Construction of *lipF* anti-sense knock out mutant of *Mycobacterium fortuitum* and study of its role in *in-vivo* infection and virulence

By

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CERTIFICATE

This is to certify that Ms. Divya has carried out the B.Tech Biotechnology dissertation project work on "**Construction of** *lipF* anti-sense knock out mutant of *Mycobacterium fortuitum* and study of its role in *in-vivo* infection and virulence" from Jaypee University of Information Technology, Solan, Himachal Pradesh under my supervision from August 2016 to May 2017. This work has not been submitted partially or wholly to any other university or institute for the award of this or any other degree or diploma.

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Date :

DECLARATION

I hereby declare that the work presented in this project dissertation entitled, "**Construction of** *lipF* anti-sense knock out mutant of *Mycobacterium fortuitum* and analysis to study its role in *in-vivo* infection and virulence" submitted as partial fulfillment of B.Tech in Biotechnology VIIIth semester was carried out at Department of Biotechnology and Bioinformatics, Jaypee University of Information Technology, Waknaghat, Himachal Pradesh. The work presented in this project dissertation is original and will remain intellectual property of Department of Biotechnology and Bioinformatics, Jaypee University of Information Technology, Jaypee University of Information is original and will remain intellectual property of Department of Biotechnology and Bioinformatics, Jaypee University of Information Technology, Waknaghat, Himachal Pradesh.

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ABSTRACT

Tuberculosis can be considered as one of the major causes of deaths throughout the world. In order to identify the genes required for the pathogenicity of *Mycobacterium tuberculosis*, a functional genomic approach has been developed. Also, these hospital acquired outbursts caused by the mycobacteria are in the picture of medical sciences for more than 20 years, yet to be diminished. Most of these outbreaks have involved the rapidly growing mycobacterial species. Among these mycobacterial species, tuberculosis is considered as the major factor of most of the deaths worldwide due to a single pathogen involved, Mycobacterium tuberculosis. Another important species in this respect belonging to the same taxon includes *Mycobacterium fortuitum* which is a non-tuberculous mycobacterium (NTM) causing various clinical symptoms such as cutaneous infections, respiratory infections, joint infections and disseminated infections in immuno-compromised patients. Out of the all the genes identified for the virulence of this pervasive disease, lipF has played a major role whose significance hasn't been taken into consideration by many researchers as such. This gene coding for lipid esterase, expresses well for the virulence of tuberculosis and, is regulated by PhoP. lipF implicates modification of the mycobacterial cell wall as an adaptive response to acid damage, also degrades host lipids during infection, making fatty acids available as building blocks for lipid biosynthesis⁻ Gene helps in compensating with all the un-favorable acidic conditions in macrophages, where mycobacterium resides. This project mainly aims on the construction of *lipF* anti-sense knock out mutant by taking *Mycobacterium fortuitum* as the base model to check its role in virulence of tuberculosis as the homology between the genomic sequences (M. fortuitum and M. tuberculosis) of the two was found to be 67% (463/696).

Objectives

- To construct *lipF* anti-sense knock out mutant of *Mycobacterium fortuitum*.
- To study the role of anti sense knock out mutants *in-vivo* infection and virulence.

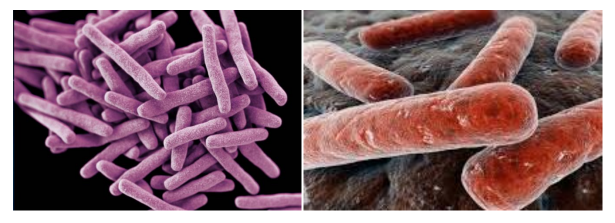
Chapter 1 Introduction

Pulmonary diseases caused by mycobacteria are increasing day by day throughout the world and hence there is an immediate need for the thorough study of the mechanisms used by these bacterial pathogens when they conquer human body cells and cause disease. In the contemporary era, the importance of the resistance acquired by these bacteria for multiple antibiotics has gained the spotlight. Yet there is no as such accurate data available because of the less number of reportable conditions in maximum countries. Ample number of mechanisms has been employed by bacteria to cause certain infectious diseases in human hosts. Though, variation occurring on the basis of geographic conditions signifies the widespread presence of these bacteria in the environment. Bacterial pathogens manifest a huge variety and range of biological entities which bind to host cell targets to facilitate diversified host responses. Different mechanisms used on the molecular level by bacteria to interact with the host can be distinctive for a particular pathogen or perpetuate beyond various divergent species. A cue to battle with these bacterial diseases is the identification and characterization of all these unconventional strategies. Some definite virulence facets are obligatory for complete pathogenicity of the bacteria irrespective of the host cell.

Tuberculosis (TB) has said to be one of the oldest accomplished human menace, which is still causing enormous number of deaths every year if talking about infectious diseases. Around 2.5-3 million people dies every year as per the recent statistics of World Health Organization(WHO) in spite of the use of attenuated vaccines and a variety of antibodies throughout the world. In 2007, approximately one third of the world's population were infected with the pathogenic *Mycobacterium tuberculosis* which is the single causative agent for tuberculosis. *M. tuberculosis* is a large non-motile and rod shaped, obligate bacterium which belongs to the family of mycobacteriaceae whereas distantly associated with actinomycetes. The bacilli of tuberculosis are 2-4 μ m in length and 0.2 -0.5 μ m in width. Because of the obligatory nature of the bacterium, in most of the cases of tuberculosis, the bacterial complexes are always found in the well-aerated upper lobes of the lungs. A number of non-pathogenic mycobacterium are constituents of usual flora of humans which are usually observed frequently in dry and oily areas. Even in the case of persons infected with HIV, tuberculosis is the foremost cause of death worldwide as per the statistics of World Health Organization 2011.

Mycobacterium tuberculosis was identified by Robert Koch in 1882, and *M. fortuitum* was identified soon afterwards. It was classified as a Non Tuberculous

Mycobacterium (NTM) and was not must studied until 1954. Edward Runyon took it upon himself to categorize all the NTMs at this time. *M. fortuitum* is now characterized as Runyon Group IV, which means, among other lab culture attributes, it is a comparatively rapid grower.



(Fig. 1.1)

(Fig. 1.2)

(Figure : Microscopic view of rod shaped Mycobacterium tuberculosis)^[30,31]

Infection in healthy humans caused by *Mycobacterium fortuitum* is rare, but exposure to large and repeated amounts of the organism can activate the immune system and cause disease. Infections most likely occur in immuno-compromised patients. Diseases and infections caused vary from each other and involve almost every organ and tissue system. The most common infections involve NTM lung disease, leprosy, local cutaneous disease, lymphadenitis, surgical site infections and injection site inflammation. Nearly all surgical site infections and injection site inflammation occur when the open wound gets contaminated under infectious or commonly used tap water.



(Fig. 1.3)

(Fig. 1.4)

(Figure : Patient suffering from tuberculosis, Lesions due to infection from M. fortuitum)^[32,33]

Lipid metabolism plays an important role in the sustenance of *M. tuberculosis* in the macrophages and out of the 250 genes involved in the lipid metabolism of bacteria *lipF* encodes a polypeptide with similarity to lipid esterases. The association of lipase production with pathogenicity has been suggested by several works concerning non-mycobacterial pathogens. *lipF* implicates modification of the mycobacterial cell wall as an adaptive response to acid damage, also degrades host lipids during infection, making fatty acids available as building blocks for lipid biosynthesis . The gene helps in compensating with all the un-favorable acidic conditions in macrophages, where bacterium resides. Hence, virulence of tuberculosis is chiefly dependent on the expression of *lipF* gene. Since, the gene is important for the pathogenesis and persistence of *M. tuberculosis* in the macrophages, this project has mainly focused on the role of *lipF* in the above said factors of tuberculosis by taking *Mycobacterium fortuitum* as the base model and construction of anti-sense knock-out mutant of *M. fortuitum*. In the further stages of the project, work done on the molecular level will be studied on various animal cell cultures and animal models, if possible.

Chapter 2 Review of Literature

2.1 Lipase Family :

Microscopic organisms of the *Mycobacterium tuberculosis complex* show remarkable abilities to subvert and to oppose the bactericidal reactions of their tainted host. These limits have driven this bacillus to colonize 33% of the total populace and to slaughter about three million individuals every year. Amid the contamination by means of the airborne course, *M. tuberculosis* bacilli are breathed in into the alveoli of the lung where they achieve macrophages. This cooperation of *M. tuberculosis* with the mononuclear phagocytes is a key stride in the pathogenesic procedure.

In vitro, M. tuberculosis has been appeared to tie either straightforwardly, or after opsonization, to a wide range of phagocyte receptors that permit the disguise of the microbes. In any case, the relative significance in vivo of these conceivable courses of section stays to be set up. It is likewise vague what ligands are included, in vivo, in the adherence. Without a doubt, numerous mycobacterial envelope atoms and in addition opsonins can intervene the in vitro official of M. tuberculosis to the phagocyte receptors, however their accessibility at the disease site (for the host ligand) or their nearness at the bacterial surface (for the mycobacterial particles) is still under scrutiny. Once phagocytosed, *M. tuberculosis* duplicates inside a particular compartment that does not ferment, conceivably in light of the avoidance of the proton ATPase from the layer of this organelle. This phagosome does not continue through the standard development pathway, and is by all accounts obstructed at a halfway phase of development. From these reviews, it appears to be certain that *M. tuberculosis* has advanced harmfulness systems that enable it to regulate the development of the phagosome and to oppose the lethal particle delivered by the macrophages.^[1] Be that as it may, the mycobacterial parts in charge of these bizarre elements stay obscure. These various reviews concerning this association between M. tuberculosis and the host underscore our poor comprehension of the mycobacterial elements, and their part in the diverse strides of the irresistible procedure.

