Secretory Phosphatases Deficient Mutant of *Mycobacterium tuberculosis* Imparts Protection at the Primary Site of Infection in Guinea Pigs

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Abstract

**Background:** The failure of *Mycobacterium bovis* Bacille Calmette-Guérin to impart satisfactory protection against adult pulmonary tuberculosis has necessitated the development of more effective TB vaccines. The assumption that the vaccine strain should be antigenically as similar as possible to the disease causing pathogen has led to the evaluation of *M.tuberculosis* mutants as candidate tuberculosis vaccines.

**Methods/Principal Findings:** In this study, we have generated a mutant of *M.tuberculosis* (Mtb Δmms) by disrupting 3 virulence genes encoding a mycobacterial secretory acid phosphatase (sapM) and two phosphotyrosine protein phosphatases (mptpA and mptpB) and have evaluated its protective efficacy in guinea pigs. We observed that Mtb Δmms was highly attenuated in THP-1 macrophages. Moreover, no bacilli were recovered from the lungs and spleens of guinea pigs after 10 weeks of Mtb Δmms inoculation, although, initially, the mutant exhibited some growth in the spleens. Subsequently, when Mtb Δmms was evaluated for its protective efficacy, we observed that similar to BCG vaccination, Mtb Δmms exhibited a significantly reduced CFU in the lungs of guinea pigs when compared with the unvaccinated animals at 4 weeks after challenge. In addition, our observations at 12 weeks post challenge demonstrated that Mtb Δmms exhibited a more sustainable and superior protection in lungs as compared to BCG. However, the mutant failed to control the hematogenous spread as the splenic bacillary load between Mtb Δmms vaccinated and sham immunized animals was not significantly different. The gross pathological observations and histopathological observations corroborated the bacterial findings. Inspite of disruption of phosphatase genes in Mtb Δmms, the lipid profiles of *M.tuberculosis* and Mtb Δmms were identical indicating thereby that the phenotype of the mutant was ascribed to the loss of phosphatase genes and the influence was not related to any alteration in the lipid composition.

**Conclusions/Significance:** This study highlights the importance of *M.tuberculosis* mutants in imparting protection against pulmonary TB.

Introduction

Tuberculosis (TB) continues to intimidate human race unabashedly and remains a major cause of morbidity and mortality throughout the world [1,2]. Every week, more than 150,000 individuals develop TB and ~30,000 human lives are lost globally due to this dreaded disease. The lethal liaison between TB and HIV infections and the emergence of various forms of drug resistant *M.tuberculosis* strains have made the situation even more precarious [3,4]. Although, the current vaccine, *Mycobacterium bovis* Bacille Calmette-Guérin (BCG) does provide protection against childhood TB especially TB meningitis, it is ineffective in providing consistent protection against the disease in adults and older people [5]. Under the best of the circumstances, it has provided 80% protection, which generally has been to the tune of 40-60% on an average.
Therefore, the need to develop a superior TB vaccine than BCG cannot be over-emphasized.

The purpose of an effective live vaccine would be best served if the vaccine strain is antigenically as similar as possible to the disease-causing pathogen in order for it to generate the host immune responses that mimic natural infection [6]. Comparative genomic studies have revealed that BCG, in comparison to M. tuberculosi s, lacks 16 defined regions (RD1-16) comprising of ~150 genes, some of which are known to encode potential antigenic determinants that could increase the immunogenicity of a vaccine [7,8]. This makes the use of attenuated M. tuberculosi s strains rather than BCG, for the generation of appropriate immune responses, an attractive idea [5,9,10]. Several M. tuberculosi s mutants have been evaluated in animal models and have resulted in varying degrees of success in imparting protection against TB when compared with BCG [11–15]. Immunization of mice with the ∆RD1∆spanCD mutant of M. tuberculosi s (an attenuated M. tuberculosi s RD1 knockout and pantothenate auxotroph) resulted in 1-2 log10 CFU lower bacillary loads in the spleens, lungs and liver when compared with the BCG. However, in bull calves, no histopathological differences were observed in the lung and lymph nodes of ∆RD1∆spanCD vaccinees when compared with the unvaccinated controls [14,15]. Similarly, mice vaccinated with ∆secA2 mutant (secA2 deletion mutant of M. tuberculosi s) exhibited significantly lower pulmonary and splenic CFU when compared with the BCG vaccinated group, however, the same vaccine performed as well as BCG in guinea pigs [11]. In contrast to these observations, Martin et al. demonstrated similar level of protection exhibited by SO2 strain (phoP deletion mutant of M. tuberculosi s) in mice, although guinea pigs vaccinated with SO2 exhibited significantly increased survival time when compared with BCG [12]. The variable results shown by the candidate vaccines and the fact that none of the current candidates has successfully made through the clinical trials reinforce the importance of keeping the pipeline full with the new candidates [16].

Among the secretory proteins of M. tuberculosi s, three phosphatases, namely, mycobacterial secretory acid phosphatase (SapM) and two phosphorylating protein phosphatases (MptpA and MptpB) have been shown to contribute to its pathogenicity [17–20]. SapM dephosphorylates phosphatidylinositol 3-phosphate (PI3P), a membrane trafficking regulatory lipid, resulting in the arrest of phagosome maturation [19]. In addition, in a study by Festjens et al., disruption of sapM locus in BCG improved its protective efficacy as a vaccine against TB [21]. The increased efficacy of the vaccine was accredited to the efficient activation and recruitment of dendritic cells to the draining lymph nodes in the absence of SapM, thus allowing successful antigen presentation and activation of the adaptive immunity by dendritic cells [21]. A recent study showed that the fbpA/sapM double mutant of M. tuberculosi s was attenuated for growth and more immunogenic in macrophages as compared to M. tuberculosi s [22].

