibits an enzyme of the branched chain amino acid biosynthetic pathway in M. tuberculosis [82, 86, 87].

Thymidylate synthase is a ubiquitous enzyme, which catalyzes the essential methylation of dUMP to dTMP, one of the four bases required for DNA synthesis. The reaction requires  $N^5$ ,  $N^{10}$ -methylene  $H_4$  foliate as a cofactor. Thymidylate synthase activity is strongly linked to the activity of the two enzymes responsible for replenishing the cellular folate pool: Dihydrofolate reductase and Serine transhydroxymethylase. Thymidylate synthase of M. tuberculosis (Rv2492) has been cloned and characterized (WO02072805) [88]. The thymidylate synthase reaction is a crucial part of the pyrimidine biosynthesis pathway, which generates dCTP and dTTP for incorporation into DNA. Inhibition of dTMP synthesis leads to a loss of DNA production, an arrest of the cell cycle and eventually a 'thymine-less' cell death. The emergence of multi-drug resistant bacteria in recent years has prompted research into the use of mycobacterial thymidylate synthase inhibitors as antibiotics. WO02072805 published in 2002 relates to the identification of a novel protein, thymidylate synthase and its use in the diagnosis, prevention and treatment of disease. Moreover, thymidylate synthase has been implicated in the pathogenicity of the M. tuberculosis and the ligands of this protein are likely to be effective in controlling disease.

Macrophages produce nitric oxide and other reactive nitrogen intermediates (RNI) to control infection by M. tuberculosis [89]. Despite the protective effect of RNI, mycobacteria persist and multiply in macrophages. Genes required for resistance against host RNI were identified using transposon mutagenesis and the mutants were screened for hyper susceptibility to acidified nitrite [90]. The study yielded seven genes with transposon insertions, including the genes required for DNA repair (uvrB) and synthesis of a flavin cofactor (fbiC) (US2004213776) [91]. Five mutants had insertions in two proteasome-associated genes that encode a proteasome-associated adenosine triphosphatase (ATPase) called Mpa (Rv2115c) and a proteasomeassociated factor called Paf (Rv2097c) [90, 91]. Unlike wild type M. tuberculosis, mutants (Rv2115c and Rv2097c) failed to grow in resting primary macrophages isolated from wild type or iNOS<sup>-/-</sup> mice. An inhibitor of the human proteasomal protease. N-[4-morpholine]carbonyl- -[1-naphthyl]-Lalanine-L-leucine boronic acid blocked proteosomal protease activity in M. tuberculosis and suppressed the growth of M. tuberculosis in culture conditions [90]. A specific inhibitor of bacterial proteasome might be useful to sensitize M. tuberculosis to the immune system if they are combined with chemotherapeutic agents that target enzymes involved in RNI resistance. US2004213776 published in 2004 provides methods for screening compounds that inhibit proteasomal and protease activity, DNA repair enzyme activity, or flavinlike co-factor synthesis enzyme activity, where the inhibitory compounds have an ability to sensitize bacteria to the antibacterial effects of oxidative/nitrosative stress.

# ENZYMES INVOLVED IN VIRULENCE AND PERSISTENCE

M. tuberculosis virulence can be understood by finding the factors that are important for the progression of tuberculosis. In most cases, essentiality of these factors for virulence has been studied either by gene knockout, global gene inactivation by transposon mutagenesis or antisense strategy. Understanding the strategies employed by M. tuberculosis for persistence would allow designing of antibiotics or inhibitors that would specifically target persistent or latent bacilli, allowing shortening of time required for chemotherapy.

Hypoxia is proposed as a key signal sensed by mycobacterium to enter into the persistent state [92]. Importantly, the dormant form of the bacterium is resistant against conventional antimycobacterials [93, 94]. To identify genes induced in dormancy, Boon et al. subjected M. bovis BCG to an oxygen-limited Wayne culture system followed by proteome analysis. Their work revealed the up-regulation of response regulator Rv3133c and three other polypeptides: crystallin and two "conserved hypothetical" proteins, Rv2623 and Rv2626c (WO0248391) [95]. The gene encoding response regulator DevR (Rv3133c/DosR) has been shown to be involved in virulence of M. tuberculosis as discussed earlier. DevR, a transcriptional regulator, also regulates the expression of three other dormancy genes [96]. Thus, the dormant mycobacterium can be targeted by inhibitors of DevR, leading to down regulation of Rv2623 and Rv2626c. WO0248391 (2002) and US2004 242471 (2004) disclose a method for the identification of an antimycobacterial agent that modulates the activity and/or expression of a protein (Rv3133c, Rv2623 and Rv2626c) expressed by a mycobacterium in non-oxygen limiting stationary phase, hypoxic stationary phase or hypoxic growth phase [95, 97].