A hereditary way to deal with recognize these mycobacterial elements is disconnect mutants influenced in one or a few stages of the irresistible procedure, and to distinguish the aggravates no longer created by these strains. Noso-comial flare-ups and pseudo-episodes brought on by the nontuberculous mycobacteria (NTM) have been perceived for over 20 years and keep on being an issue. Comparative sequence analysis of the *M. tuberculosis* genome has uncovered that it contains 250 proteins required in lipid digestion contrasted with just 50 in *Escherichia coli*. Among these proteins, a group of 21 carboxyl ester hydrolases, called *Lip* (A to W, aside from K and S), have been commented on as putative esterases or lipases, in light of the nearness of the agreement succession GXSXG normal for individuals from the α/β hydrolase overlap family. Inside this family, the current gem structure of the *M. tuberculosis* antigen 85C (Ag85C), a mycolyltransferase required for survival of mycobacteria, alongside that of the noncatalytic *M. tuberculosis* MPT51 protein (FbpC1), which is included in mycobacteria pathogenicity, have uncovered that they have the same α/β hydrolase crease. In this way, a nitty gritty biochemical portrayal of all individuals from the *Lip* family ought to be performed past the computational investigation. For a long time, it was by and large expected that lipases are ineffectively dynamic against solvent esters and turn out to be notably dynamic when the dissolvability furthest reaches of the substrate is surpassed, a wonder called interfacial actuation. Conversely, esterases don't share this conduct and show their maximal movement on esters in arrangement. Sadly, biochemical investigations of a few lipases have demonstrated that the interfacial enactment wonder can't be viewed as a general (and adequate) administer to separate between a lipase and esterase. An unmistakable qualification amongst lipases and esterases was built up as of late from the examination of the K estimations of these two classes of carboxyl ester hydrolases utilizing halfway dissolvable triacylglycerols and vinyl esters as substrates. Where a lipase can likewise be viewed as an esterase, the K values speak to a solid foundation to separate between these two classes of catalysts portrayal, the Lip expression could be befuddled to allude to a NLH, and the first promotion number ought to be utilized to maintain a strategic distance from disarray.^[7]

The *Mycobacterium tuberculosis* genome contains an inquisitively high number of proteins required in the absorption of lipids having a place with the *Lip* family, including distinctive nonlipolytic and lipolytic hydrolases.^[7] In one of the studies based on the ruinous characteristics of lipolytic mixes from microorganisms with extraordinary highlight on *Mycobacterium tuberculosis*, it was seen that in vivo, triacylglycerols as joining bodies are accessible in tubercle bacilli in the lungs. This pathogenic bacterium has a lipase quality (*Lip*) family, which is conveyed and differentially controlled under a grouping of in vitro conditions. Next to no exploration work has been finished on these lipolytic chemicals. A prevalent understanding of lipolytic mixes in mycobacteria would incite develop new frameworks for tuberculosis treatment.^[3]Among the diverse metabolic pathways included, lipid assimilation is thought to be one of the essential pathways taken by mycobacteria, and this pathway in this way ought to be considered in detail. Various irregular lipids, including

phthiocerol dimycocerosates, mycolic destructive, glycolipids, polyketides and glycans, have been seen to be required in the ruinous tendency and pathogenicity of *M. tuberculosis*.^[2]

2.2 *Lip* family qualities of *M. tuberculosis* and their plausible capacity :^[3]

Gene Product	Theoretical Molecular Mass	Predicted subcellular localization	Activity Type	Probable Role/ Function	Maximum Identity
LIPC (Rv0220)	44.3	Cytoplasmic	Probable esterase	Carboxylesterase type B	77% with alpha/beta hydrolase
LIPD (Rv1923)	47.2	Periplasmic	Similar to esterases, beta-lactamases	Defense mechanisms	69% with β-lactamase
LIPE (Rv3775)	45.3	Periplasmic	Carboxylic-ester hydrolase	Defense mechanisms	79% with β-lactamase
LIPF (Rv3487c)	29.4	Cytoplasmic	Unknown but similar to esterases and lipases	intermediary metabolism and respiration	75% with alpha/beta hydrolase
LIPG (Rv0646c)	32.9	Cytoplasmic	Homology to hydrolases	Lipid transport and metabolism	80% with alpha/beta hydrolase
LIPH (Rv1399c)	33.9	Cytoplasmic	Possible lipase	Lipid transport and metabolism	70% with alpha/beta hydrolase fold
LIPI (Rv1400c)	34.0	Cytoplasmic	Possible lipase	Intermediary metabolism and respiration	69% carboxylesterase family protein
LIPJ (Rv1900c)	49.7	Cytoplasmic	Similarity to esterases	Alkaloid biosynthesis II	75% with alpha/beta hydrolase fold
LipK, MBTJ(Rv2385)	32.9	Cytoplasmic	Probable esterase/acetyl hydrolase	intermediary metabolism	71% with alpha/beta hydrolase fold
LIPL (Rv1497)	45.8	Cytoplasmic	Probable esterase	Intermediary metabolism and respiration	70% with β-lactamase
LIPM (Rv2284)	46.7	Cytoplasmic	Probable esterase	Lipid transport and metabolism	78% with alpha/beta hydrolase fold
LIPN (Rv2970c)	40.1	Cytoplasmic	Lipase-like enzyme	Lipid transport and metabolism	77% with alpha/beta hydrolase fold
LIPO (Rv1426c)	46.1	Cytoplasmic	Possible esterase, also similar human arylacetamide deacetylase	Lipid transport and metabolism	79% with alpha/beta hydrolase fold
LIPP (Rv2463)	42.8	Periplasmic	Similar to esterases from Pseudomonas spp.	Defense mechanisms	76% β-lactamase
LIPQ (Rv2485c)	45.2	Cytoplasmic	Carboxylesterase	Intermediary metabolism and respiration	74% with alpha/beta hydrolase fold
LIPR (Rv3084)	32.6	Cytoplasmic	Similar to acetyl- hydrolase	intermediary metabolism and respiration	68% with alpha/beta hydrolase fold
LipS (MesT, Rv3176c)	35.2	Cytoplasmic	Similarity to esterases/ lipases/epoxide hydrolase	Virulence, detoxification, adaptation	99% with amidase
LIPT (Rv2045c)	56.1	Cytoplasmic	Probable carboxylesterase	Lipid transport and metabolism	71% Carboxyesterase
LIPU (Rv1076)	31.7	Cytoplasmic	Unknown but very similar to esterases and lipases	Alpha/beta hydrolase	76% with alpha/beta hydrolase fold
LIPV (Rv3203)	23.6	Cytoplasmic	Unknown but shows some similarity to lipases	intermediary metabolism and respiration	71% with alpha/beta hydrolase fold
LIPW (Rv0217c)	32.2	Cytoplasmic	Esterase	Alkaloid biosynthesis II	75% with alpha/beta hydrolase fold
LipX, PE Family Protein (Rv1169c)	10.8	Cytoplasmic	Esterase/lipase	Hydrolases or acyltransferases	73
LIPY (Rv3097c)	45.0	Inner-membarane	Triacylglycerol lipase, PE-PGRS family	Lipid transport and metabolism	99% with PE-PGRS family
LipZ (Rv1834)	31.6	Cytoplasmic	Probable hydrolase	Intermediary metabolism	76% with hydrolase

Gene Product	M. smegmatis	M. marinum	M. leprae	M. abscessus	M. bovis
LIPC (Rv0220)	71% with α, β hydrolase fold	79%	_	_	100%
LIPD (Rv1923)	family protein 69% with β-lactamase	80%	64% esterase	68% with putative lipase/β-lactamase	99% with putative lipase LipD
LIPE (Rv3775)	72% with $\beta\text{-lactamase}$	81%	79% with probable hydrolase	74%	100%
LIPF (Rv3487c)	_	72%	67% with probable esterase	_	100%
LIPG (Rv0646c)	73% with hydrolase, α , β hydrolase fold	79%	75% with probable hydrolase	68%	100%
LIPH (Rv1399c)	70% with esterase	78%		_	100%
LIPI (Rv1400c)	71% with esterase	77%	67% with probable lipase	67% with LipH	100%
LIPJ (Rv1900c)	_	75% with lignin peroxidise	71% with probable esterase	_	99% with putative lipoprotein
LipK, MBTJ(Rv2385)	_	74% with acetyl hydrolase	-	65% with putative lipase	100% with putative acetyl hydrolase
LIPL (Rv1497)	68% with β -lactamase	81%	69% with probable esterase	68% with putative lipase	99% with putative esterase
LIPO (Rv1426c)	72% with peptidase	79%	-	_	100% with putative esterase
LIPP (Rv2463)	74% with putative carboxylesterase	81%	69% with probable esterase	71%	100%
LIPQ (Rv2485c)	66% with peptidase	76%	-	_	100% with putative carboxlesterase
LIPR (Rv3084)	_	_	-	_	99% with probable acetyl hydrolase
LipS (MesT, Rv3176c)	_	_	_	_	99% with amidase
LIPT (Rv2045c)	71% with para- nitrobenzyl esterase	79% with carboxylesterase	70%	71% with putative carboxylesterase	100% with putative carboxylesterase
LIPU (Rv1076)	69% with esterase	81%	76% with putative esterase	_	100% with putative lipase & 73% with putative LipF
LIPV (Rv3203)	_	79%	_	_	100%
LIPW (Rv0217c)	73% with α, β hydrolase fold	67% with esterase/ lipase	_	70%	99%
LipX, PE Family Protein (Rv1169c)	_	_	_	_	100% with PE fam- ily protein
LIPY (Rv3097c)	_	80% with PE-PGRS family protein	_	_	99% with PE-PGRS family protein
LipZ (Rv1834)	_	76% with hydrolase	_	_	100% with putative hydrolase

2.3 Role of *lipF* in the virulence of *M. tuberculosis* :

The part of *lipF* quality of *M. tuberculosis* has been ensnared in pathogenesis and its promoter has been appeared to be upregulated by acidic anxiety. This announcement was demonstrated in a review where a 59 base combine corrosive inducible insignificant promoter locale was segregated in which a putative -10 area was distinguished. To research the -10 area of the *lipF* promoter, mutational examination was finished. Transformations inside this particular district prompt an effective abatement of promoter movement, while a change outside of this area does not influence promoter action. Another review says that the recombinant Rv3487c (*lipF*), subsequent to denaturing refinement and refolding, hydrolyses the substrate phosphatidylcholine and creates diacylglycerol by expelling the phosphocholine. One of the researcher in 2007 filtered recombinant Rv0183 quality item indicating lipase

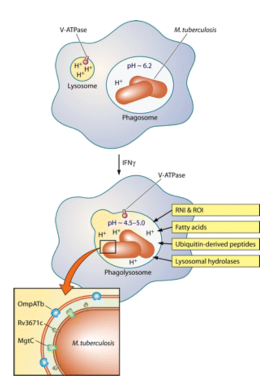
movement and revealed that this compound demonstrates an ideal action at pH values extending from 7.5 to 9.0 utilizing mono-olein as substrate.^[4]

lipF encodes a polypeptide with closeness to lipid esterases. The relationship of lipase creation with pathogenicity has been recommended by a few works concerning non-mycobacterial pathogens. Another match of researcher in 1995 demonstrated that the generation of a lipase by *V. cholerae* was prompted after contamination in a baby mouse demonstrate. Be that as it may, an invalid mutant for the quality encoding this lipase did not give off an impression of being constricted in an opposition measure. Different pathogens, for example, *Pseudomonas cepacia* and *Pseudomonas aeruginosa*, were appeared to emit lipases. Proteins refined from these living beings were appeared to have a few impacts in vitro on eukaryotic cells. They restrain the phagocytic elements of alveolar macrophages and regulate the arrival of incendiary middle people by various cells of the invulnerable framework . Another conceivable capacity for these lipases may be to corrupt the host lipid to give unsaturated fats to the pathogen.^[1]

2.4 Acidic nature of *M. tuberculosis*:

In vitro investigations of microscopic organisms at low pH are educational on the grounds that they can show whether the microorganisms are probably going to be acid safe or delicate amid disease. These reviews can likewise distinguish bacterial elements that give security against low pH and may do as such also in the host condition. In this manner, to start understanding whether *M. tuberculosis* opposes acid in vivo, it is valuable to first audit survival of *M. tuberculosis* in acid in vitro. Critically, in any case, the translation of in vitro studies is confounded by the perceptions that survival of numerous microscopic organisms in acid is reliant on the way of life conditions, for example, bacterial thickness and piece of the test medium. These factors likewise drastically impact the survival and development of mycobacteria at low pH. When all is said in done, the quickly developing, saprophytic mycobacteria become over a more extensive pH run than the pathogenic, moderate developing mycobacteria. This may mirror that the situations in which saprophytic mycobacteria dwell, for example, soil and water, are frequently acidic. Strikingly, mycobacterial species were discovered extraordinarily enhanced in greatly acidic volcanic shake at pH 1. With people being its lone regular habitat and inward breath its most normal course of passage into the body, M. tuberculosis does not have to keep up such a high resilience for acidic conditions. E. coli additionally shows more prominent acid resistance at high densities, and a cell-cell contact-based instrument seems, by all accounts, to be included.