MptpA has been demonstrated to block phagosome-lysosome fusion by inhibiting V-ATPase trafficking to the mycobacterial phagosome [23–25]. It has been reported that mptpA mutant of M. tuberculosi s was impaired for survival/growth in THP-1 macrophages and phagosomes harboring the mutant strain exhibited increased phagosome-lysosome fusion [23]. It has been previously reported that M. tuberculosi s devoid of MptpB activity was impaired for survival in IFN-γ activated macrophages and in guinea pigs [26]. In another study, it was shown that MptpB inhibits ERK ½, p38 signaling pathways and caspase 3 activity, thus subverting the host immune response to infection [27]. The importance of MptpB in the intracellular survival of M. tuberculosi s was also demonstrated in a study in which specific inhibitors against MptpB were shown to inhibit mycobacterial survival within murine macrophages [17,27].

In this study, by deleting the function of three virulence genes, namely, mptpA (Rv2234), mptpB (Rv0153c) and sapM (Rv3310), we have developed the mutant MtbΔmms and evaluated its protective efficacy in guinea pig model of experimental tuberculosis.

Materials and Methods

Bacterial strains and growth conditions

The bacterial strains and plasmids used in this study are listed in Table 1. M. bovis BCG (Danish strain) was obtained from BCG laboratories, Chennai, India. M. tuberculosi s H37Rv (ATCC No. 25618) used for challenge was procured from Dr. J. S. Tyagi, AIIMS, New Delhi, India. Mycobacterial strains were grown to mid-log phase in MBTH9 medium supplemented with 1X albumin-dextrose-catalase (ADC), 0.5% glycerol and 0.05% Tween 80. PBS stocks were prepared and stored at -80°C till further use. The CFU of stocks was enumerated by plating appropriate dilutions in duplicates on MBTH11 agar supplemented with 1X oleic acid-albumin-dextrose-catalase (OADC) and 0.5% glycerol. E.coli strains, XL-1 Blue (Stratagene) and HB101 (Life Technologies) were used for cloning purposes. Kanamycin and Chloramphenicol were used at 25 μg/ml and 30 μg/ml, respectively. Hygromycin was used at 50 μg/ml for M. tuberculosis and at 150 μg/ml for E.coli.

Generation of MtbΔmms Mutant of M. tuberculosis

To generate MtbΔmms mutant of M. tuberculosis, a portion of mptpA and sapM was deleted in the genome of MtbΔmptpB [26] and replaced with Kanamycin resistance cassette and Chloramphenicol resistance cassette, respectively. For the generation of MtbΔmptpBΔmptpA double gene mutant, primers were designed to amplify (i) 156 bp of 5’ proximal end of mptpA along with 1135 bp of immediate upstream region of mptpA (amplicon I) and (ii) 167 bp of 3’ distal end of mptpA along with 1240 bp of immediate downstream region of mptpA (amplicon II). The amplicons I and II were PCR amplified and cloned into the vector pLITmus-38 (New England Biolabs) to generate the vector pLITΔA (with a deletion of 169 bp from the central region of mptpA ORF). The Kanamycin resistance gene was excised out from pSD5 as an NheI-BstEII fragment, end-repaired and cloned into NdeI digested, end-repaired pLITΔA to generate pLIT3ΔAK. The vector pLIT3ΔAK was pretreated with alkali [28] and 2 μg of DNA was electroporated into MtbΔmptpB electrocompetent cells to generate a double gene mutant, namely, MtbΔmptpBΔmptpA.
Table 1. Bacterial strains, plasmids, cell line and primers used in this study.

| Strains/Plasmids/Cell line/Primers | Description | Reference |
|------------------------------------|-------------|-----------|
| **Strains**                        |             |           |
| E.coli XL-1 Blue                   | endA1 gyrA96 (naiR) thi-1 recA1 relA1 lac glvIV F4 [-Tn10 proA+ lacIQ Δ (lacZ M15) hsdR17 (K- mK+)] | Stratagene, Heidelberg, Germany |
| E.coli HB101                       | F+(gpt-proA) Δ lacI2 lacY1 (mcrC-mmn) rpsL20 (Strr) xyZ5 mtl-1 recA13 | Life Technologies, CA, USA |
| MtbΔmptpB                         | M.tuberculosis Erdman mptpB mutant | [26] |
| MtbΔmptpBΔmptpA                    | M.tuberculosis mptpB and sapM mutant | This Study |
| MtbΔmmtms                         | M.tuberculosis mptpA, mptpB and sapM mutant | This Study |
| M.tuberculosis H37Rv (ATCC No. 25618) | Virulent strain of M.tuberculosis | Dr. J. S. Tyagi, AIIMS, New Delhi, India |
| M.bovis BCG Danish                 | Vaccine strain against tuberculosis | BCG laboratories, Chennai, India |
| **Plasmids**                       |             |           |
| pLitV/A                            | pLitmus vector carrying the amplicons of mptpA with a deletion of 169 bp from the central region of ORF | This study |
| pLitV/AK                           | pLitmus vector with kanamycin resistance gene cassette flanked with mptpA amplicons | This study |
| pYUBΔSapM                          | Cloning vector with Hygromycin resistance gene cassette flanked with sapM amplicons | unpublished data |
| pYUB.CATtrrnΔSapM                  | Cloning vector with Chloramphenicol resistance gene under mycobacterial trm promoter flanked with sapM amplicons | This study |
| pYUB.CATtrrnΔmptpA                 | A derivative of pSD5 containing Chloramphenicol resistance gene under mycobacterial trm promoter | [51] |
| **Cell line**                      |             |           |
| THP-1                              | The Human acute monocytic leukemia cell line | NCCS, Pune, India |
| **Primers**                        |             |           |
| F-mptpA                            | 5′-gtgctgatccgctgtaacctcattc-3′ | This study |
| R-mptpA                            | 5′-caacctgcggaagatagcag-3′ | This study |
| F-sapM                             | 5′-atgcctggaatccgagctg-3′ | This study |
| R-sapM                             | 5′-cagtagccatcgaatcggg-3′ | This study |