As discussed above, the drugs available against tuberculosis are unable to eliminate dormant bacilli. The persistent bacilli can get reactivated and cause disease following immune system perturbations. The resuscitation of dormant bacilli would make the bacterium susceptible to anti-mycobacterial drugs and would lead to successful elimination of dormant bacilli. The discovery of new class of pheromones, which stimulate the resuscitation of dormant bacilli has provided opportunities for treatment of persistent mycobacterial infection. Resuscitation factor (Rpf) is a secreted growth factor, which is required for the growth of vegetative cells in minimal media at very low inoculum densities, as well as for the resuscitation of dormant cells. M. tuberculosis contains five genes whose predicted products resemble Rpf from Micrococcus luteus (WO9855624) [98]. The Rpf-like proteins of M. tuberculosis (RpfA-E) were cloned and were shown to stimulate bacterial growth in laboratory culture at picomolar concentrations [99]. The Rpflike proteins of M. tuberculosis show cross-species activity as they were shown to stimulate the growth of the slowgrowing organism, M. bovis BCG and two fast-growing

organisms, *M. smegmatis* and *M. luteus*. Moreover, expression of Rpf in *M. smegmatis* could also stimulate its growth in minimal medium [99]. WO9855624 published in 1998 describes Rpf, their cognate receptors and inhibitors or mimetics. Pharmaceutical compositions and methods based on the Rpf and their receptors/convertases are also described in the patent.

Zhang et al. have also shown that dormant bacilli can be resuscitated by spent culture supernatant [100]. The components present in the culture supernatant having resuscitation activity were found to be phospholipids and peptide fragments of Rv1174c (WO0245736) [101]. Phosphatidyl-Lserine and a dioleoyl phosphatidyl-L-serine, both of which are precursors of phosphatidylethanolamine and phosphatidylcholine had significant resuscitation activity for the 6-month-old M. tuberculosis H37Ra cells over medium control as judged by colony forming units (CFU) assay. Phosphatidyl-L-serine allowed small inocula (10<sup>-7</sup> dilution) to form visible growth and CFU on plates, whereas the control culture grew only at 10<sup>-5</sup> dilution. Taken together, these results suggest that the phospholipids not only resuscitated old tubercle bacilli but also allowed small inocula to initiate growth in liquid culture. MALDI mass spectroscopic analysis of culture supernatant fractions and N-terminal amino acid sequencing revealed a peptide identical to the 29th to 58th aminoacids of a hypothetical protein Rv1147c of M. tuberculosis with unknown function [100]. WO0245736 published in 2002 provides the media for growth enhancement and resuscitation of mycobacteria. In this patent, diagnostic kits and treatment methods utilizing spent culture supernatant and cell extracts are also provided.

Alkyl hydroperoxide reductase, subunit C (AhpC) of M. tuberculosis (MtAhpC, WO9954479) [102] is a member of peroxidases, the peroxyredoxins found in many organisms. MtAhpC can detoxify hydroperoxides and protect against reactive nitrogen intermediates [103, 104] and specifically catalyzes the conversion of peroxynitrite (OONO) to nitrite [105]. Enhanced expression of MtAhpC is observed both in INH resistant KatG-deficient strains [106] as well as in INHsensitive strains when challenged with the drug [107]. The crystal structure of Cys176-Ser point mutant of MtAhpC suggested a model for the peroxidase reaction that includes the generation of a large internal cavity, which encloses the reaction center [108]. The intricacies of the mechanism supported by the structural details might provide a structural framework for the design of inhibitors with potential therapeutic applications. MtAhpC together with AhpD, dihydrolipoamide dehydrogenase, and dihydrolipoamide succinyltransferase (US2003190325) comprise an NADHdependent peroxidase and peroxynitrite reductase system, which provides support to the antioxidant defense of M. tuberculosis [109, 110]. The AhpD crystal structure revealed a trimer with two catalytic sulfhydryl groups; Cys-130 and Cys-133, in which each of the subunits had an identical but novel protein fold [111]. Structural details can support for designing inhibitors with potential utility as antitubercular agents. WO9954479 published in 1999 relates to the use of alkyl hydroperoxide reductase subunit C encoding gene, proteins or polypeptides to confer resistance against antimicrobial reactive nitrogen intermediates. AhpC can be used to screen drugs that inhibit the activity of AhpC and sensitize M. tuberculosis to RNI produced by host cells. Alternatively, therapeutics can be developed to treat gastric infection. AhpC can be useful in vaccines to prevent infection by M. tuberculosis, while the antibodies raised against this protein can be employed in passive immunization of infected person. These proteins, antibodies and DNA molecules may also be utilized in diagnostic assays to detect M. tuberculosis in tissue or body fluids. US2003190325 published in 2003 relates to the methods of inhibiting AhpD, dihydrolipoamide dehydrogenase and dihydrolipoamide succinyltransferase in the infected person to make the pathogen susceptible to antimicrobial reactive nitrogen intermediates or reactive oxygen intermediates. Methods of producing an AhpD crystal suitable for X-ray diffraction as well as methods for designing a compound suitable for treatment or prevention of tuberculosis are also disclosed.