Defensive variables emitted by E. coli may likewise assume a part in its imperviousness to acidic. The in vitro perceptions of M. tuberculosis in acidic condition welcomes the hypothesis that the bacterium may be profoundly vulnerable to the low pH of the phagolysosome, especially on the off chance that one considers the bacterial thickness (that is, number of microorganisms per unit liquid volume) in a phagosome "low." However, one may likewise consider the bacterial thickness in a phagosome to be to a great degree high. More critical, executing of *M. tuberculosis* at pH 4.5 is significantly affected by the creation of the medium in a way that can be viewed as artifactual. Due to their inclination to bunch, mycobacteria are usually developed in cleansers to take into account scattered development and readiness of moderately uniform bacterial suspensions for trial contemplates. Free unsaturated fats are lethal to *M. tuberculosis*, especially at low pH. It was accounted for that an assortment of strains of *M. tuberculosis* are impervious to executing at a pH of 4.5 in phosphatecitrate cradle. The bacilli are likewise ready to keep up a close nonpartisan intrabacterial pH when set in phosphate-citrate support at pH 4.5, demonstrating that they can counter the passage of protons. In this way, in basic support M. tuberculosis opposes phagolysosomal centralizations of corrosive.^[6]



(Figure 2.1 *M. tuberculosis inside the macrophage. In resting macrophages, M. tuberculosis impedes phagosome development and lives in a somewhat acidic compartment. Enactment with IFN-\gamma brings about phagosome development and phagosome-lysosome combination. This opens the microscopic organisms to have gotten push including protons from the*

vacuolar ATPase, RNI and ROI, free unsaturated fats, ubiquitin-inferred peptides, and lysosomal hydrolases. M. tuberculosis opposes fermentation with the assistance of the Rv3671c-encoded layer bound serine protease, the putative magnesium transporter MgtC, and the pore-shaping M. tuberculosis external layer protein (OmpATb). The correct systems by which these proteins give acid resistance stay to be distinguished)^[6]

The previously mentioned perceptions laid ground for the showing that when lysosomes combine with *M. tuberculosis* containing phagosomes in immunologically enacted macrophages, the phagolysosomal pH tumbles to 4.5 to 5.0. Various reviews have affirmed these perceptions. In one of the reviews where fluorescent mannosylated dots were utilized, it was accounted for that the pH of the macrophage phagolysosome falls quickly, inside 15 to 60 min, to a pH somewhat underneath 5.0. Along these lines, the pH of the macrophage phagolysosome seems to shift with the immunological condition of actuation of the macrophage and the way of the phagocytic molecule however by and large achieves a pH related with suspension of development of numerous facultative intracellular bacterial pathogens in soup culture. Absolutely, overabundance protons can harm DNA, proteins, and lipids and upset biochemical responses. Regardless of whether phagosomal acid is really a noteworthy bactericidal effector mechanism of macrophages is hard to set up.^[5]

2.5 *PhoPR* : two component regulation :

Bacterial two-component regulatory systems (2CRs) are key administrative frameworks used to react to natural changes adjusting gene expression because of outer jolts to produce adaptive reactions. Prokaryotic 2CRs are exceptionally rationed, and in their least difficult structures comprise of a sensor histidine kinase (HK) and an effector reaction controller (*RR*). Because of particular flags the HK autophosphorylates and afterward exchanges this phosphate to the *RR*. Much of the time the enactment of the *RR* prompts adjustment of quality expression through DNA official, empowering bacterial adjustment to the underlying natural jolt. *M. tuberculosis* has 12 finish 2CRs, a modest number contrasted with other bacterial species with comparative genome sizes. *PhoPR* is one of only a handful few mycobacterial 2CRs that are moderately all around characterized, with a key administrative part in controlling cell divider sythesis and harmfulness. *PhoP*, the *RR*, emphatically controls a few noteworthy procedures, including oxygen consuming and anaerobic breath, lipid digestion, the prompt and persevering hypoxic reactions. It is evaluated that *PhoPR* results in a

decreased capacity to duplicate inside macrophages and changes in the extent of acyl types of mannosylated lipoarabinomannan (manLAM), a basic cell divider part. ManLAM is known to assume a key part in pathogenicity and immunemodulation, recommending that *PhoP* is a key controller of harmfulness. The weakening brought about by erasure of the *PhoPR* is severe to the point, that cancellation strains are under thought in current immunization strain trials. In examination, generally little is thought about *PhoR*, the HK sensor of this complex. *PhoP* and *PhoR* are co-transcribed and the operon is decidedly auto-managed with *PhoP* authoritative to direct rehashes in its own promoter. *PhoR* is anticipated to be a basic film protein with an outside area required in detecting outer signs.

The particular signs to which *PhoR* reacts are obscure, in spite of the fact that *PhoPR* flagging was as of late connected to pH detecting, since it controls the declaration of the AprABC (Acid and Phagosome Regulated) locus in *M. tuberculosis*. In addition, recent work uncovered that a SNP found in *PhoR* in *Mycobacterium bovis* disengages brings about pleiotropic consequences for lipid generation, discharge and harmfulness.

The *M. tuberculosis PhoP* quality has been appeared to encode one of the segments of a TCS (*PhoP/R*) assuming a noteworthy part in virulence. Inactivation of *PhoP* results in a weakened mutant not able to duplicate in creature models and in cells refined in vitro, yet ready to persevere, dissimilar to auxotrophs. This prompted the development of live hopeful antibodies against tuberculosis in view of the inactivation of *PhoP*. Investigations of the genomes of known avirulent strains, for example, the BCG antibody or the H37Ra lessened clinical variation, indicated transformations in *PhoP* or *PhoR* bringing about lost capacity, accordingly affirming the part of *PhoP/PhoR* in virulence.

The *M. tuberculosis PhoP* controller has a place with the *PhoB/OmpR* subfamily, the biggest one among reaction controllers. Individuals from this subfamily have two areas, a N terminal administrative space and a C terminal DNA binding area (likewise called the effector area). Numerous individuals from this subfamily have been examined in detail, and most reports have demonstrated that these reaction controllers tie DNA as a dimer perceiving couple rehash restricting locales at the 35 district of the managed promoter.^[8]

2.6 Epidemiology of *M. fortuitum* and clinical significance :

2.6.1 Mycobacteria fortuitum:

Mycobacteria other than *Mycobacterium tuberculosis complex* and *Mycobacterium leprae* are by and large alluded to as Non Tuberculous Mycobacteria (NTM). Human sickness coming about because of NTM contamination is ordered into four particular clinical disorders: aspiratory infection, lymphadenitis, cutaneous malady, and dispersed illness. Among these, unending aspiratory malady is the most widely recognized limited clinical condition. Thinks about have exhibited that sickness inferable from NTM is on the ascent. NTM are in charge of an expanding extent of mycobacterial illness in many created and creating nations.^[8]

Non Tuberculous Mycobacteria are omnipresent creatures and are much of the time disconnected from natural sources, including surface water, tap water, and soil. In like manner, the segregation of NTM species from a respiratory example is lacking confirmation for the nearness of NTM lung sickness. A few patients are contaminated with NTM without confirmation of aspiratory illness. Such disease may demonstrate colonization or transient contamination. *Mycobacterium fortuitum* is one of the quickly developing mycobacteria (RGM), which are recognized from other NTM by their capacity to frame settlements in under one week and there in vitro imperviousness to antimycobacterials. *M. fortuitum* is a generally detached creature from respiratory examples in clinical research facilities in numerous nations. To date, in any case, the clinical importance of this living being has not been all around examined.^[9]

M. fortuitum contamination can be a nosocomial (healing center obtained) illness. Surgical locales may end up noticeably tainted after the injury is uncovered straightforwardly or by implication to sullied tap water. Other conceivable wellsprings of *M. fortuitum* disease incorporate embedded gadgets, for example, catheters, infusion site abscesses, and debased endoscopes. Late production on Rapidly developing Mycobacteria (RGM) is accessible gives the accompanying parts of RGM: (i) its sources, inclining variables, clinical signs, and associative contagious contaminations; (ii) the dangers of misdiagnoses in the administration of RGM diseases in dermatological settings; (iii) the conclusions and results of treatment reactions in like manner and remarkable contaminations in immuno-compromised and immuno-competent patients; (iv) traditional versus flow sub-atomic strategies for the location of RGM; (v) the fundamental standards of a promising MALDI-TOF MS, inspecting convention for cutaneous or subcutaneous sores and its potential for the exact separation of *M. fortuitum*, *M. chelonae*, and *M. abscessus*; and (vi)

changes in RGM disease administration as depicted in the current 2011 Clinical and Laboratory Standards Institute (CLSI) guidelines, including interpretation criteria of molecular methods and antimicrobial drug panels and their break points [minimum inhibitory concentrations (MICs)], which have been highlighted for the initiation of antimicrobial therapy.^[10]

2.6.2 Clinical significance :

M. fortuitum and *M. chelonae* have been one of the reason for abscesses and post surgical injury diseases.^{[11].} They are spreaded by airborne, clean or polluted faucet water. In one review the post agent wounds had been at first mended agreeably after surgery however it was seen after a time of 12-28 days they ended up noticeably erythematous and began releasing discharge in little amount which later handed overflowing and serous over character. These injuries did not have impact of anti-toxins on them.

Twisted contaminations after operation set aside some opportunity to show up, it by and large happened as talked about before when the operation scar separates and a non-recuperating shallow ulcer creates with releasing sinus.^[12] Sethi et al announced seven patients with M. fortuitum diseases post laproscopic tubectomies.^[13] There was development of mellow inconvenience, induration, with/without neighborhood torment, swelling and sero-sanguineous release which happened from a moment opening over the worked scar for past 2-3 weeks, after the onset of the disease. Suction examples demonstrated no living being on gram stain were sterile for oxygen consuming and anaerobic creatures. Subsequently, all these sterile examples must be recolored by ZN strategy for corrosive quick bacilli (AFB) and subjected to culture on LJ media^[14] Surgical site contaminations because of *M. fortuitum* are all around reported, and are extraordinarily connected with cardio thoracic surgery.^[15] Deferred wound mending, chronicity of disease with delayed course of costly anti-toxins makes it a genuine nosocomial contamination. The wellspring of disease is mostly sullying of twisted because of any methods. On the off chance that there is appropriate observation of natural culture from faucet water, bowl and operation theater (OT) it may not diminish development of NTM. Some healing center gained diseases or say nosocomial contaminations with this living being incorporate diseases of embedded gadgets extraordinarily catheters and infusion site abscesses. There ought to be appropriate and strict cleansing of all OT hardware and legitimate hand washing, purification and offices must be under taken to avert wound contaminations. To begin with line anti-tubercular drugs like ethambutol and rifampicin have

a cidal impact against the living being however as a rule blend of antibacterial operators like amikacin, fluorinated quinolones, doxycycline, imipenem and clarithromycin are utilized.^[16] In India, there is requirement for high level of doubt for recognizable proof of *M. fortuitum* as few instances of *M. fortuitum* contaminations are being under announced because of low doubt of its nearness among clinicians and microbiologists.^[17] Particularly in instances of constant post agent wounds. The quantity of diseases brought on by *M. fortuitum* is on the ascent.^[16] A few clinical elements like osteomyelitis, catheter contaminations, postsurgical diseases, peritonitis and aspiratory diseases have additionally been accounted for as signs of its disease. The shortcoming of one's resistant framework can likewise impact the hazard calculate of a man getting an illness because of *M. fortuitum* presentation. The primary target gatherings of mycobacterioses (malady because of atypical mycobacteria) are the elderly, and the HIV-contaminated with traded off insusceptibility. Along these lines, individuals with stifled insusceptible framework have a more prominent possibility of building up a contamination. Intrigue has been kept up in the astute "atypical" mycobacteria in connection to their capacity to bring about illness in the immuno-compromised people, and progressively in their conceivable part in adjusting invulnerable reactions like the major pathogenic species.^[18]

2.7 Pathogenesis of *M. fortuitum* :

M. fortuitum contamination can bring about different clinical disorders. It is an exceptional reason for NTM lung infection. Neighborhood cutaneous sickness, osteomyelitis, joint contaminations, and visual ailment (eg, keratitis, corneal ulcers) may happen after injury *M. fortuitum* disease is an uncommon reason for separated lymphadenitis. Dispersed malady, more often than not with scattered skin sores and delicate tissue sores, happens solely in the setting of serious immuno-suppression, particularly AIDS. Endocarditis has been recorded.