For the generation of MtbΔmmtms mutant of M. tuberculosis, sapM was disrupted in MtbΔmptpBΔmptpA. For this, we employed a modified pYUBΔsapM vector generated in our laboratory (Table 1) by replacing hygromycin resistance cassette in pYUBΔsapM with the Chloramphenicol resistance gene expressed under the mycobacterial trm promoter, CATtrrn. Briefly, primers were designed to amplify (i) ~700 bp amplicon comprising of ~200 bp of 5′ proximal end of sapM and ~500 bp of immediate upstream region of sapM (amplicon I) and (ii) ~700 bp amplicon comprising of ~200 bp of 3′ distal end of sapM and ~500 bp of immediate downstream region of sapM (amplicon II). Both amplicons were cloned into pYUB854 flanking the hygromycin resistance cassette to generate pYUBΔsapM. The hygromycin resistance cassette in pYUBΔsapM was replaced with CATtrrn (The CATtrrn was PCR amplified from pVR1, Table 1) resulting in pYUB.CATtrrnΔsapM. A linear Allelic Exchange Substrate (AES), ΔsapM::CATtrrn was then excised out as KpnI/SpeI fragment and electroporated into MtbΔmptpBΔmptpA to generate the mutant which was designated as MtbΔmmtms (mptpA, mptpB and sapM).

**Confirmation of deletion of mptpA and sapM by Southern hybridization**

To confirm the deletion of mptpA in MtbΔmptpBΔmptpA, the genomic DNA was isolated from the parental strain (MtbΔmptpB) and the mutant strain (MtbΔmptpBΔmptpA) followed by the digestion of 2 μg of DNA with PvuII. The deletion of sapM in MtbΔmmtms was confirmed by isolating the genomic DNA from MtbΔmptpBΔmptpA and MtbΔmmtms, followed by digestion of 2 μg of DNA with PstI. DNA was electrophoresed through 1% agarose gel followed by depurination, denaturation and neutralization of DNA within the agarose gel. DNA was then transferred onto positively charged nylon membrane by capillary transfer overnight and immobilized by UV radiation. 200 bp region at 5′ termini of mptpA and sapM were amplified for the generation of probe. The probe labeling, subsequent pre-hybridization, hybridization and detection were performed as described in the DIG High Prime DNA Labeling and Detection Starter Kit II (Roche Applied Science, IN, USA).

**Lipid profile analysis**

**Isolation of mycolic acids.** Mycolic acids were extracted from M. tuberculosis as well as MtbΔmmtms as described previously [29]. Briefly, mycobacterial strains were grown in 50 ml of MB7H9 supplemented with 1X ADC to an A600 of 1.0. The culture was harvested, heat killed (95°C for 1 hr) and then saponified with 6 ml of 20% tetrabutylammonium hydroxide at 100°C, overnight, to hydrolyze the mycolic acids from the cell wall. Free mycolic acids so generated were methylated by adding 1:1 dichloromethane methyleuchlorid and 300 μl of methyl iodide to form mycolic acid methyl esters. Upon phase separation, the lower organic layer was collected, dried and re-suspended in diethyl ether (3 ml). This lipid suspension was centrifuged at 2500 rpm for 2-3 min and the supernatant was collected and dried. The crystals thus formed, were suspended in 900 μl of a mixture of toluene and acetonitrile (2:1). The solution was transferred to a microcentrifuge tube followed by addition of 600 μl of acetonitrile to the suspension. The suspension was then frozen at -20°C overnight. The solution was centrifuged at 12000 rpm at 4°C for 15 min. Finally, the pellet was suspended in 500 μl of diethyl ether and transferred to a small glass tube and evaporated with liquid nitrogen. The equivalent amount of mycolic acids extracted from...
M. tuberculosis as well as MtbΔmms, suspended in diethyl ether were spotted on a thin layer chromatography (TLC) plate (Merck, TLC Aluminium sheets silica gel 60), chromatographed in hexane: ethylacetate (95: 5, v/v) seven times and visualized by staining with 20% sulphuric acid in ethanol followed by charring.

**Extraction of polar and apolar lipids.** Mycobacterial lipids were extracted as described previously [30]. Briefly, 50 ml of mycobacterial cultures, grown in MB7H9 supplemented with 1X ADC, were harvested at an A600nm of 1.0 and heat killed (95°C for 1 hr). Apolar lipids were extracted by adding 2 ml of methanolic solution of 0.3% sodium chloride and 1 ml of petroleum ether (60-80°C) to the cell pellet. The cell suspension was mixed end-over-end for 30 min followed by centrifugation at 2500 rpm for 10 min. The upper layer consisting of apolar lipids was collected in a separate vial and 1 ml of petroleum ether was added to the lower layer, vortexed and mixed end-over-end for 15 min. The cell suspension was again centrifuged to recollect the upper layer. The upper layers comprising of apolar lipids were pooled and dried at 60°C.

Further, the polar lipids were extracted by adding 2.3 ml of chloroform: methanol: 0.3% sodium chloride solution (90: 100: 30, v/v/v) to the bottom layer. The cell suspension was mixed end-over-end for 60 min followed by centrifugation at 2500 rpm for 10 min. Polar lipids, present in the supernatant fraction, were collected and the pellet was further treated twice with 750 μl of chloroform: methanol: 0.3% sodium chloride (50: 100: 40, v/v/v) to obtain all polar lipids. The supernatants from these three extractions were pooled and further extracted with 1.3 ml of chloroform and 1.3 ml of sodium chloride. The lower layer comprising of polar lipids was collected into a fresh glass tube and dried at 60°C. Equivalent amounts of polar and apolar lipids suspended in chloroform: methanol (2: 1, v/v) from both M. tuberculosis and MtbΔmms strains were then spotted on TLC plates and analysed for different lipid fractions by using different solvent system as described in Table S1. TLC plates were developed by dipping in 10% phosphomolybdate or spraying with 2% orcinol in 10% sulphuric acid (for solvent C) followed by charring.