During infection, many pathogens encounter deprivation of certain essential nutrients and cofactors. Mutations in the genes encoding enzymes in the biosynthetic/degradative pathways and acquisition systems for some of these factors have helped in deciphering new drug targets. Earlier it has been reported that M. tuberculosis shifts from a metabolism that preferentially uses carbohydrate when growing in vitro to one that utilizes fatty acids when growing in the host, which is supported by the fact that over 200 genes were annotated to be involved in fatty acid metabolism [9]. One such enzyme is isocitrate lyase (Icl, WO0233118) [112] that converts isocitrate to succinate in the glyoxylate shunt and thus helps to survive on acetate or fatty acids as the sole carbon source. Icl activity increases dramatically in the stationary phase of M. tuberculosis growth [113] and its mRNA levels increase during macrophage infection and in the lungs of infected mice [114-116]. It has also been demonstrated that Icl is important for survival of M. tuberculosis in the lungs of mice during the persistent phase of infection (2-16 weeks), but is not essential during the acute phase (0-2 weeks) [117]. Given its potential as a drug target against persistent infections, its structure was solved without ligand and in complex with two inhibitors [118]. Covalent modification of an active site residue, Cys 191, by the inhibitor 3-bromopyruvate traps the enzyme in a catalytic conformation with the active site completely inaccessible to solvent. These inhibitor-bound structures not only help to establish key residues in the active site, but enable to pinpoint interactions, which are essential in forming the closed conformation of the enzyme and are likely to be a key to successful drug discovery. WO0233118 published in 2002 discloses the importance of the glyoxylate shunt in the persistent phase of various infectious agents, including mycobacteria and the identification of targets for drug development. Crystals and three-dimensional structures of M. tuberculosis Icl, without ligand and in complex with two inhibitors that can be used in the design of inhibitors and therapeutic agents are also disclosed.

Protein export is an important aspect of bacterial pathogenesis. Research on diverse bacterial pathogens has demonstrated that the majority of virulence factors are secretory proteins [119, 120]. In bacteria, the majority of exported proteins are translocated by the Sec system, which recognizes the signal sequence of a pre-protein and uses ATP and the proton motive force to mediate protein translocation across the cytoplasmic membrane. SecA (WO9626276) [121] is an essential protein component of this system, containing the molecular motor that facilitates translocation. There are two homologues of secA in mycobacteria, secA1 and secA2. Using an allelic-exchange strategy in M. smegmatis, it was demonstrated that secA1 is the essential housekeeping protein whereas secA2 is an accessory factor for the secretion [122]. The topography of SecA of M. tuberculosis and its ATP binding sites are highly conserved, whereas its membrane insertion domains are species specific [123]. The crystal structure of SecA1 revealed that each subunit of the homodimer contains a motor domain and a translocation domain. The structure predicts that SecA can interact with the SecYEG pore and function as a molecular ratchet that uses ATP hydrolysis for physical movement of the preprotein [124]. A deletion of the secA2 gene in M. tuberculosis led to loss of virulence because SecA2 mediates secretion of SodA, a virulence factor of M. tuberculosis [125]. WO9626276 published in 1996 relates to an isolated nucleic acid encoding a SecA protein of M. tuberculosis. The invention includes the mutant SecA protein of M. tuberculosis and provides methods of screening for putative virulence factors translocated by SecA.

### CURRENT AND FUTURE DRUG DEVELOPMENTS

The availability of *M. tuberculosis* genome sequence [9] and recent advances in understanding the molecular basis of host pathogen interaction have opened new avenues for the development of novel antimycobacterial drugs. Promising new drug candidates such as PA-824 (nitroimidazopyran) have entered phase I clinical trials (www.tballiance.org). PA-824 exhibits bactericidal activity against both actively growing and static M. tuberculosis. The potency of PA-824 is attributed to its ability to get activated by M. tuberculosis F420 cofactor and inhibit the synthesis of protein and cell wall lipid [126-128]. Diarylquinoline (R207910), another molecule reported by Johnson and Johnson Pharmaceutical Research and Development, has a unique spectrum of potent and selective anti-mycobacterial activity under in vitro conditions. The molecule is active against both drugsensitive and drug-resistant M. tuberculosis and it was suggested that the drug targets the proton pump of adenosine triphosphate (ATP) synthase [129]. Plasma levels associated with efficacy of R207910 in mice were well tolerated in healthy human volunteers. Another drug, tetrahydrobenzothiophene (AX20017) developed by Axxima Pharmaceuticals AG, Germany, was shown to inhibit the kinase activity of PknG of M. tuberculosis. Chemical targeting of PknG led to the localization of *M. bovis* BCG into lysosomes causing bacterial lysis within macrophages [70]. AX20017 represents a promising candidate for the development of a class of drugs that would target the intracellularly residing mycobacteria.

In the past several years, it has been realized that controlling tuberculosis needs two issues to be addressed, drug resistance and persistence. A better understanding of the biology of tubercle bacilli, development in mycobacterial genetic tools, high throughput drug screening and structure based drug designing have increased the prospect of identifying novel anti-tubercle agents to combat drug resistant and persistent organisms. In short, investigation of *M. tuberculosis* pathogenesis has entered a new era and it is anticipated that the global challenge of tuberculosis will be surmounted in near future.

#### ACKNOWLEDGEMENTS

We thank Mr. Adesh Kumar Saini, Ms. Noor Jailkhani and Dr. Beena Pillai for their critical comments. Financial support is provided by SMM 0003 (CSIR).