Surgical-site diseases because of *M. fortuitum* contamination are very much reported, particularly in relationship with cardiothoracic surgery. The source is as often as possible tainting of the injury, specifically or by implication, with colonized faucet water. Other nosocomial diseases with this living being incorporate contaminations of embedded gadgets (e.g. catheters, etc.) and infusion site abscesses. Pseudo-flare-ups have been related with debased endoscopes. Late episodes have additionally been depicted in immuno-competent has after utilization of debased whirlpool footbaths in nail salons.^[19]

There are three critical perceptions that are made in regards to pathogenesis of NTM diseases:

1. Disseminated NTM contaminations happened in patients who were tainted by HIV after the estimation of CD4 T-lymphocyte esteem diminishes beneath 50/l, proposing that particular T-cell items or exercises are required for mycobacterial resistance.^[20,21]

2. In second gathering where patients were HIV-uninfected, transformations in interferon (IFN)- γ and interleukin (IL)- 12 blend and reaction pathway. (IFN- γ receptor 1 [IFN γ R1], IFN- γ receptor 2 [IFN γ R2], IL-12 receptor β 1 subunit [IL12R β 1], the IL-12 subunit p40 [IL12p40], the flag transducer and activator of interpretation 1 [STAT1], and the atomic variable $\beta\beta$ fundamental modulator [NEMO]) have been related with hereditary disorders of scattered NTM.

3. There is likewise a relationship between bronchiectasis, nodular aspiratory NTM contaminations and a specific body habitus, transcendently in postmenopausal ladies (e.g., pectus excavatum, scoliosis, mitral valve prolapse)^[18]

Non tuberculosis mycobacteria are known to bring about illnesses in immuno skilled people too. The instruments by which they taint and remain in macprophages are not that outstanding. So in one review THP-1 macrophages were contaminated with *M. fortuitum*, *M. celatum*, *M. abscessus* and *M. tuberculosis*. Comes about demonstrated that moderate developing mycobacteria got access to the THP-1 macrophages in contrast with other quickly developing mycobacteria. It can be along these lines presumed that quickly developing microorganisms were effectively phagocytosed. Reasonability (CFUs/ml assurance) test was performed to check whether intracellular mycobacteria were alive. Quickly developing microbes indicated 2-log of reduction in CFUs of *M. abscessus* display inside THP-1 macrophages . Number of *M. fortuitum* stayed same between various conditions however it expanded by 2-log from 6h to 48h post disease. Quickly developing mycobacteria likewise harmed monolayer trustworthiness of THP-1 macrophage. Uprightness of THP-1 macrophage monolayer diminished when *M. abscessus* and *M. fortuitum* were available. It was likewise watched that moderate producers did not adjusted morphology of THP-1. Along these lines obviously macrophage monolayer was harmed at 48h post contamination.

Quickly developing mycobacteria instigated the generation of ROS by THP-1 macrophages at 6 and 24 h post contamination. Conversely, no ROS could be recognized with the moderate developing mycobacteria, *M. celatum* and *M. tuberculosis*, even after 24 hour of

contamination. It is plausible that the large amounts of ROS prompted by quickly developing mycobacteria could be related with the broad cell harm.

Cytokines assume vital part in regulating resistant reaction and furthermore in assurance obviously of disease. After time of 24 hours quickly developing microbes indicated abnormal state of cytokine. The quickly developing mycobacteria and *M. celatum* incited the generation of elevated amounts of IL-8 at 6 and 24 h post contamination, while no perceptible levels of this cytokine were found within the sight of *M. tuberculosis*.

Macrophage obliteration can likewise be an outcome of a strong generation of receptive oxygen species (ROS) in THP-1 cells tainted with M. abscessus and M. fortuitum, as has been accounted for beforehand with various host cell/pathogen frameworks. These comes about affirm the theory that quickly developing mycobacteria initiate an extremely intense safe reaction when contrasted with run of the mill pathogenic mycobacteria, (for example, *M. tuberculosis*) This review gives prove that development rate may be connected, sometimes, to the intracellular survival of mycobacteria and the safe reaction that they incite in THP-1 macrophages. Development rate, be that as it may, is not by any means the only determinant of the result of the cooperation of mycobacteria macrophages; different elements, for example, envelope cell lipids and the specific destructiveness elements of every mycobacterium ought to be additionally considered. There is a recommendation that the capacity to square responsive oxygen species generation by moderate developing mycobacteria is an invulnerable avoidance methodology that putatively advances their survival and cytokine creation in the host, even in NTM species. At long last, the information gives understanding into the novel components that *M. celatum* utilizations to hold on inside its host cell, which ought to further be described keeping in mind the end goal to pick up learning about the pathogenic NTM species that cause sickness in immuno able patients. M. fortuitum lives significantly in soil and water and can flourish even in most antagonistic conditions. Human contaminations because of this microorganisms have been found in each segment of the world this is on account of it is discovered wherever and duplicates when supplement substance and temperature are at right level. High centralization of nourishment is a bit much for improvement of *M. fortuitum* rather high temperatures upgrade nearness of these microscopic organisms. The way that it is observed in soil turns out to be valuable since it helps in the breakdown of natural matter in soil, in this way recharging the dirt with supplements.^[23,24]

2.8 Virulence of *M. fortuitum* :

"Virulence" is gotten from the Latin words "infection" ("harm") and "lentus"("fullness") signifying "loaded with toxin", and the expression "infection" might be identified with the Sanskrit word "visham," signifying "harm". The term virulence is utilized to describe the relative limit of a microorganism to bring about infection. It is the normal for harmfulness that recognizes pathogens, from non-pathogens. It is the capacity of a microorganism to bring about ailment because of the statement of specific microbial qualities. Such qualities, or virulent elements, have been characterized traditionally as segments of a pathogen that hinder destructiveness when erased, yet not feasibility. Microbial characteristics, for example, the container of *S. pneumoniae*, the poisons of *C. diphtheriae* and *Vibrio cholerae*, and the M protein of gathering *A. Streptococcus*, are reliable with this definition.

As it was realized that *M. fortuitum* is pathogenic in this way there are numerous destructiveness calculates yet to be found. It goes under family "Mycobacterium" which additionally incorporates *Mycobacterium tuberculosis*, the subsequent specialist of tuberculosis. Studing the pathogenicity of its assorted is undoubtly testing. Destructiveness being the fundamental component of pathogenicity can be measured at the season of disease of macrophages and creatures, utilizing different examines and different procedures can be upgraded to produce mutants as in *M. tuberculosis*. Thus, diverse methodologies can be joined in same route as of plenty to recognize virulence determinant of *M. fortuitum*.

A superior instrument TnphoA is utilized to produce influenced mutants in secretory or film crossing proteins. Replication of TnphoA contains transposon of Tn5 which encodes kanamycin and promoter less basic Gene of E.coli alkane phosphates which needs flag peptide arrangements.^[25] Antacid phosphatase is dynamic just when sent out on the cell surface. Action of antacid phosphate is not distinguished until and unless the coordination of TnphoA has happened in casing into the qualities coding for secretory proteins. It is a superior columnist framework for the determination of clones having joining in secretory qualities. TnphoA is a framework accommodated insertional mutagenesis of secretory proteins and contemplating of these proteins as harmfulness determinant(s). Since by and large all bacterial proteins involved as destructiveness variables are extracellular, surface related, or periplasmic, consequently utilization of TnphoA ought to give a solid enhancement to inclusion changes that influence pathogenic properties of microscopic

organisms. Utilizing the transposon $in^{[25]}$ ought to bring about the recognizable proof of layer or secretory proteins which assumes a part in destructiveness. Distinguishing proof of qualities of *M. fortuitum* brings about the ID of their homologues in *M. tuberculosis* as appeared with other atypical mycobacterial strains like *M. marinum* and *M. smegmatis*.

A murine disease demonstrate for *M. fortuitum* indicated nearness of consistent CFU over broadened timeframes which was created in one review where the model was abused for the distinguishing proof of destructiveness determinant of tireless in *M. fortuitum*. As these microscopic organisms are less pathogenic and are quickly developing when contrasted with other moderate developing Mycobacteria, it give preferred standpoint of working effectively with it. This model and demonstrated bacterial load to be consistent over developed timeframe:

• Immunological profiling (Thl and Th2) of the model at different phases of pathogenesis was done, including reactivation.

• Histo-pathological examination of kidney tissues at various phases of disease

• Identification of quality succession influencing harmfulness of *M. fortuitum* in the wake of screening of library of TnphoA mutants in the murine disease model was done it prompted distinguishing proof of an extremely lessened mutant.

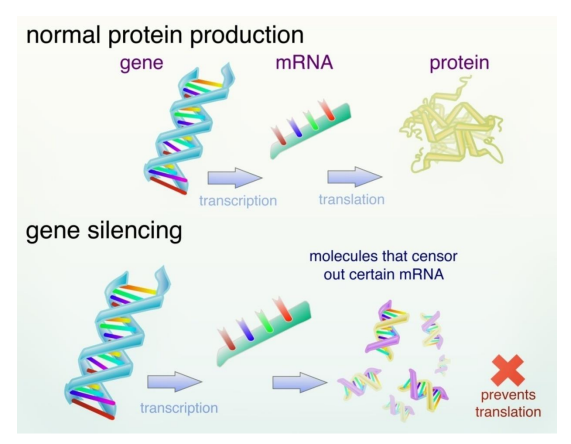
Cloning and sequencing of the disturbed quality taken after by complementation examination of the mutant indicated inclusion of the upset quality in destructiveness and constancy of M. *fortuitum*, bearing abnormal state of homology with administrative grouping Rv3291c of M. *tuberculosis*. Contemplates on destructiveness in mycobacteria other than the M. *tuberculosis* complex have been completed before and homologues of M. *ulcerans* toxin and M. *avium* subsp. avium mig (macrophage initiated quality) in M. *tuberculosis* have been effectively distinguished.^[26]

Focal points in distinguishing virulent components incorporate the arrangement of focuses for new potential therapeutics and the recognizable proof of qualities that can be changed to permit the advancement of new immunizations. At a more basic level, the distinguishing proof of bacterial destructiveness figures and additionally the host elements required at various phases of contamination and illness is basic for the explanation of mycobacterial pathogenesis. Improvement of a reproducible and solid model for the

investigation of industrious disease ought to demonstrate important data towards understanding the science of mycobacterial idleness.