For the detection of trehalose monomycolate (TMM), trehalose dimycolate (TDM) and sulfolipids (SL), 5 μCi of 14C-acetate was added to 10 ml of log phase culture of both M. tuberculosis and MtbΔmms strains, separately. Cultures were then harvested after 18 hrs of radioactive pulse and apolar lipids were extracted from a methanolic solution of 0.3% sodium chloride and petroleum ether as described above. The organic phase was suspended in chloroform: methanol (2: 1, v/v). Approximately, 25,000 counts from the samples belonging to each strain were spotted on the TLC plate followed by chromatography in the appropriate solvents (Table S1). The lipids were visualized with a Typhoon FLA 700 Phosphorimager.

**Comparison of the growth of MtbΔmms and the parental strain in human macrophages**

Human monocytic THP-1 cells were cultured in complete RPMI-1640 medium [containing 10% heat inactivated FBS and 1% antibiotic-antimycotic mix] (GIBCO Grand Island, NY, USA) and were differentiated to macrophages by the addition of 30 nM Phorbol 12-myristate 13-acetate (PMA, Sigma) for 16 hrs at 37°C, 5% CO2. Cells were washed with complete RPMI medium and rested for 2 hrs in fresh medium without antibiotic-antimycotic mix before infection. For infection, 5 x 10^6 macrophages were infected with 5 x 10^5 mycobacteria to achieve an MOI of 1:1 in 24 well plates for 4 hrs in triplicates [31]. Following infection, the extracellular bacteria were removed by overlaying the cells with RPMI medium containing 200 μg/ml amikacin for 2 hrs. At designated time points, day 0 (4 hrs), 2, 4 and 6, macrophages were lysed by the addition of 0.025% SDS and intracellular bacteria were enumerated by plating appropriate dilutions on MB7H11 agar. Colonies were counted after 4 weeks of incubation at 37°C and the data was expressed as CFU/ml.

**Experimental animals**

Pathogen-free outbred female guinea pigs (200-300 g) of the Duncan-Hartley strain were procured from Disease Free Small Animal House Facility, Lala Lajpat Rai University, Hissar, India. The animals were housed in individually ventilated cages and were provided with food and water ad libitum in a BSLIII facility at University of Delhi South Campus (UDSC), New Delhi, India.

**Ethics statement**

Guinea pig experiments included in this manuscript were reviewed and approved by the Institutional Animal Ethics Committee of University of Delhi South Campus, New Delhi, India (Ref. No. IAEC/AKTBiochem/UDSC/24.08.2010). All animals were routinely cared for, according to the guidelines of CPCSEA (Committee for the Purpose of Control and Supervision of Experiments on Animals), India. Guinea pigs were vaccinated intradermally with mycobacterial strains by injecting not more than 100 μl and were euthanized, whenever required, by CO2 asphyxiation and all efforts were made to ameliorate animal suffering.

**Influence of deletion of phosphatase genes on the pathogenicity of M. tuberculosis**

To evaluate whether the MtbΔmms mutant was sufficiently attenuated for its use as a vaccine, animals (n=6) were inoculated intradermally (i.d.) with 5 x 10^6 bacilli of either M. tuberculosis or MtbΔmms or BCG in 100 μl saline. Animals were euthanized at 4 weeks and 10 weeks post inoculation by CO2 asphyxiation. Lungs, liver and spleen were scored for gross pathological damage such as tissue involvement, areas of inflammation, extent of necrosis and number/size of tubercles due to infection. The scores given to these organs were graded from 1-4 and were based on the modified Mitchison scoring system [32]. For histopathological evaluation, the right lung and a portion of left dorsal lobe of liver were removed and fixed in 10% buffered formalin. 5 μm thick sections of formalin fixed, paraffin embedded lung tissues were stained with haemotoxylin and eosin (H & E). The tissues were coded and the coded samples were evaluated by a certified pathologist having no knowledge of the experimental groups. Left caudal lung lobe and caudal portion of spleen were aseptically removed for the measurement of the bacillary...
load. The specific portions of lungs and spleen were weighed and homogenized separately in 5 ml saline by using a polytron homogenizer. Appropriate dilutions of the homogenates were plated on to MB7H11 agar plates in duplicates and incubated at 37°C for 3-4 weeks. The number of colonies was counted and expressed as mean log_{10} CFU/organ.

**Evaluation of protective efficacy of MtbΔmms against M.tuberculosis infection**

Guinea pigs were divided into 3 groups (n=8) and the animals were immunized intradermally with 5 x 10^5 CFU of either (i) BCG or (ii) MtbΔmms in 100 μl of saline. In the control group, guinea pigs were injected with 100 μl of saline. Twelve weeks post immunization, guinea pigs were infected with a low dose of virulent M.tuberculosis via the respiratory route in an aerosol chamber (Inhalation Exposure System, Glascol Inc.), pre calibrated to deliver 10-30 bacilli in lungs per animal. Guinea pigs were euthanized at 4 weeks and 12 weeks after challenge and evaluated for bacterial load, gross pathological and histopathological changes in various organs as described in the previous section. A significant reduction in these parameters in vaccinated animals was considered as a protective effect of the vaccine.

**Statistical analyses**

For comparison between the groups, Non-parametric Kruskal–Wallis test followed by Mann-Whitney U-test, One-way analysis of variables (ANOVA) with Tukey post-test, Two-way ANOVA with Bonferroni multiple comparison test and student’s t-test were employed, wherever appropriate. Differences were considered significant when p<0.05. For statistical analyses and generation of graphs, Prism 5 software (Version 5.01; GraphPad Software Inc., CA, USA) was used.