#### REFERENCES

- Medical Research Council. Streptomycin treatment of pulmonary tuberculosis. Medical Research Council investigation. Br Med J 1948: 2: 769-82.
- [2] Pyle MM. Relative numbers of resistant tubercle bacilli in sputa of patients before and during treatment with streptomycin. Proc Staff Meet Mayo Clin 1947; 22: 465-72.
- [3] Youmans GP, Williston EH, Feldman WH, Hinshaw HC. Increase in resistance of tubercle bacilli to streptomycin: a preliminary report. Proc Staff Meet Mayo Clin 1946; 21: 126-7.
- [4] Medical Research Council. Treatment of pulmonary tuberculosis with streptomycin and para-aminosalicylic acid. Medical Research Council investigation. Br Med J 1950; 2: 1073-86.
- [5] Gilman AG. In: Gilman AG Ed, Antimicrobial agents, The pharmacologic basis of therapeutics. Pergamon Press, New York. 1990; 1061-62.
- [6] Goldberger MJ. Antituberculous agents. Med Clin North Am 1988; 72: 661-8.
- [7] Iseman MD. Treatment of multidrug-resistant tuberculosis. N Engl J Med 1993; 329: 784-91.
- [8] Duncan K, Sacchettini JC. In: Hatfull GF, Jacobs WRJ Ed, Approaches to tuberculosis drug development. Molecular genetics of mycobacteria. Washington DC, ASM Press. 2000; 297-307.
- [9] Cole ST, Brosch R, Parkhill J, et al. Deciphering the biology of Mycobacterium tuberculosis from the complete genome sequence. Nature 1998; 393: 537-44.
- [10] Dover LG, Cerdeno-Tarraga AM, Pallen MJ, Parkhill J, Besra GS. Comparative cell wall core biosynthesis in the mycolated pathogens, *Mycobacterium tuberculosis* and *Corynebacterium diphtheriae*. FEMS Microbiol Rev 2004; 28: 225-50.
- [11] Brennan PJ, Nikaido H. The envelope of mycobacteria. Annu Rev Biochem 1995; 64: 29-63.
- \*[12] Barry, C. E.: Yuan, Y. US5573915 (1996).
- [13] Yuan Y, Lee RE, Besra GS, Belisle JT, Barry CE 3rd. Identification of a gene involved in the biosynthesis of cyclopropanated mycolic acids in *Mycobacterium tuberculosis*. Proc Natl Acad Sci USA 1995; 92: 6630-4.
- [14] Cole ST, Eiglmeier K, Parkhill J, et al. Massive gene decay in the leprosy bacillus. Nature 2001; 409: 1007-11.
- [15] Quemard, A., Labesse, G., Daffe, M., et al.: WO03082911A2 and WO03082911A3 (2004).
- [16] Banerjee A, Sugantino M, Sacchettini JC, Jacobs WR Jr. The *mabA* gene from the *inhA* operon of *Mycobacterium tuberculosis* encodes a 3-ketoacyl reductase that fails to confer isoniazid resistance. Microbiology 1998; 144: 2697-704.
- [17] Kremer L, Dover LG, Carre're S, et al. Mycolic acid biosynthesis and enzymic characterization of the -ketoacyl-ACP synthase A-condensing enzyme from Mycobacterium tuberculosis. Biochem J 2002; 364: 423-30.
- [18] Marrakchi H, Lane'elle G, Que'mard A. InhA, a target of the antituberculous drug isoniazid, is involved in a mycobacterial fatty acid elongation system, FAS-II. Microbiology 2000; 146: 289-96
- [19] Schaeffer ML, Agnihotri G, Volker C, Kallender H, Brennan PJ, Lonsdale JT. Purification and biochemical characterization of the Mycobacterium tuberculosis beta-ketoacyl-acyl carrier protein synthases KasA and KasB. J Biol Chem 2001; 276: 47029-37.
- [20] Ducasse-Cabanot S, Cohen-Gonsaud M, Marrakchi H, et al. In vitro inhibition of the Mycobacterium tuberculosis betaketoacyl-acyl carrier protein reductase MabA by isoniazid. Antimicrob Agents Chemother 2004; 48: 242-9.