2.9 Gene Silencing and Knock Out :

Gene silencing: Gene silencing is by and large used to portray direction of gene expression. A specific cell averts articulation of certain quality this is characterized as Gene Silencing. For research planned quality quieting can happen either amid interpretation or interpretation. It is thought to be like quality knockout however it has some distinction for sure. Amid quality knockout quality is totally expelled from genome and have no expression yet in gene silencing, expression is for the most part decreased. Hence gene silencing is considered as quality knockdown.



(Figure 2.2 : Mechanism of gene silencing)^[34]

<u>Gene knockout</u>: A gene knockout is a hereditarily built creature that conveys at least one genes in its chromosomes that have been made out of commission (have been "knock out" of the organism). This is accomplished for research purposes. Otherwise called knockout living beings or just knockouts, they are utilized as a part of finding out about a gene that has been sequenced, however which has an obscure or not completely known capacity. Scientists draw

derivations from the contrast between the knockout life form and ordinary people. The term likewise alludes to the way toward making such a living being, as in "knocking out" a gene. Knockout is expert through a mix of systems, starting in the test tube with a plasmid, a bacterial artificial chromosome or other DNA construct, and continuing to cell culture. Singular cells are hereditarily changed with the build and for knockouts in multi-cell life forms - at last intertwined with an undifferentiated cell from an early incipient organism. The construct is designed to recombine with the objective gene, which is proficient by joining arrangements from the gene itself into the develop. Recombination then happens in the locale of that arrangement inside the gene, bringing about the inclusion of a foreign gene to disrupt the gene. With its sequence intruded on, the altered gene by and large will give rise to a nonfunctional protein, in the event that it is interpreted or translated by any means. A restrictive knockout permits quality cancellation in a tissue particular way. Since recombination is an uncommon occasion on account of most cells and most constructs, the sequence decided for insertion is usually a reporter. This empowers simple determination of cells or people in which knockout was effective.

Gene Knockout-Directed Gene Disruption Directed gene inactivation involves the inclusion of an antibiotic resistance cassette in center of the gene of interest, trailed by allelic substitution of the chromosomal gene by the transformed one, bringing on disturbance of the gene capacity in the host cell. Gene knockout is normally a two-stage handle including incorporation of plasmid containing desired disrupted gene by a solitary hybrid occasion (Campbell-sort combination) into the genome at the area of homology. In the second hybrid occasion, the plasmid spine is extracted by methods for recombination, bringing about the desired gene disruption. Another strategy for directed gene inactivation, named specific transduction, utilizing a bacteriophage delivery system has been portrayed and is the present technique for decision for the directed mutation.^[27] This system is exceptionally productive in light of the fact that basically the majority of the beneficiary cells can be transduced and the selection is powerful.

2.10 Antisense Technology:

In Antisense technology, artificially – created corresponding atoms search out and tie to messenger RNA (mRNA), hindering the last stride of protein generation. mRNA is the nucleic acid molecule that conveys hereditary data from the DNA to the next cell apparatus required in the protein generation. By Binding to mRNA, the antisense drugs hinder and

restrain the generation of particular illness related proteins. "Sense" alludes to the first arrangement of the DNA or RNA particle. "Antisense" alludes to the reciprocal grouping or complementary sequence of the DNA or RNA molecules.

The fundamental thought is that if an oligo-nucleotide (a short) RNA or DNA atom integral to a mRNA created by a quality) can be brought into a cell, it will particularly tie to its objective mRNA through the stunning specificity of correlative based blending a similar mechanism which ensures the constancy of DNA replication and of RNA translation from the gene. This coupling shapes a RNA dimmer in the cytoplasm and stops protein synthesis. This happens in light of the fact that the mRNA no longer has entry to the ribosome and cytoplasm by ribonucleotide. Along these lines, the presentation of short chain of DNA reciprocal to mRNA will prompt a particular decrease, or blockage, of protein blend by a specific gave. In actuality, the gene will be turned off.^[28] A Sense strand is a 5' to 3' mRNA atom or DNA particle. The complementary strands or mirror strand to the sense is called an antisense. Antisense technology is the procedure in which the antisense strand hydrogen bonds with the focused on sense strand. At the point when an antisense strand ties to a mRNA sense strand, a cell will perceive the twofold helix as outside to the cell and continue to debase the faculty mRNA particle in this way keeping the generation of undesired protein. Despite the fact that DNA is now a twofold stranded protein, antisense innovation can be connected to it constructing a triplex development. A DNA antisense molecule must be around seventeen packs to function, and roughly thirteen bases for a RNA particle RNA antisense strands can be either catalytic, or non catalytic. The catalytic antisense strands, likewise called ribozymes, which will cleave the RNA molecule at specific sequences. A Non catalytic RNA antisense strand squares assist RNA preparing, i.e. changing the mRNA strand or translation. The correct mechanism of an antisense strand has not been resolved. The present theories incorporate. "Blocking RNA grafting, accelerating degradation of the RNA particle, and keeping introns from being spliced out of the mRNA, obstructing the exportation of mRNA into the cytoplasm, hindering translation, and the triplex arrangement in DNA.^[29]

Antisense technology includes utilization of complementary RNA, which ties to the mRNA of a particular gene along these lines preventing its translation. Antisense technology is especially helpful for silencing of fundamental genes as their knockout mutants can't survive, and in frameworks where gene inactivation is troublesome. Regulatable acetamide/acetamidase framework is for the most part utilized for restrictively controlling the generation of antisense RNA in mycobacteria. Uses of the antisense strategy were in *M bovis* to bring down the levels of AhpC and in *M. tuberculosis* H37Rv to decrease the measures of *SodA*. As of late, the antisense approach was utilized to diminish the level of *sigA*, and Rv3303c in *M. tuberculosis*.

Chapter 3 Materials And Methods

lipF gene with full length of 927 base pair having an open reading frame (orf) of 729 base pair is present in the genome of *Mycobacterium tuberculosis*. The confirmation of this gene in *M. fortuitum* is done by PCR. Multiple sets of primers were designed from flanking region of *lipF* constant domain regions (CDS) and then PCR was done using the *M. fortuitum* genome. The PCR products were sequenced and checked for the homology with the *lipF* of *M. tuberculosis*. After doing the homology search, the longest sequence having the maximum homology was determinded which is yet to be submitted to NCBI. *pMV261* vector was used to construct the antisense mutants of *M. fortuitum lipF*. The orientation is to be confirmed using *BamHI* site in *pMV261* vector. Presence of full length gene and its orientation was confirmed by restriction digestion and colony PCR using the primers designed. Obtained sense / antisense mutants are yet to be electroporated in *M. fortuitum*.

3.1 Primer designing for *lipF* sequence of *M. tuberculosis* :

Primers were designed using following method manually:

- 1. Full genomic sequence of *lipF* gene of *M. tuberculosis* was taken and full length primers of *M. tuberculosis* were designed using software primer 3.
- 2. For the forward primer, nucleotides of the sequence for the region that binds to the ORF was taken and full length primer was designed.
- 3. Similarly, for reverse primer designing, reverse complement of the bases were taken to get the PCR amplification including the stop codon using software primer 3.

3.2 Genomic DNA isolation of *M. fortuitum* :

Chemical requirements: TES buffer, lysozyme, buffer phenol, chloroform: isoamylalcohol, isopropanol (chilled), ethanol, autoclaved distilled water, culture of *M. fortuitum*

Equipments used: Centrifuge, centrifuge tubes, micro-centrifuge tubes, incubator, discard box, gel doc, pipettes, and tips.

Procedure:

- 1. The culture was taken and pelleted in 50 ml centrifuge tube at 7000 rpm for 15 minutes.
- 2. The supernatant was discarded and 2 mL TES buffer was added to pellet.
- Then it was incubated at 80^oC for about 1 hour. After that 2mg/ml of lysozyme was added to it i.e. 80μL and incubated at 37^oC for 1 hour.

- 4. 1.5 % SDS and 100 μ g/mL proteinase K was added and incubated at 50^oC for 1 hour.
- 5. Buffer phenol was added in each tube and they were centrifuged at 12000g for 15 minutes.
- 6. Then chloroform: isoamylalcohol was added in 24:1 ratio in each tube and centrifuged them at 12000g for 15 minutes.
- 7. The aqueous layer was transferred to fresh tubes. Again chloroform: isoamylalcohol was added and centrifuged at 12000g for 10 minutes.
- 8. The aqueous layer was transferred to fresh tubes and equal volume of chilled isopropanol was added.
- 9. It was incubated at 4° C for overnight and then centrifuged at 12000g for 15 minutes. Supernatant was removed and 1mL 70% ethanol was added and centrifuged at 12000g for 15 minutes.
- 10. Again, supernatant was discarded and ethanol was evaporated. 100 µL of autoclaved distilled water was added.
- 11. It was then placed in ice for about 2 hours for proper suspension of DNA.
- 12. Finally, electrophoresis was done to check the DNA in 0.8% agarose gel.

3.3 PCR amplification of genomic DNA of *M. fortuitum* by full length *lipF* primer of *M. tuberculosis*:

Chemical requirements: Isolated genomic DNA of M. fortuitum, forward primer, reverse primer, green master mix, nuclease free water.

Equipments used: Thermocylcer, spin, PCR - vials, discard box, gel doc, pipettes, and tips. Procedure :

1. PCR reaction was carried out with the following mixture:

-	gDNA	- 2.0 µL
-	Forward Primer	- 0.5 μL
-	Reverse Primer	- 0.5 μL

- Master Mix - 5.0 µL
- Nuclease free water $-2.0 \,\mu L$ T (1 10 I

2. PCR conditions were kept as follows:

-	Initial Denaturation	- 95 ⁰ C
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- Denaturation $-95^{\circ}C$
- Annealing $-58^{\circ}C / 60^{\circ}C / 62^{\circ}C$
- Extension $-72^{\circ}C$
- Final Extension $-72^{\circ}C$

Total number of cycles -35

3. Electrophoresis of the PCR product was done on 1.5% agarose gel.

3.4 Primer designing for orf region of *lipF* sequence of *M. fortuitum* :

For PCR based plasmid cloning, primers were designed using following method manually:

- 1. For the forward primer, primers were designed using software Primer 3 and *BamHI* restriction site (GGATCC) was added to the 5' end of this primer.
- Similarly, for reverse primer designing, again primers were designed using software Primer 3. Finally *BamHI* restriction enzyme was added to the same but in a reverse complement order.

3.5 PCR amplification (gradient) of genomic DNA of *M. fortuitum* using designed *lipF* primers :

Chemical requirements: Isolated genomic DNA of *M. fortuitum*, forward primer, reverse primer, green master mix, nuclease free water.