**Results**

**Functional disruption of mptpA and sapM in MtbΔmptpB and characterization of the multigene mutant**

To generate triple gene mutant of M.tuberculosis, we first disrupted mptpA in MtbΔmptpB (published from our laboratory previously, [26]) to generate MtbΔmptpBΔmptpA (Figure 1A). Deletion of mptpA was confirmed by three approaches (1). PCR by using mptpA gene specific primers (Table 1, Figure 1B). In the case of MtbΔmptpB, a 0.5 kb amplicon representing the complete mptpA gene was amplified as expected, while in the case of MtbΔmptpBΔmptpA, an amplicon of 2.0 kb was observed indicating the disruption of mptpA by Kanamycin resistance cassette (2). Southern hybridization. In the case of MtbΔmptpB strain, the probe hybridized to a 0.5 kb (lane 1) PvuII fragment whereas disruption of mptpA gene by Kanamycin resistance cassette resulted in a signal at 2.4 kb (lane 2) in the MtbΔmptpBΔmptpA strain (Figure 1C) (3). Nucleotide sequencing. Both 0.5 kb and 2.0 kb amplification products were DNA sequenced that further confirmed the disruption of mptpA in MtbΔmptpBΔmptpA. Further, the sapM gene was deleted in MtbΔmptpBΔmptpA by employing linear AES to generate MtbΔmms (Figure 1D). The triple gene mutant was confirmed by PCR by employing sapM gene specific primers (Table 1, Figure 1E). The primers yielded an amplicon of 0.9 kb in MtbΔmptpBΔmptpA, however, the deletion of sapM gene resulted in a PCR amplicon of 1.5 kb in MtbΔmms (Figure 1E) (2). Southern hybridization. The probe in the MtbΔmptpBΔmptpA hybridized to a 3.0 kb (lane 1) PvuII fragment whereas disruption of sapM gene by Chloramphenicol resistance cassette resulted in a signal at 1.5 kb (lane 2) in the MtbΔmms strain (Figure 1F) (3). Nucleotide sequencing. Both 0.9 kb and 1.5 kb amplification products were DNA sequenced that further confirmed the disruption of sapM in MtbΔmms. Deletion of mptpA and sapM was further confirmed by immunoblot analysis by using polyclonal antibodies raised against MptpA and SapM. As shown in Figure 1G and Figure 1H, we did not observe any expression of MptpA and SapM in MtbΔmms.

**Disruption of Phosphatases Does Not Alter the Lipid Profile of MtbΔmms**

To ascertain whether the disruption of phosphatase genes had any influence on the lipid composition of the mutant, we performed a total lipid analysis of the parental as well as the mutant strain by TLC. M.tuberculosis and MtbΔmms were analysed for the well known characteristic lipids of the tubercle bacillus. The apolar and polar lipid fractions were extracted and assayed for phthiocerol dimycocerosate (PDIM), triacylglycerol (TAG), mycolic acids, free fatty acids, diacylglycerol (DAG), diacyltrehalose (DAT), trehalose monomycolate (TMM), trehalose dimycolate (TDM), glucose monomycolate (GMM), sulfolipids (SL), phosphatidylinositol (PI) and phosphatidylinositol mannoside (PIMs) and phospholipids (P) by TLC. Equivalent amounts of apolar as well as polar lipids from both M.tuberculosis and MtbΔmms were spotted on TLC plates and analysed for different lipid fractions (Table S1).

TLC analysis of the lipids of M.tuberculosis and MtbΔmms exhibited a similar and usual lipid profile with respect to the mycobacterial lipid components. M.tuberculosis produces three classes of mycolic acids: alpha-, keto- and methoxy- mycolic acids [33]. Analysis of total mycolic acids extracted from both M.tuberculosis and MtbΔmms by single dimension TLC exhibited that there was no significant difference in total or alternate types of mycolic acids (Figure 2A). In addition, we observed similar accumulation of structural variants of DIM and TAGs as described by Giovannini et al [34] (Figure 2B). Two dimensional TLC indicated the equivalent presence of both apolar (DAG, TMM, TDM, SL, GMM, DAT) and polar lipids (PIMs, PI and P) in both M.tuberculosis and MtbΔmms as described previously by Bhatt et al [35] (Figure 2C, Figure 2D and Figure 2E). Hence, our observation demonstrated that the lipid profile of M.tuberculosis and MtbΔmms was similar with no notable differences.