- [21] Cohen-Gonsaud M, Ducasse S, Hoh F, Zerbib D, Labesse G, Quemard A. Crystal structure of MabA from *Mycobacterium* tuberculosis, a reductase involved in long-chain fatty acid biosynthesis. J Mol Biol 2002; 320: 249-61.
- [22] Lety MA, Nair S, Berche P, Escuyer V. A single point mutation in the *embB* gene is responsible for resistance to ethambutol in *Mycobacterium smegmatis*. Antimicrob Agents Chemother 1997; 41: 2629-33
- [23] Telenti A, Philipp WJ, Sreevatsan S, et al. The emb operon, a gene cluster of Mycobacterium tuberculosis involved in resistance to ethambutol. Nat Med 1997; 3: 567-70.
- \*[24] Jacobs, Jr.W. R., Musser, J. M., Telenti, A.: US6015890 (2000).
- [25] Belanger AE, Besra GS, Ford ME, et al. The embAB genes of Mycobacterium avium encode an arabinosyl transferase involved in cell wall arabinan biosynthesis that is the target for the antimycobacterial drug ethambutol. Proc Natl Acad Sci USA 1996; 93: 11919-24.
- [26] Khoo KH, Douglas E, Azadi P, et al. Truncated structural variants of lipoarabinomannan in ethambutol drug-resistant strains of Mycobacterium smegmatis. Inhibition of arabinan biosynthesis by ethambutol. J Biol Chem 1996; 271: 28682-90.
- [27] Wolucka BA, McNeil MR, de Hoffmann E, Chojnacki T, Brennan PJ. Recognition of the lipid intermediate for arabinogalactan/arabinomannan biosynthesis and its relation to the mode of action of ethambutol on mycobacteria. J Biol Chem 1994; 269: 23328-35.
- [28] Alcaide F, Pfyffer GE, Telenti A. Role of *embB* in natural and acquired resistance to ethambutol in mycobacteria. Antimicrob Agents Chemother 1997; 41: 2270-3.
- [29] Strominger JL. In: Gunzalus IC, Stanier RY Ed, Biosynthesis of bacterial cell walls. The bacteria, vol. III. Washington DC, Academic Press, Inc. New York, NY Press. 1962; 413–70.
- [30] Julius M, Free CA, Barry GT. Alanine racemase (*Pseudomonas*). Methods Enzymol 1970; 17: 171-6.
- [31] Neuhaus FC. The enzymatic synthesis of D-alanyl-D-alanine. Purification and properties of D-alanyl-D-alanine synthesase. J Biol Chem 1962; 237: 778-86
- [32] Barletta , R. G., Barletta-C, O.: US2003133952A1 (**2003**).
- [33] Ishihama A. Role of the RNA polymerase alpha subunit in transcription activation. Mol Microbiol 1992; 6: 3283-8.
- [34] Russo FD, Silhavy TJ. Alpha: the Cinderella subunit of RNA polymerase. J Biol Chem 1992; 267(21): 14515-8.
- [35] Ebright RH, Busby S. The Escherichia coli RNA polymerase alpha subunit: structure and function. Curr Opin Genet Dev 1995; 5: 197-203.
- [36] Levin ME, Hatfull GF. Mycobacterium smegmatis RNA polymerase: DNA supercoiling, action of rifampicin and mechanism of rifampicin resistance. Mol Microbiol 1993; 8: 277-85.
- \*[37] Healy, J. M., Bodorova, J., Lam, K. T., Lesoon, A. J.: EP0956347A1 and EP0956347A4 (**2002**).
- [38] Wosten MM. Eubacterial sigma-factors. FEMS Microbiol Rev 1998; 22: 127-50.
- \*[39] Balganesh, M., Sharma, U.: WO9638478A1 (1996).
- [40] Collins DM, Kawakami RP, de Lisle GW, Pascopella L, Bloom BR, Jacobs WR Jr. Mutation of the principal sigma factor causes loss of virulence in a strain of the *Mycobacterium tuberculosis* complex. Proc Natl Acad Sci USA 1995; 92: 8036-40.
- [41] Hu Y, Coates AR. Transcription of two sigma 70 homologue genes, sigA and sigB, in stationary-phase *Mycobacterium tuberculosis*. J Bacteriol 1999; 181: 469-76.
- [42] Lam, K. T.: US6355469 (2002).
- [43] Raman S, Song T, Puyang X, Bardarov S, Jacobs WR Jr, Husson RN. The alternative sigma factor SigH regulates major components of oxidative and heat stress responses in Mycobacterium tuberculosis. J Bacteriol 2001; 183: 6119-25.
- [44] Kaushal D, Schroeder BG, Tyagi S, et al. Reduced immunopathology and mortality despite tissue persistence in a Mycobacterium tuberculosis mutant lacking alternative sigma factor, SigH. Proc Natl Acad Sci USA 2002; 99: 8330-5.
- [45] Geiman DE, Kaushal D, Ko C, et al. Attenuation of late-stage disease in mice infected by the Mycobacterium tuberculosis mutant lacking the SigF alternate sigma factor and identification of SigF-dependent genes by microarray analysis. Infect Immun 2004; 72: 1733-45.