Equipments used: Thermocylcer, spin, PCR - vials, discard box, gel doc, pipettes, and tips. Procedure :

- 1. PCR reaction was carried out with the following mixture:
 - gDNA 2.0 μL
 - Forward Primer $-0.5 \ \mu L$
 - Reverse Primer $-0.5 \,\mu L$
 - Master Mix $-5.0 \,\mu L$
 - Nuclease free water 2.0 μL Total - 10 μL
- 2. PCR conditions were kept as follows:

-	Initial Denaturation	- 95 ⁰ C
-	Denaturation	- 95 ⁰ C
-	Annealing	- 58°C / 60°C / 62°C
-	Extension	- 72 ⁰ C

- Final Extension - 72^oC

Total number of cycles -35

3. Electrophoresis of the PCR product was done on 1.5% agarose gel.

3.6 TA Cloning of the amplified PCR product of *lipF* of *M. fortuitum* :

Chemical requirements: Amplified PCR product of *M. fortuitum*, DNA Ligase, pGEM-T easy vector, buffer, nuclease free water.

Equipments used: Thermocylcer, spin, vials, discard box, gel doc, pipettes, and tips.

Procedure :

1. Cloning reaction of the PCR product was carried as follows :

-	pGmEt easy	-1μL
-	Buffer	- 5 µL
-	Nuclease free water	-1μL
	Total	- 10 µL

2. The above reaction was kept at 16° C for overnight.

3.7 Transformation of ligated product into competent DH5a E.coli cells :

3.7.1 Preparation of competent cells :

Chemical requirements: Transformation buffer-1 (RbCl, MnCl₂.4H₂O, Potassium acetate, CaCl₂.2H₂O, 15% glycerol), transformation buffer-2 (MoPs, RbCl, CaCl₂.2H₂O, 15% glycerol), LB.

Equipments used: Centrifuge, PCR-vials, micro-centrifuge tubes, vortex, incubator, discard box, gel doc, pipettes and tips.

Procedure:

- Single colony was picked and inoculated in 10 mL LB. It was incubated at 37^oC for overnight.
- 1 ml of overnight culture was added to 100 mL pre-warmed LB medium and incubated with constant shaking at 37^oC.
- 3. The culture was cooled in ice for 5 minute and transferred to sterile round bottom centrifuged tube.
- The cells were collected by centrifuged at low speed 4000g for 5 minute at 4^oC.
 Supernatant was carefully discarded. The cells were always kept in ice.
- 5. The cells were gently re-suspended in transformation buffer 1 and kept on ice for an additional 90 minute.
- Cells were collected by centrifuging for 5 minute at 4000 rpm at 4^oC. The supernatant was discarded.
- 7. The cells were then carefully resuspended in 4 ml ice cold transformation buffer 2.
- 8. Aliquots of 100-200 μ L were prepared in sterile centrifuge tubes and stored at -80^oC.

3.7.2 Transformation of TA cloned genomic DNA with competent cells of *E.coli* :

Chemical requirements: Ligated product, competent cells, LB.

Equipments used: PCR - vials, micro-centrifuge tubes incubator, water bath, petri plates. Procedure:

- 1. Competent cells were thawed in ice. 15μ L of ligated product was added in 200 μ L of competent cells.
- 2. It was mixed by swirling and incubated in ice for 30 minutes.
- 3. After that, it was incubated in water bath set at 42^oC for 90 minutes. Then, it was chilled on ice for about 5-15 minute.
- 4. Then, 800μ L of LB was added to the vials, and incubated for 45 minute a 37^{0} C.
- 5. After 45 minutes of incubation, the cells were spread on selection plate with appropriate antibiotics.

3.8 Screening and selection of transformed mutants on Ampicillin resistant plates along with the preparation of master plate:

Chemical requirements: LB Agar plates, ampicillin, IPTG, X-gal, transformed mutant cells, distilled water.

Equipments used: Petri-dishes, weighing balance, autoclave, micro-centrifuge tubes, centrifuge, pipette, incubator, tips.

Procedure :

- 1. Culture plates of LB Agar were prepared along with the standard concentration of ampicillin. IPTG and X-gal were spread on the same culture plates.
- 2. 100μ L of transformed cells were spread on LB agar plates.
- 3. Plates were kept in incubator for overnight at 37^{0} C.
- 4. White colonies observed the very next day were transferred to master plate containing same concentration of ampicillin.
- 5. Plates were stored at 4^{0} C.

3.9 Colony PCR of *lipF* transformants :

This step was performed for the confirmation of transformed mutants on the plate.

Chemical requirements: Isolated colony of transformed mutant from the master plate, forward primer, reverse primer, green master mix, nuclease free water.

Equipments used: Thermocylcer, spin, vials, discard box, gel doc, pipettes, and tips. Procedure :

1. PCR reaction was carried out with the following mixture:

-	gDNA -	1 isolated colony
-	Forward Primer	- 0.5 μL
-	Reverse Primer	- 0.5 μL
-	Master Mix	- 5.0 μL
-	Nuclease free water	- 4.0 μL
	Total	- 10 μL

- 2. PCR conditions were kept as follows:
 - Initial Denaturation 95⁰C
 - Denaturation $-95^{\circ}C$
 - Annealing $-58^{\circ}C$
 - Extension 72^oC
 - Final Extension $-72^{\circ}C$

Total number of cycles – 35

3. Electrophoresis of the PCR product was done on 1.5% agarose gel.

3.10 Plasmid isolation from the *lipF* transformants from master plate :

3.10.1 Inoculation of seed culture followed by primary culture :

Chemical requirements: LB nutrient media, distilled water, ampicillin, *lipF* transformed colony from master plate.

Equipments used: Flask, test-tubes, autoclave, incubator, petri-plates, pipette, tips.

Procedure:

- 1. Seed culture was inoculated in 10mL LB nutrient media with one isolated colony of *lipF* transformant taken from the master plate.
- 2. Test tubes were kept at 37^{0} C for overnight growth.
- 3. Next day, 1mL of the growth culture was inoculated in 100mL LB nutrient media in two flasks separately.
- 4. After overnight growth, plasmid was isolated manually and from kit from the cultures.

3.10.2.1 Plasmid isolation from *lipF* transformants primary culture (Manually) :

Chemical requirements: Primary culture, GTE (Glucose tris EDTA), ALS (Alkaline lysis buffer), SDS-NaOH solution, acetate solution, isopropanol, 96% ethanol, 70% ethanol, autoclaved distilled water.

Equipments used: Centrifuge, micro-centrifuge tubes, discard box, pipette, tips.

Procedure :

- 1. The culture was centrifuged at 7000 rpm for 10 minutes.
- 2. Then pellet was dried properly.
- 3. 300µL of GTE (Glucose tris EDTA), (ALS I) was added and vortexed.
- 4. 600µL of SDS-NaOH (ALS II) solution was added and mixed gently.
- 5. It was then incubated at 37^{0} C for 5 minutes.
- 6. 450µL of acetate solution (ALS III) was added.
- 7. It was gently mixed for 5-6 times.
- It was then incubated in ice for 30 minutes and centrifuged at 7000 rpm for 25 minutes at 4⁰C.
- Supernatant was transferred to the fresh eppendorf or tarson tube and 270µL of isopropanol was added (or in 1:1 ratio)
- 10. Samples were incubated at -20° C for overnight.
- 11. The next day, samples were centrifuged at 12000 rpm for 15 minutes at 4^{0} C.

- 12. Supernatant was discarded and 1mL of 96% ethanol was added to each sample.
- 13. Centrifugation was done at 10,000 rpm for 12 minutes at 4^oC and again supernatant was discarded.
- 14. 1 mL of 70% ethanol was added to each sample and centrifuged at 10,000 rpm for 5 minutes at 4°C.
- 15. Supernatant was removed and 1mL of 70% ethanol was added to each eppendorf again.
- 16. Centrifugation is done at 10,000 rpm for 3 minutes at 4^oC and the pellet is dried properly afterwards to evaporate ethanol.
- 17. 50μL of distilled autoclaved water was added to each sample and kept in ice for 1-2 hours for proper suspension of plasmid.
- 18. Plasmid from all micro-centrifuge tubes are pooled together and stored at 20° C.
- 19. Electrophoresis was done in 0.8% agarose gel.

Similarly, plasmid pMV261 was also isolated (manually).

3.9.2.2 Plasmid isolation from *lipF* transformants primary culture (Using kit):

Chemical requirements: Primary culture, plasmid isolation kit, autoclaved distilled water. *Equipments used*: Centrifuge, vortex, centrifuge tubes, micro-centrifuge tubes, discard box, pipette, tips.

Procedure:

- 1. Primary culture was centrifuged at 7000 rpm for 10 minutes at 4^oC and supernatant was discarded afterwards.
- 2. Pellet was dried properly and dissolved in 500µL of suspension buffer.
- Culture was vortexed and 500µL of lysis buffer was added. The solution was mixed smoothly.
- 4. 650µL of neutralization buffer was added to the solution and mixed properly.
- 5. Centrifugation was done at 7000 rpm for 10 minutes at 4° C.
- Supernatant was transferred into the gel column provided in the kit and centrifuged at 10,000 rpm for 2 minutes at 4^oC.
- 500µL of wash buffer was added to it and again it was centrifuged at 10,000 rpm for two minutes.
- 8. The above step was performed twice.
- 9. Column was again spun at 10,000 rpm for 2 minutes.

- 10. 40µL of autoclaved distilled water was added to it and it was incubated at room temperature for 10 minutes.
- 11. Centrifugation was done at 10,000 rpm for 2 minutes.
- 12. Again 30µL of autoclaved distilled water was added to it and it was incubated at room temperature for 10 minutes for the proper suspension of plasmid in the autoclaved distilled water.
- 13. Centrifugation was done again at 10,000 rpm for 2 minutes.
- 14. Finally, the microcentrifuge tubes were stored at -20° C.
- 15. Electrophoresis was done on 0.8% agarose gel.

3.11 Digestion of *lipF* plasmid and *pMV261* plasmid with *BamHI* and *EcoRV*:

Chemical requirements: lipF plasmid, cut smart buffer, enzymes (*BamHI* and *EcoRV*), nuclease free water.

Equipments used: Vials, spin, ice-box, incubator, discard box, pipette, tips.