**MtbΔmms exhibits a severe growth defect in human THP-1 macrophages**

Next, we compared the growth characteristics of MtbΔmms and the parental strain in MB7H9 medium and in THP-1 cells. As shown in Figure 3A, we did not observe any difference in
Figure 1. Construction and characterization of MtbΔmms. (A) Disruption of mptpA in MtbΔmptpB by homologous recombination. The figure depicts the deletion of mptpA by mptpA::kan AES to generate MtbΔmptpBΔmptpA. Solid bar (a) depicts the region of probe binding for Southern hybridization. kan- Kanamycin resistance cassette. (B) Confirmation of disruption of mptpA in MtbΔmptpBΔmptpA was carried out by PCR by employing mptpA gene specific primers. A 0.5 kb amplification product was obtained with MtbΔmptpB DNA as template (lane 2) and 2.0 kb amplification product was obtained with MtbΔmptpBΔmptpA DNA as template (lane 3). 100 bp ladder (lane 1) and λHindIII ladder (lane 4). (C) Confirmation of mptpA deletion in MtbΔmptpBΔmptpA by Southern hybridization. 200 bp at 5′ termini of mptpA was used as probe (a). Genomic DNA of MtbΔmptpB as well as MtbΔmptpBΔmptpA was digested with PvuII, separated on a 1.0% agarose gel, transferred to nylon membrane and probed with DIG-labeled DNA. The MtbΔmptpB strain (lane 1) showed a hybridization signal at 0.5 kb as expected. The presence of 2.4 kb band in the case of MtbΔmptpBΔmptpA (lane 2) confirmed that allelic exchange had occurred at the mptpA locus. (D) Disruption of sapM in MtbΔmptpBΔmptpA by homologous recombination. The figure depicts the deletion of sapM by sapM::CATtrrn AES to generate MtbΔmms. Solid bar (b) depicts the region of probe binding for Southern hybridization. CATtrrn-Choloramphenicol resistance cassette. (E) Confirmation of disruption of sapM in MtbΔmms was carried out by PCR by employing sapM gene specific primers. A 0.9 kb amplification product was obtained with MtbΔmptpBΔmptpA DNA as template (lane 2) and 1.5 kb amplification product was obtained with MtbΔmms mutant DNA as template (lane 3). 100 bp ladder (lane 1) and λHindIII ladder (lane 4). (F) Confirmation of sapM deletion in MtbΔmms by Southern hybridization. 200 bp at 5′ termini of sapM was used as probe (b). Genomic DNA of MtbΔmptpBΔmptpA as well as MtbΔmms strain was digested with PstI, separated on a 1.0% agarose gel, transferred to nylon membrane and probed with DIG-labeled DNA. The MtbΔmptpBΔmptpA strain (lane 1) showed a hybridization signal at 3.0 kb as expected. The presence of 1.5 kb band in the case of MtbΔmms strain (lane 2) confirmed that allelic exchange had occurred at the sapM locus. (G) Confirmation of mptpA deletion in MtbΔmms by immunoblot analysis. 50 μg of culture filtrate proteins of MtbΔmptpB (lane 1) and MtbΔmms (lane 2) were loaded onto a 12.5% polyacrylamide gel and subjected to electrophoresis. MptpA was detected by immunoblot analysis by using anti-MptpA polyclonal antiserum. MptpA protein migrated as an 18 kDa protein band (lane 1). Disruption of mptpA in MtbΔmms was confirmed by the absence of this 18 kDa band (lane 2). (H) Confirmation of sapM deletion in MtbΔmms by immunoblot analysis. 50 μg of culture filtrate proteins of MtbΔmptpB (lane 1) and MtbΔmms (lane 2) were loaded onto a 12.5% polyacrylamide gel and subjected to electrophoresis. SapM was detected by immunoblot analysis by using anti-SapM polyclonal antiserum. SapM protein migrated as a 28 kDa protein band (lane 1). Disruption of sapM in MtbΔmms was confirmed by the absence of this 28 kDa band (lane 2). doi: 10.1371/journal.pone.0077930.g001
the growth characteristics of Mtb Δmms and the parental strain in MB7H9, however, a significant difference was observed in the growth kinetics between these two strains in THP-1 macrophages. We observed that Mtb Δmms displayed a significantly reduced ability (~2.89 fold difference) to infect macrophages in comparison to the parental strain. Moreover, while M. tuberculosis continued to grow normally for 6 days, MtbΔmms exhibited no sign of growth during this time period demonstrating that the deletion of 3 phosphatases rendered the mutant completely incapable of growing in the macrophages (***p<0.001) (Figure 3B). These results demonstrate the importance of mptpA, mptpB and sapM in the growth and survival of M.tuberculosis in the human macrophages.

Deletion of phosphatase genes leads to the attenuation of M. tuberculosis

To evaluate whether the deletion of three phosphatases had rendered the MtbΔmms mutant sufficiently attenuated for its use as a vaccine, animals were inoculated with either M. tuberculosis or BCG or MtbΔmms strain (Figure 4A). At 4 weeks post inoculation, we observed the maximum bacillary load of 3.96 log_{10} CFU in the lungs of M. tuberculosis infected animals, as compared to negligible bacillary load of 0.28 log_{10} CFU in the lungs of BCG treated animals. No bacilli were detectable in the lungs of Mtb Δmms inoculated animals (Figure 4B). However, when the splenic bacillar counts were analyzed, we observed 5.68 log_{10} CFU, 0.36 log_{10} CFU and 3.99 log_{10} CFU in the animals inoculated with M. tuberculosis, BCG and MtbΔmms, respectively (Figure 4C). Hence, during this initial phase, Mtb Δmms showed some growth in the spleens of animals, although it was ~70 fold less as compared to the parental strain. At 10 weeks post inoculation, a bacillary...
load of 4.69 log₁₀ CFU was observed in the lungs of *M. tuberculosis* infected animals. However, at this time point, no bacilli were recovered from the lungs of the animals inoculated with either MtbΔmms or BCG (Figure 4D). In spleens, we observed a bacillary count of 5.45 log₁₀ CFU in *M. tuberculosis* infected animals. However, no bacilli were recovered from the spleens of animals inoculated with either MtbΔmms or BCG (Figure 4E).

Although, MtbΔmms exhibited some growth in the spleens of the inoculated animals, the bacilli were recovered only during the initial phase (4 weeks post inoculation) and the bacillary load was only 1.4% of that observed in the case of *M. tuberculosis* infected animals (70 fold fewer bacilli in MtbΔmms inoculated animals). Further, on extending the time post inoculation, no MtbΔmms bacilli were recovered in spleens as well as in lungs. Thus, based on these observations, it appeared that as a result of deletion of the phosphatase genes, MtbΔmms was sufficiently attenuated for growth in the host tissues and could be safely used as a vaccine candidate.

Deletion of phosphatase genes renders *M. tuberculosis* incapable of causing pathology in guinea pigs at 10 weeks post inoculation

The gross pathological changes observed in the organs of the animals at 4 weeks and 10 weeks post inoculation with MtbΔmms were commensurate with the bacillary load observed. At 4 weeks post inoculation, the extent of damage observed in the case of *M. tuberculosis* inoculated animals was found to be maximum amongst all the groups with numerous small sized tubercles along with scattered areas of necrosis in all the organs (score: 2 in lungs and liver and 3 in spleen), indicating progressive pulmonary and extra-pulmonary disease (Figure 5A). However, in the case of BCG inoculation, no pathology was observed in the lungs, liver or spleen as expected (score: 1 in all the organs). In the case of inoculation with MtbΔmms, most of the animals displayed negligible lung and hepatic pathology (score: 1) with predominantly scanty and extremely small necrotic lesions. However, in the case of spleen, MtbΔmms inoculated animals were allotted intermediate score (score: 2) in comparison to other two groups. This indicated that MtbΔmms inoculation resulted in some pathological damage to spleens, although, the damage was considerably less in comparison to the *M. tuberculosis* infected animals (Figure 5A). When the animals were evaluated at 10 weeks post inoculation, in the case of *M. tuberculosis* infected animals, as expected, the extent of damage was more than that observed at 4 weeks (score: 3 in lungs, 4 in liver and 3-4 in spleen) with extensive involvement and numerous large sized tubercles effacing the entire organs (Figure 6A). However, the animals inoculated with either BCG or MtbΔmms displayed normal lungs, liver and spleen phenotype (score: 1) with no pathological damage (Figure 6A).