- Chen P, Ruiz RE, Li Q, Silver RF, Bishai WR. Construction and characterization of a Mycobacterium tuberculosis mutant lacking the alternate sigma factor gene, sigF. Infect Immun 2000; 68:
- [47] DeMaio J, Zhang Y, Ko C, Young DB, Bishai WR. A stationaryphase stress-response sigma factor from Mycobacterium tuberculosis. Proc Natl Acad Sci USA 1996; 93: 2790-4.
- \*[48] Demaio, J., Young, D. B., Bishai, W. R., Zhang, Y.: WO9735611A1 (1997).
- [49] Molle V, Palframan WJ, Findlay KC, Buttner MJ. WhiD and WhiB, homologous proteins required for different stages of sporulation in Streptomyces coelicolor A3(2). J Bacteriol 2000; 182: 1286-95.
- \*[50] Soni, V., Khandrika, L. P., Agrawal, P.: US6645505 (2003).
- [51] Steyn AJ, Collins DM, Hondalus MK, Jacobs WR Jr, Kawakami RP, Bloom BR. Mycobacterium tuberculosis WhiB3 interacts with RpoV to affect host survival but is dispensable for in vivo growth. Proc Natl Acad Sci USA 2002; 99: 3147-52.
- Sassanfar, M., Schimmel, P. R.: US5756327 (1998). [52]
- Sassanfar M, Kranz JE, Gallant P, Schimmel P, Shiba K. A [53] eubacterial Mycobacterium tuberculosis tRNA synthetase is eukaryote-like and resistant to a eubacterial-specific antisynthetase drug. Biochemistry 1996; 35: 9995-10003.
- [54] Sutherland R, Boon RJ, Griffin KE, Masters PJ, Slocombe B, White AR. Antibacterial activity of mupirocin (pseudomonic acid), a new antibiotic for topical use. Antimicrob. Agents Chemother 1985; 27: 495-8.
- [55] Fuller AT, Banks GT, Mellows G, Barrow KD, Woolford M, Chain EB. Pseudomonic acid: an antibiotic produced by Pseudomonas fluorescens. Nature 1971; 234: 416-7.
- Martinis, S. A., Sassanfar, M., Kim, S., Lee, S. H., Schimmel, P. [56] R.: US5798240 (1998).
- Kim S, Jo YJ, Lee SH, Motegi H, Shiba K, Sassanfar M, [57] Martinis SA. Biochemical and phylogenetic analyses of methionyl-tRNA synthetase isolated from a pathogenic microorganism, Mycobacterium tuberculosis. FEBS Lett 1998; 427: 259-62.
- [58] Schimmel P, Tao J, Hill J. Aminoacyl tRNA synthetases as targets for new anti-infectives. FASEB J 1998; 12:1599-609.
- [59] Martinis, S. A., Zhang, J., Schimmel, P. R.: US5656470 (1997).
- [60] Lowrie DB, Aber VR, Jackett PS. Phagosome-lysosome fusion and cyclic adenosine 3':5'-monophosphate in macrophages infected with Mycobacterium microti, Mycobacterium bovis BCG or Mycobacterium lepraemurium. J Gen Microbiol 1979; 110: 431-41
- Voelkel, H., Schultz, J., Linder, J.: WO02092805A1 and [61] WO02092805C1 (2002).
- [62] Volkel, H., Schultz, J., Linder, J.: US2005048603 (2005).
- Ketkar AD, Shenoy AR, Kesavulu MM, Visweswariah SS. [63] Suguna K. Purification, crystallization and preliminary X-ray diffraction analysis of the catalytic domain of adenylyl cyclase Rv1625c from Mycobacterium tuberculosis. Acta Crystallogr D Biol Crystallogr 2004; 60: 371-3.
- [64] Shenoy AR, Srinivasan N, Subramaniam M, Visweswariah SS. Mutational analysis of the Mycobacterium tuberculosis Rv1625c adenylyl cyclase: residues that confer nucleotide specificity contribute to dimerization. FEBS Lett 2003; 545: 253-9.
- \*[65] Tyagi, A. K., Singh, R., Rao, V., et al.: WO2005005639A2 and WO2005005639A3 (2005).
- [66] Koul A, Choidas A, Treder M, Tyagi AK, Drlica K, Singh Y, Ullrich A. Cloning and characterization of secretory tyrosine phosphatases of Mycobacterium tuberculosis. J Bacteriol 2000; 182:5425-32.
- [67] Singh R, Rao V, Shakila H, et al. Disruption of mptpB impairs the ability of Mycobacterium tuberculosis to survive in guinea pigs. Mol Microbiol 2003; 50:751-62.
- [68] Av-Gay Y, Everett M. The eukaryotic-like Ser/Thr protein kinases of Mycobacterium tuberculosis. Trends Microbiol 2000;
- [69] Cowley S, Ko M, Pick N, et al. The Mycobacterium tuberculosis protein serine/threonine kinase PknG is linked to cellular glutamate/glutamine levels and is important for growth in vivo. Mol Microbiol 2004; 52: 1691-702.
- [70] Walburger A, Koul A, Ferrari G, et al. Protein kinase G from pathogenic mycobacteria promotes survival within macrophages. Science 2004; 304: 1800-4.