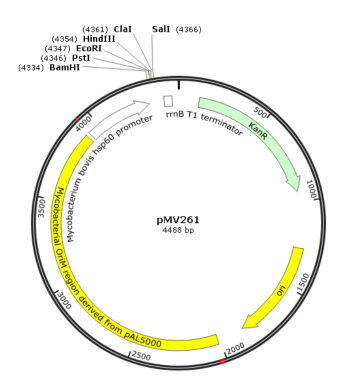
Procedure :

1. The following reaction was prepared in a vial separately for *lipF* and *pMV261* with both the enzymes:

-	Nuclease free water	-	3.5µL
-	Cut smart buffer (10X)	-	1.0µL

- Plasmid 5.0µL
- Enzyme 0.5μL Total - 10.0μL
- 2. The vials were kept at 37C for 3 hours reaction.
- 3. Electrophoresis was done at 0.8% agarose gel.
- 4. After observing successful results, bulk/ mass reaction was prepared with different parameters:
 - Nuclease free water 6.0µL
 - Cut smart buffer (10X) $6.0\mu L$
 - Plasmid 45µL
 - Enzyme 3.0µL
 - Total 60µL

5. Again, the vials were kept at 37°C for 3 hours digestion reaction and stored at -20°C afterwards.



(Figure 3.1 Vector map of pMV261)^[35]

Chapter 4 Results

4.1 Primers designed for the *lipF* full length sequence :

Forward Primer : 5' TTACCASGTGGCCTCT 3' Reverse Primer : 3' ATGCATGCCGATGGC 5'

4.2 Blast results of *lipF* sequence of *M. tuberculosis* and *M. fortuitum* :

Score		Expect Identities	Gaps
232 bits((256)	1e-64 463/696(67%)	29/696(4%)
Query	80	GTGGCCTCTCGAATGTACTCACCGATCTGGCGCAACGAGCGCTTCGCTTCGGCCACCATC	139
Shict	737	GTGGCGTCGCGAATGAACTGCCCGATCTGACGCAACGAACG	678
50)00	151	GIGGCGICGCGAALGAACIGCCCGAICIGACGCAACGAACGGGICGCIICGGGCACCAGC	070
Query	140	GGCGACGCGAGCTGGAATACGTGCATCTGACCGGGCCAGATCCTGAGCTCGACGGGAACG	199
Sbjct	677	GGTGTGGCGAGTTGGAAAAGATGAGCCTGACCGGGCCAAACCCGTACCTCGGCACAGACG	618
Query	200	CCGGCCGCAGCCAGCATGTGCGCGGCTTTCCGCGCATCGCTGAGCAGAGCCTCCGAGCCC	259
Sbjct	617	CCTGCCGCCGCCAGCTTGCCGGCGCCCAGCTGCGCGTCGTGCAGCAGCACTTCGGAGCCG	558
010 714	260	GATGCGTGGATCAACGTGCGAGGCAGACCGGGCTCGATGTGGTCGAGAGGTTCGTAGACG	319
Query	200		519
Sbjct	557	GAAACGTGAATAAGTGTCGGCGGCAAGCTGGATTCGATATGGTCGAGCGGCTCATAGAGG	498
Query	320	TCCTCCCCCTTGCGGGCGGCCGCCC-GCTCGATGAGCTCGAC	360
Sbjct	497	IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	441
2			

Query	361	CAGAGCATCGAAGGCCTTGGGCGGGAACATCGCGTCGGTGTGGATGTTGGGGTGGTTGGC	420
Sbjct	440	AAG-GCATCGAATGCCCGCGCCGGAAACATCGCGTCGGTCCCGATGTTGGGATGGTCCTG	382
Query	421	CCGTGATTCGTTGTCGATCTCGAACAACGGCGACATCGTCWCCAGCGCGGCAGGCATCTC	480
Sbjct	381	I I I III I II I IIIIIII II II I I I I	323
Query	481	TCCCTCRTCG-ACCAGGCGCTCGGCCAGCGCCAGCGACRGGTAGCYGCYCGCCGAGTC	537
Sbjct	322	CGTCGTCGCACTGCAGCCGCTGCGCAAGCGCGAGCGCAAGGTAACCACCCGCGGAATC	265
Query	538	GCCGGYCARCACGATCTGGTCCGGCWGGTARCCGGTCAGCCGCAGCCAGCGGTAGGCGTC	597
Sbjct	264	IIII IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	205
Query	598	GTAACAGTCKTYGATCGCCGTGCCCACCGAGTGCTTGGGCACCATCMKGTAATCGACGAC	657
Sbjct	204	II IIIII I II IIII I IIII III IIII III IIII	145
Query	658	CAGGACCGGGCTGTCGGCATATCCCGACAGCGCCGTGACGATCCCGGAGTGCGTGTTGGC	717
Sbjct	144		85
Query	718	YCCGCWGGMCAKGAACRCAYCGCCGTG <mark>Y</mark> AAG <mark>WW</mark> GAG 753	
Sbjct	84	CCCGCACATGACAAACGCGCCGCCGTGCAAATAGAG 49	
Score		Expect Identities	Gaps
22.9 bit	s(24)	0.096 12/12(100%)	0/12(0%)

```
Query 330 TGCGGGCGGCCG 341
```

4.3 *lipF* sequence of *M. fortuitum* obtained after Sanger sequencing (to be submitted to NCBI):

TTACCASGTGGCCTCTCGAATGTACTCACCGATCTGGCGCAACGAGCGCTTCGCT TCGGCCACCATCGGCGACGCGAGCTGGAATACGTGCATCTGACCGGGCCAGATC CTGAGCTCGACGGGAACGCCGGCCGCAGCCAGCCAGCATGTGCGCGGGCTTTCCGCGCA TCGCTGAGCAGAGCCTCCGAGCCCGATGCGTGGATCAACGTGCGAGGCAGACCG GGCTCGATGTGGGTCGAGAGGTTCGTAGACGTCCTCCCCCTTGCGGGGGGGCGGCCGCC GCTCGATGAGCTCGACCAGAGCATCGAAGGCCTTGGGCGGGAACATCGCGTCGG TGTGGATGTTGGGGTGGTTGGCCCGTGATTCGTTGTCGATCTCGAACAACGGCGA CATCGTCACCAGCGCGGCAGGCATCTCTCCCTCGTCGACCAGGCGCCAGC GCCAGCGACAGGTAGCCGCCGCGCGAGTCGCCGGCCAACACGATCTGGTCCGGC GGCCAGCGACAGGTCAGCCGCCGCGGCAGGCCGCGGCAACACGGCCGCG WGGTARCCGGTCAGCCGCCGCGCGGCAGGCCAGCGCGCGAGCCAGGCCTGG TCGGCCACCGAGTGCTTGGGCACCATCCGGTAATCGACGACCAGGACCGGGCTG TCGGCATATCCCGACAGCGCCGTGACGATCCCGGAGTGCGTGTTGGCCCCGCAG GMCAKGAACGCACCGCCGTGCAAGTAGAGGATGWCGCTGCGTTTCYCATCGRY GGGCAGCACGCCATCGGCATGCAT

According to the IUPAC nomenclature, changes were made in the above sequence:

- 677 nucleotide W replaced with T
- 678 nucleotide W replaced with A
- 673 nucleotide Y replaced with C
- 665 nucleotide Y replaced with C
- 662 nucleotide R replaced with G
- 657 nucleotide K corresponds to C whereas as per IUPAC nomenclature
 K refers to either G or T

- 654 nucleotide M corresponds to T whereas as per IUPAC nomenclature
 M refers to either A or C
- 652 nucleotide W replaced with A
- 646 nucleotide Y replaced with C
- 573 nucleotide K replaced with G
- 572 nucleotide M replaced with C
- 537 nucleotide Y replaced with C
- 535 nucleotide K replaced with G
- 496 nucleotide R corresponds to T whereas as per IUPAC nomenclature R refers to either A or G
- 491 nucleotide W corresponds to C whereas as per IUPAC nomenclature
 W refers to either A or T
- 474 nucleotide R replaced with A
- 471 nucleotide Y replaced with C
- 456 nucleotide Y replaced with C
- 453 nucleotide Y replaced with C
- 446 nucleotide R replaced with A
- 415 nucleotide R replaced with G
- 389 nucleotide W replaced with A

Amino acid sequence for the above nucleotide sequence obtained :

3'5' Frame 1

Met HADGVLPXDXKRSXILYLHGGAFLXCGANTHSGIVTALS GYADSPVLVVDYR Met VPKHSVGTAIDDCYDAYRWLRLTXY XPDQIVLAGDSAGGYLSLALAERLVDEGE Met PAALVT Met SP LFEIDNESRANHPNIHTDA Met FPPKAFDALVELIERAAARKG EDVYEPLDHIEPGLPRTLIHASGSEALLSDARKAAH Met LAAA GVPVELRIWPGQ Met HVFQLASP Met VAEAKRSLRQIGEYIRE AXW Stop

4.4 Primers designed for the *lipF* sequence of *M*. *fortuitum* :

- Forward Primer : 5' **GGATCC**ATGCATGCCGATGGCGTG 3'

- Reverse Primer : 3' **GGATCC**TTACCACGTGGCCTCTCG 5'

4.5 Gradient PCR for the amplification of *lipF* sequence of *M. fortuitum* using manully designed primers :

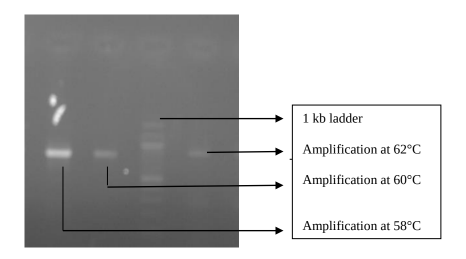


Fig. 4.1 Electrophoresis of gradient PCR amplicons of gDNA isolated from M. fortuitum lipF showed the best amplification at 58°C, hence it was always amplified at 58°C afterwards.

4.6.1 Screening of transformed cells on LB agar plates (ampicillin resistant):

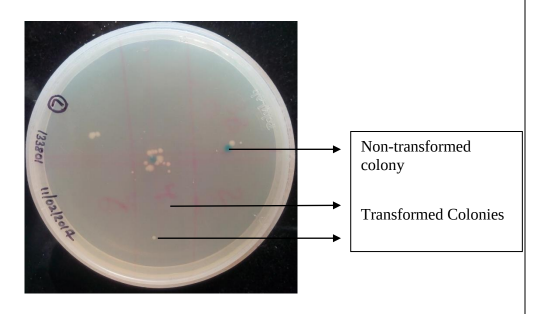


Fig. 4.2 Blue-white screening of transformant mutant strains on LB agar ampicillin plate containing X-gal + IPTG

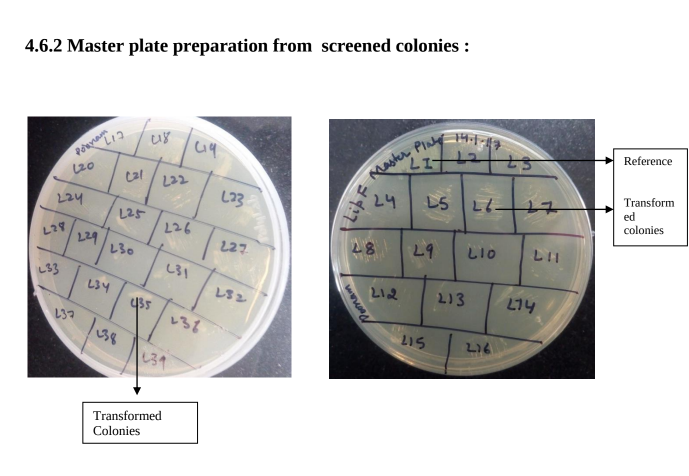
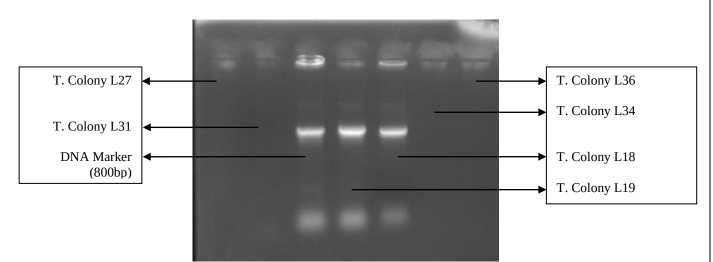


Fig. 4.3 Master plates of transformant mutant strains on LB agar ampicillin plate containing X-gal + IPTG