To evaluate the histopathological changes in the lungs and liver of guinea pigs inoculated with *M. tuberculosis*, BCG or MtbΔmms, the tissue sections were stained with haematoxylin and eosin. At 4 weeks post inoculation, the lungs of *M. tuberculosis* infected animals exhibited granulomatous infiltration with caseating necrotic granulomas effacing the pulmonary parenchyma (Figure 5B). Inoculation with either BCG or MtbΔmms resulted in a negligible granulomatous

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**Figure 3. Influence of deletion of phosphatase genes on the growth and survival of *M. tuberculosis* in vitro.** (A) Growth of *M. tuberculosis* and MtbΔmms in broth culture. *M. tuberculosis* and MtbΔmms strains were cultured separately in MB7H9 medium at 37°C/200 rpm for 13 days. *M. tuberculosis* and MtbΔmms exhibited similar growth pattern in MB7H9 broth culture. (B) Growth of *M. tuberculosis* and MtbΔmms in human THP-1 macrophages. THP-1 cells were infected with *M. tuberculosis* or MtbΔmms mutant separately at an MOI of 1:1. The number of intracellular viable bacteria was determined for 6 days. The figure represents the CFU data of *M. tuberculosis* and MtbΔmms obtained on MB7H11 plates. MtbΔmms mutant exhibited a significant attenuation in its growth when compared with *M. tuberculosis*, till 6 days post infection. The experiment was repeated thrice with three independent samples each time. **p<0.001 (Two-way ANOVA). The values are represented as the means (±SE).**

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infiltration in lungs when compared with *M. tuberculosis* infected animals. Liver of *M. tuberculosis* infected animals displayed moderate involvement of tissues with few granulomatous lesions. However, BCG or *MtbΔmms* inoculated animals exhibited a normal hepatic tissue organization with clearly visible portal triad (Figure 5B). At 10 weeks post inoculation, no bacterial colony was obtained from the lung and spleen homogenates of BCG or *MtbΔmms* inoculated animals on MB7H11 agar. Each data point represents the Log_{10} CFU value for an individual animal (n=6) and the bar depicts mean (±SE) for each group. **p<0.01 (One way ANOVA).

Figure 4. Bacillary load in the organs of guinea pigs post inoculation with various strains. (A) Experimental protocol for evaluating the attenuation of *MtbΔmms*. (B, C) Bacillary load in the lungs and spleens of guinea pigs inoculated with *M. tuberculosis* (Mtb), BCG or *MtbΔmms* at 4 weeks post inoculation. (D, E) Bacillary load in the lungs and spleens of guinea pigs inoculated with Mtb, BCG or *MtbΔmms* at 10 weeks post inoculation. At 10 weeks post inoculation, no bacterial colony was obtained from the lung and spleen homogenates of BCG or *MtbΔmms* inoculated animals on MB7H11 agar. Each data point represents the Log_{10} CFU value for an individual animal (n=6) and the bar depicts mean (±SE) for each group. **p<0.01 (One way ANOVA).

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Figure 5. Gross pathology and histopathology of guinea pig organs inoculated with various strains at 4 weeks. (A) Representative photographs of lungs, liver and spleen of individual animal (n=6) euthanized at 4 weeks post inoculation with either *M. tuberculosis* (Mtb), BCG or *MtbΔmms*. The graphical representation of gross scores of various organs is also shown alongside. The bar depicts median for each group. *p<0.05; **p<0.01 and ***p<0.001 (Kruskal Wallis test). (B) The representative photomicrographs of the lung and liver tissues stained with Haematoxylin-Eosin staining (H&E). The figure depicts 20x magnification of representative photomicrographs of the organs of animals inoculated with either Mtb, BCG or *MtbΔmms* euthanized at 4 weeks post inoculation. The scale bars depict 500 μm for the lung as well as liver sections. *MtbΔmms* inoculated animals displayed negligible lung and hepatic pathology. However, in the case of spleens, *MtbΔmms* inoculation resulted in some pathological damage, although, the damage was considerably less in comparison to the *M. tuberculosis* infected animals.

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normal hepatic tissue organization with clearly visible hepatic lobules (Figure 6B).

Mutant of *M. tuberculosis* as Vaccine
MtbΔmms vaccination limits \textit{M. tuberculosis} multiplication in the lungs of guinea pigs