- Av-Gay, Y., Drews, S. J., Cowley, S.: WO03074728 (2003).
- [72] Sharma K, Chandra H, Gupta PK, et al. PknH, a transmembrane Hank's type serine/threonine kinase from Mycobacterium tuberculosis is differentially expressed under stress conditions. FEMS Microbiol Lett 2004; 233: 107-13.
- [73] Molle V, Kremer L, Girard-Blanc C, Besra GS, Cozzone AJ, Prost JF. An FHA phosphoprotein recognition domain mediates protein EmbR phosphorylation by PknH, a Ser/Thr protein kinase from Mycobacterium tuberculosis. Biochemistry 2003; 42: 15300-9.
- [74] Sassetti CM, Boyd DH, Rubin EJ. Genes required for mycobacterial growth defined by high density mutagenesis. Mol Microbiol 2003: 48: 77-84.
- [75] Drews SJ, Hung F, Av-Gay Y. A protein kinase inhibitor as an antimycobacterial agent. FEMS Microbiol Lett. 2001; 205: 369-
- Tyagi, J. S., Kapur, V.: EP1472339A1 (2004). \*[76]
- [77] Dasgupta N, Kapur V, Singh KK, et al. Characterization of a two-component system, devR-devS, of Mycobacterium tuberculosis. Tuber Lung Dis 2000; 80: 141-59.
- [78] Saini DK, Malhotra V, Dey D, Pant N, Das TK, Tyagi JS. DevR-DevS is a bona fide two-component system of Mycobacterium tuberculosis that is hypoxia-responsive in the absence of the DNA-binding domain of DevR. Microbiology 2004; 150: 865-
- [79] Malhotra V, Sharma D, Ramanathan VD, et al. Disruption of response regulator gene, devR, leads to attenuation in virulence of Mycobacterium tuberculosis. FEMS Microbiol Lett 2004; 231: 237-45
- [80] Sim, E., Payton, M., Sinclair, J.: WO9961625A1 (1999).
- Upton AM, Mushtaq A, Victor TC, et al. Arylamine N-[81] acetyltransferase of Mycobacterium tuberculosis is polymorphic enzyme and a site of isoniazid metabolism. Mol Microbiol 2001: 42: 309-17.
- Grandoni, J.: US5998420 (1999). [82]
- LaRossa RA, Schloss JV. The sulfonylurea herbicide [83] sulfometuron methyl is an extremely potent and selective inhibitor of acetolactate synthase in Salmonella typhimurium. J Biol Chem 1984; 259: 8753-7.
- [84] Chaleff RS and Mauvais J. Acetolactate synthase is the site of action of two sulphonyl urea herbicides in higher plants. Science 1984: 224: 1443-5.
- [85] Grandoni JA, Marta PT, Schloss JV. Inhibitors of branchedchain amino acid biosynthesis as potential antituberculosis agents. J Antimicrob Chemother 1998; 42: 475-82.
- [86] Grandoni, J.: AU2451397 (1997).
- [87] Grandoni, J.: WO9737660 (1997).
- [88] Fagan, R. J., Phelps, C. B., Gutteridge, A.: WO02072805A2 and WO02072805A3 (2003).
- [89] Nathan C, Shiloh MU. Reactive oxygen and nitrogen intermediates in the relationship between mammalian hosts and microbial pathogens. Proc Natl Acad Sci USA 2000; 97: 8841-8.
- [90] Darwin KH, Ehrt S, Gutierrez-Ramos JC, Weich N, Nathan CF. The proteasome of Mycobacterium tuberculosis is required for resistance to nitric oxide. Science 2003; 302: 1963-6.
- Darwin, K. H., Nathan, C. F.: US2004213776A1 (2004). \*[91]
- [92] Wayne LG, Sohaskey CD. Nonreplicating persistence of Mycobacterium tuberculosis. Annu Rev Microbiol 2001; 55:
- [93] Wayne LG, Sramek HA. Metronidazole is bactericidal to dormant cells of Mycobacterium tuberculosis. Antimicrob Agents Chemother 1994; 38: 2054-8.
- Wayne LG, Hayes LG. An in vitro model for sequential study of [94] shiftdown of Mycobacterium tuberculosis through two stages of nonreplicating persistence. Infect Immun 1996; 64: 2062-9.
- Dick, T., Calvin, B. K. K.: WO0248391A2 and WO0248391A [95] (2002)
- Park HD, Guinn KM, Harrell MI, et al. Rv3133c/dosR is a [96] transcription factor that mediates the hypoxic response of Mycobacterium tuberculosis. Mol Microbiol 2003; 48: 833-43.Esser, E., Wedel, H.: US2004242471 (2004).
- [97] Esser, E., Wedel, H.: US2004242471 (2004).
- [98] Mukamolova, G., Kaprelyants, A. S., Young, D. I., Kell, D. B., Young, M.: WO9855624 (1998).