4.7 Colony PCR of *lipF* transformed cells :

Fig. 4.4 Electrophoresis of colony PCR amplicons of lipF transformed cells

4.8 Plasmid isolation of transformed mutants (*lipF and pMV261*) :

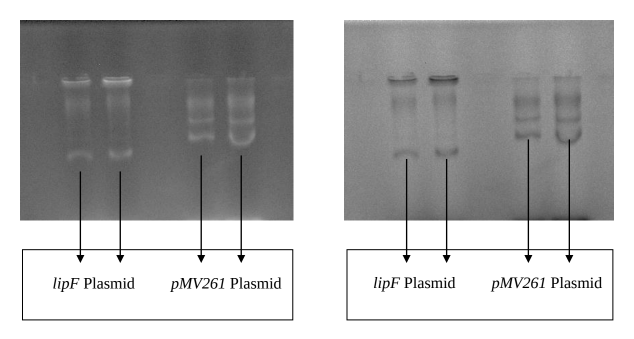
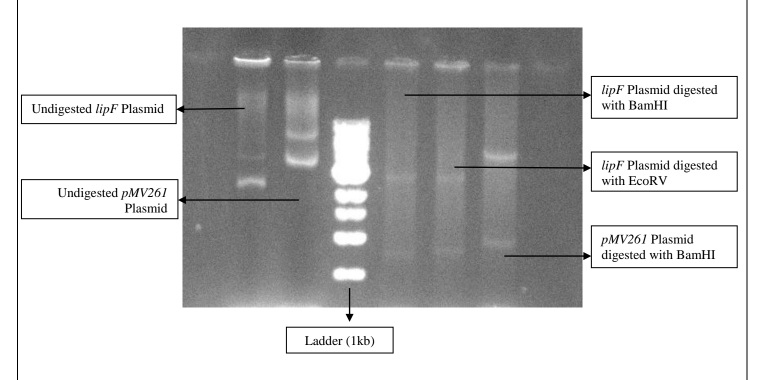


Fig. 4.5 Electrophoresis of isolated plasmid from lipF transformed mutants and pMV261



4.8 Digestion of *lipF* plasmid with *BamHI* and *EcoRV*:

Fig. 4.6 Electrophoresis of digested plasmid of lipF transformant with BamHI and EcoRV and pMV261 with BamHI

Chapter 5 Discussion and Conclusion

Mycobacterium tuberculosis was recognized by Robert Koch in 1882, and *M. fortuitum* was distinguished soon afterwards. It was classified as a Non Tuberculous Mycobacterium (NTM) and until 1954, not a particular study was done over it. Edward Runyon took it upon himself to categorize all the NTMs at this time. *M. fortuitum* is now characterized as Runyon Group IV, which means, among other lab culture attributes, it is a comparatively rapid grower.

Tuberculosis is one of the greatest threat to mankind even in today's era as no effective vaccine or antibiotic has been produced yet to cure this disease completely. And the only single pathogen responsible for causing the disease is *Mycobacterium tuberculosis*. This single pathogen has been showing a lot of changes in its system which has led to the emergence of multiple drug resistant strains followed by extensively drug resistant strains. In a country like India, where public health sector is yet in the emerging stage, need of the hour is to study the various pathogenic mechanisms showed by this bacteria in order to end this disease as soon as possible. On the other hand, *Mycobacterium foruitum* infection in healthy humans is rare, but exposure to large and repeated amounts of the organism can overwhelm the immune system and cause disease. Infections most likely occur in immunocompromised patients. Mortality is very rare, but death may come from extensive pulmonary or disseminated disease in immune-compromised patients.

This Project study shows the importance of lipase esterase gene in the pathogenesis of tuberculosis. lipF is an important gene required for pathogenesis and persistence of *Mycobacterium tuberculosis*. Disruption of the *M. tuberculosis lipF* gene reduces survival of *M. tuberculosis* under persistent conditions. Though *in-vivo* analysis of anti sense construct of *lipF* is yet to be done in the next phase of the project, but as of now, *lipF* plays an important role in survival and proliferation under different acidic conditions, thus can be a potential drug target in case of tuberculosis, in both antibiotic formulation as well as for vaccine production.

Appendix

6.1 Bacteriological media

All the media were prepared in Milli RO grade water and autoclaved at 15 pounds per square inch for 15 min. unless otherwise indicated.

6.2 LB Broth (Luria Bertani Broth)

Tryptone	10 g
Yeast extract	5 g
NaCl	10 g

The components were dissolved in 950ml Milli RO water and the pH adjusted to 7.5 with 5N NaOH and the volume adjusted to 1000 ml with Milli RO water. Agar at a concentration of 1.5 % was added whenever solid medium was required.

6.3 Nutrient Broth

Peptone	5 g
Yeast extract	1.5 g
Beef extract	1.5 g
NaCl	5 g

The components were dissolved in 950ml Milli RO water and the pH adjusted to 7.5 with 5N NaOH and the volume adjusted to 1000 ml with Milli RO water.

6.4 Nutrient Agar Tween80 (NAT)

Nutrient Broth	13 g
Tween 80	2 ml

The components were dissolved in 950ml Milli RO water and the pH adjusted to 7.5 with 5N NaOH and the volume adjusted to 1000 ml with Milli RO water. Agar at a concentration of 1.5 % was added whenever solid medium was required.

6.5 Middle brook (MB) 7H9 broth

Disodium Phosphate

2.5gm

Monopotassium Phosphate	1.0gm
L-Glutamic Acid	0.5gm
Ammonium Sulfate	0.5gm
Sodium Citrate	0.1gm
Magnesium Sulfate	50.0mg
Ferric Ammonium Citrate	40.0mg
Zinc Sulfate	1.0mg
Copper Sulfate	1.0mg
Pyridoxine	1.0mg
Calcium Chloride	0.5mg
Biotin	0.5mg
Glycerol	2.0ml
Bovine Albumin	5.0gm
Dextrose	2.0gm
Beef Catalase	3.0mg
Tween80 (0.15%-0.18%)	1.5ml
Glycerol (0.5%)	5ml
Milli RO water	900ml

All the components were mixed using Milli RO water and the pH should be around 6.8. The final volume adjusted to 1000 ml and then sterilised

The components were dissolved in 950ml Milli RO water and the pH adjusted to 7.2 with 5N NaOH and the volume adjusted to 1000 ml with Milli RO water.

6.6 Reagents for Acid Fast Staining

1. Carbol fuchsin (Primary stain)

Basic fuchsin	3g
Phenol	5%
Ethanol (96%)	10 ml

Mixed 10 ml of Basic fuchsin to 90 ml of phenol and the solution was filtered through Whatman filter paper no. 1.

2. Acid alcohol (Decolorizer)

HCL (conc.) 3 ml

Ethanol (96%) 97 ml

3. Malachite green solution (Counter stain)

Malachite green 0.25 g in Milli RO water

6.7 Antibiotics and Substrates

All antibiotic solutions were filter sterilized by a 0.22 μ m filter (Millipore) and stock solutions were stored at -20^oc for long-term use.

Reagent	Stock solution	Final Conc.	Final Conc.
		(in E. coli)	(in Mycobacterium)
Ampicillin	5 mg/ml in H ₂ O	50 µg/ml	-
Kanamycin	5 mg/ml in H ₂ O	50 µg/ml	25 μg/ml
Cycloheximide	5 mg/ml in H ₂ O	100 µg/ml	100 μg/ml
Streptomycin	5 mg/ml in H ₂ O	50 μg/ml	25 μg/ml
X-gal	40 mg/ml in DMF	40 µg/ml	40 μg/ml

6.8 Reagents and Buffers

All the reagents and buffers for DNA and protein work were prepare in Milli Q grade water and sterilized by autoclaving for 15 minutes at 15-psi pressure unless otherwise indicated

6.8.1 Commonly used Buffers

i. Tris HCL buffer

Tris-HCL buffer of desired strength was prepared by dissolving appropriate amount of Tris in distilled water and adjusting the pH with concentrated HCl. For bacteriological work 10 mM Tris-HCl (pH 8.0) was used.

ii. Ethylene diamine tetra acetic acid (EDTA)

0.5 M solution of disodium salt of EDTA was prepared in Milli Q, pH adjusted to 8.0 with NaOH pellets and stored at 4^{0} c.

iii.	Norr	nal Saline		
	NaC	l	8.50 g	
	Milli	RO water	1000 ml (final volume)	
iv.		en Normal Saline		
	0.02% Tween -80 was added to normal saline.			
6.8.2	Reage	nts for Genomic DNA	isolation from Mycobacterium	
	i. TE Buffer			
		Tris-HCl (pH 8.0)	10 mM	
		EDTA	1 mM	
	ii. Tris EDTA Saline (TES) Buffer		TES) Ruffer	
		TRIS-HCL (pH 8.0)	10 mM	
		EDTA	1mM	
		NaCl	150 mM	
	iii. Lysozyme			
		Lysozyme	50 mg/ml in Milli Q	
	iv. Proteinase K			
	1	Proteinase k	20 mg/ml in Milli Q	
	v. Buffer Phenol			
	Molten phenol containing 0.1% 8-hydoxyquinoline was equilibrated w			
	1MTris-HCl (pH 8.0) and twice with 0.1 Tris-HCl (pH 8.0) till the pH > 7.8			
	and then it is stored submerged in 10 mM Tris-HCl (pH 8.0) in dark b			
	4^{0} c away from direct light.			
	vi. Chloroform: Isoamyl alcoholSolution contains 24 parts chloroform and 1 part Isoamyl alcohol. The solution		yl alcohol	
			-	
		is stored in dark bottle at 4° c.		

6.8.3 Buffer for Plasmid Isolation from E. coli

i.	Glucose Tris EDTA Buffer (G	TE)
	TRIS-HCL (pH 8.0)	25 mM
	EDTA (pH 8.0)	10 mM
	Glucose	50 mM
ii.	NaOH-SDS Mix	
	NaOH	0.2 N
	SDS	1.0%

iii. Acetate Mix

Solution contains 3 volumes of 3 M sodium acetate and 4 volume of 7.5 M ammonium acetate.

6.8.4 Buffers for Electrophoresis

i. TAE Buffer (50 X)

Tris Base	242 g
Glacial Acetic Acid	57.1 ml
0.5 M EDTA (pH 8.0)	100 ml
Final Volume	1000 ml

ii. TBE Buffer (50 X)

Tris Base	54 g
Boric acid	27.5 g
0.5 M EDTA	20 ml
Final Volume	1000 ml

iii. Tris-glycine

Tris Base	3.0 g
Glycine	14.4 g

SDS	2.0 g
Final Volume	1000 ml

6.8.5 Buffer for Transformation

i.

MOPS Buffer (pH 6.5)	100 mM
RbCl	10 mM

ii. Transformation Buffer II (TFB II)

Transformation Buffer 1 (TFB I)

MOPS Buffer (pH 6.5)	100 mM
RbCl	10 mM
CaCl ₂	50 mM

6.8.6 Buffer for Gel Loading

i. 6X dye for agarose gel electrophoresis :

Bromophenol Blue	0.25%
Sucrose	40%

The volume was then made up to 10 ml with R Nase free water. For loading onto the gel the buffer was added to a final concentration of 1x to the sample, the mixture incubated at 650c for 3-5 min. and chilled on ice. Before loading 11 of EtBr (1mg/ml) was added to the sample.

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