- [99] Mukamolova GV, Turapov OA, Young DI, Kaprelyants AS, Kell DB, Young M. A family of autocrine growth factors in Mycobacterium tuberculosis. Mol Microbiol 2002; 46: 623-35.
- [100] Zhang Y, Yang Y, Woods A, Cotter RJ, Sun Z. Resuscitation of dormant *Mycobacterium tuberculosis* by phospholipids or specific peptides. Biochem Biophys Res Commun 2001; 284: 542-7.
- [101] Zhang, Y.: WO0245736 (2002).
- [102] Nathan, C. F., Xie, Q.: WO9954479A2, WO9954479A3 and WO9954479C2 (2000).
- [103] Chen L, Xie Q, Nathan C. Alkyl hydroperoxide reductase subunit C (AhpC) protects bacterial and human cells against reactive nitrogen intermediates. Mol Cell 1998; 1: 795-805.
- [104] Master SS, Springer B, Sander P, Boettger EC, Deretic V, Timmins GS. Oxidative stress response genes in *Mycobacterium tuberculosis*: role of ahpC in resistance to peroxynitrite and stage-specific survival in macrophages. Microbiology 2002; 148: 3139-44.
- [105] Bryk R, Griffin P, Nathan C. Peroxynitrite reductase activity of bacterial peroxiredoxins. Nature 2000; 407: 211-5.
- [106] Sherman DR, Mdluli K, Hickey MJ, et al. Compensatory ahpC gene expression in isoniazid-resistant Mycobacterium tuberculosis. Science 1996; 272: 1641-3.
- [107] Wilson M, DeRisi J, Kristensen HH, et al. Exploring druginduced alterations in gene expression in Mycobacterium tuberculosis by microarray hybridization. Proc Natl Acad Sci USA 1999; 96: 12833-8.
- [108] Guimaraes BG, Souchon H, Honore N, et al. Structure and mechanism of the alkyl hydroperoxidase AhpC, a key element of the Mycobacterium tuberculosis defense system against oxidative stress. J Biol Chem 2005; 280: 25735-42.
- [109] Bryk R, Lima CD, Erdjument-Bromage H, Tempst P, Nathan C. Metabolic enzymes of mycobacteria linked to antioxidant defense by a thioredoxin-like protein. Science 2002; 295: 1073-
- [110] Nathan, C. F., Lima, C. D., Bryk, R.: US2003190325A1 (2003).
- [111] Nunn CM, Djordjevic S, Hillas PJ, Nishida CR, Ortiz de Montellano PR. The crystal structure of *Mycobacterium tuberculosis* alkylhydroperoxidase AhpD, a potential target for antitubercular drug design. J Biol Chem 2002; 277: 20033-40.
- \*[112] Sacchettini, J. C., Mckinney, J. D. Russell, D. G., *et al.*: WO0233118A2 and WO0233118A3 (**2003**).
- [113] Wayne LG, Liu KY. Glyoxalate metabolism and adaptation of Mycobacterium tuberculosis to survival under anaerobic conditions. Infect Immun 1982; 37: 1042-9.
- [114] Dubnau E, Fontan P, Manganelli R, Soares-Appel S, Smith I. Mycobacterium tuberculosis genes induced during infection of human macrophages. Infect Immun 2002; 70: 2787-95.
- [115] Graham JE, Clark-Curtiss JE. Identification of *Mycobacterium tuberculosis* RNAs synthesized in response to phagocytosis by

- human macrophages by selective capture of transcribed sequences (SCOTS). Proc Natl Acad Sci USA 1999; 96: 11554-9
- [116] Shi L, Jung YJ, Tyagi S, Gennaro ML, North RJ. Expression of Th1-mediated immunity in mouse lungs induces a Mycobacterium tuberculosis transcription pattern characteristic of nonreplicating persistence. Proc Natl Acad Sci USA 2003; 100: 241-6.
- [117] McKinney JD, Bentrup KHZ, Munoz-Elias EJ, et al. Persistence of Mycobacterium tuberculosis in macrophages and mice requires the glyoxalate shunt enzyme isocitrate lyase. Science 2000; 406: 735-8.
- [118] Sharma V, Sharma S, Hoener zu Bentrup K, et al. Structure of isocitrate lyase, a persistence factor of Mycobacterium tuberculosis. Nat Struct Biol 2000; 7: 663-8.
- [119] Finlay BB, Falkow S. Common themes in microbial pathogenicity revisited. Microbiol. Mol Biol Rev 1997; 61: 136-69
- [120] Miller JF, Cossart P. Bacterial pathogenesis: before the postgenomic era. Curr Opin Microbiol 1999; 2: 15-7.
- \*[121] Schmidt, M. G., Owens, M. U., King, H. C., Quinn, F. D.: WO9626276A1 (1996).
- [122] Braunstein M, Brown AM, Kurtz S, Jacobs WR Jr. Two nonredundant SecA homologues function in mycobacteria. J Bacteriol 2001; 183: 6979-90.
- [123] Owens MU, Swords WE, Schmidt MG, King CH, Quinn FD. Cloning, expression, and functional characterization of the Mycobacterium tuberculosis secA gene. FEMS Microbiol Lett 2002; 211: 133-41.
- [124] Sharma V, Arockiasamy A, Ronning DR, et al. Crystal structure of Mycobacterium tuberculosis SecA, a preprotein translocating ATPase. Proc Natl Acad Sci USA 2003; 100: 2243-8.
- [125] Braunstein M, Espinosa BJ, Chan J, Belisle JT, Jacobs WR Jr. SecA2 functions in the secretion of superoxide dismutase A and in the virulence of *Mycobacterium tuberculosis*. Mol Microbiol 2003; 48: 453-64.
- [126] Stover CK, Warrener P, VanDevanter DR, et al. A small-molecule nitroimidazopyran drug candidate for the treatment of tuberculosis. Nature 2000; 405: 962-6.
- [127] Tyagi S, Nuermberger E, Yoshimatsu T, et al. Bactericidal activity of the nitroimidazopyran PA-824 in a murine model of tuberculosis. Antimicrob Agents Chemother 2005; 49: 2289-93.
- [128] Lenaerts AJ, Gruppo V, Marietta KS, et al. Preclinical testing of the nitroimidazopyran PA-824 for activity against Mycobacterium tuberculosis in a series of in vitro and in vivo models. Antimicrob Agents Chemother. 2005; 49: 2294-301.
- [129] Andries K, Verhasselt P, Guillemont J, et al. A diarylquinoline drug active on the ATP synthase of Mycobacterium tuberculosis. Science 2005; 307: 223-7.