As the MtbΔmms mutant appeared to be safe for its use as a vaccine candidate, we next evaluated its protective efficacy against \textit{M. tuberculosis} challenge. For this, guinea pigs were vaccinated with either BCG or MtbΔmms and were infected with 10-30 \textit{M. tuberculosis} bacilli by aerosol route at 12 weeks post vaccination. As a control, one group of guinea pigs was sham immunized. Following 4 and 12 weeks after challenge, animals were euthanized and bacillary load in the lungs and spleens was determined (Figure 7A). At 4 weeks after challenge, the sham immunized animals exhibited 6.30 log_{10} CFU in lungs. The BCG vaccinated animals exhibited a significantly reduced CFU in lungs (4.43 log_{10} CFU) indicating 1.87 log_{10} CFU reduction (\(*p<0.05\)) as compared to the sham-immunized animals (Figure 7B). MtbΔmms vaccinated animals exhibited a bacillary load of only 3.60 log_{10} CFU in lungs indicating that the mutant also significantly reduced the pulmonary load by 2.70 log_{10} CFU (\(*\ast\ast\ast p<0.001\)) in comparison to the sham immunized animals (Figure 7B). Further, the sham immunized animals exhibited a splenic bacillary load of 5.37 log_{10} CFU (Figure 7C), while, the BCG vaccinated animals exhibited a splenic bacillary load of only 1.62 log_{10} CFU. This significant reduction in splenic bacillary load by 3.75 log_{10} CFU (\(*\ast\ast\ast p<0.001\)) demonstrated a tight control of hematogenous spread of bacilli by BCG. The splenic bacillary load in the case of MtbΔmms vaccinated animals (4.73 log_{10} CFU) was 0.64 log_{10} CFU less when compared with the sham immunized animals but the difference was not significant (Figure 7C).

On extending the time period between challenge and euthanasia to 12 weeks, the sham immunized animals exhibited a bacillary load of 6.15 log_{10} CFU in lungs (Figure 7D). Immunization with BCG resulted in 4.57 log_{10} CFU in lungs as compared to the sham immunized animals, however, the difference in the pulmonary bacillary load between BCG and sham immunized animals was statistically not significant and with the extension of time, the ability of BCG to impede bacillary multiplication met with a considerable decline. In contrast, MtbΔmms vaccinated animals exhibited only 3.16 log_{10} CFU in lungs, thus indicating a significantly reduced bacillary load by 2.99 log_{10} CFU (\(*p<0.05\)) in comparison to the sham immunized animals. These observations demonstrated that MtbΔmms exhibited a more sustainable and superior protection as compared to BCG. The splenic bacillary load in the case of sham immunized animals was 4.92 log_{10} CFU (Figure 7E). Although, the splenic bacillary loads in the cases of BCG vaccination and MtbΔmms vaccination were 3.30 log_{10} CFU and 4.21 log_{10} CFU, respectively, these were not significantly different in comparison to the splenic bacillary load observed in the sham immunized animals. Thus, our observations indicated that MtbΔmms was not able to exhibit a significant control on the hematogenous spread at either 4 weeks or 12 weeks after challenge.

\textbf{MtbΔmms vaccination imparts protection from pathological damage in lungs}

At 4 weeks after challenge, the sham immunized animals exhibited severe pathology in lungs characterized by the presence of numerous large and small sized tubercles (score: 4 in lungs). However, hepatic and splenic tissues exhibited moderate involvement (score: 2 in liver and 2-3 in spleen) (Figure 8A). In contrast, BCG vaccinated animals displayed significantly reduced gross lesions in the organs when compared with the unvaccinated animals (score: 2 in lungs and 1 in liver and spleen). In the case of immunization with MtbΔmms, the animals exhibited moderately inflamed lungs

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Figure 6. Gross pathology and histopathology of guinea pig organs inoculated with various strains at 10 weeks. (A) Representative photographs of lungs, liver and spleen of individual animal (n=6) euthanized at 10 weeks post inoculation with either \textit{M. tuberculosis} (Mtb), BCG or MtbΔmms. The graphical representation of gross scores of various organs is also shown alongside. The bar depicts median for each group. \(*\ast\ast\ast p<0.001\) (Kruskal Wallis test). (B) The representative photomicrographs of the lung and liver tissues stained with Haematoxylin-Eosin staining (H&E). The figure depicts 20x magnification of representative photomicrographs of the organs of animals inoculated with either Mtb, BCG or MtbΔmms euthanized at 10 weeks post inoculation. The scale bars depict 500 μm for the lung as well as liver sections. MtbΔmms inoculated animals displayed complete restoration of pulmonary and hepatic parenchyma with negligible pathological damage to the tissues at 10 weeks post inoculation.

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and spleen (score: 2 in both the organs) with minimal hepatic tissue destruction (score: 1) (Figure 8A). On extending the period between challenge and euthanasia to 12 weeks, we observed an overall increase in the gross pathological damage to the organs of sham immunized animals as characterized by extensive involvement of tissue with numerous large tubercles and scattered areas of necrosis in both lungs and liver (score: 3-4). In addition, a marked discoloration of spleen with numerous large and small sized tubercles and occasional attrition of capsular structure was also observed in most of the sham immunized animals (score: 4) (Figure 9A). BCG immunized guinea pigs exhibited moderate involvement of lung and splenic tissues with small sized tubercles effacing the entire tissues (score: 2-3 in lungs and 1-2 in spleen). However, liver of these animals exhibited normal architecture (score: 1). In case of immunization with MtbΔmms, the animals exhibited a lung phenotype similar to BCG immunized animals (score: 2-3). However, in the case of spleen and liver, vaccination with MtbΔmms resulted in an enhanced pathology when compared with the BCG vaccination (score: 2-3 in spleen and 3-4 in liver) (Figure 9A).

Histopathological analyses of lung and liver sections further substantiated the gross pathological observations. At 4 weeks after challenge, sham immunized animals exhibited several discrete necrotic tubercles occupying 30-40% of the lung sections (Figure 8B). Vaccination with BCG or MtbΔmms prevented pulmonary damage as was evident from the presence of only a moderate granulomatous infiltration and well preserved alveolar spaces, when compared with the sham immunized animals. Sham immunized animals displayed inflammation of hepatic tissues with scattered areas of cellular infiltration. BCG vaccinated animals exhibited minimal involvement of hepatic tissues; however, MtbΔmms immunized guinea pigs exhibited moderate involvement with granulomatous infiltration (Figure 8B). Further, at 12 weeks after challenge, as expected, lungs of the sham immunized animals exhibited extensive granulomatous infiltration with multi-focal coalescing granulomas along with prominent central coagulative necrosis (Figure 8B). BCG immunized animals exhibited the presence of scattered areas of inflammation with discrete granulomas along with necrotic centre in lungs. MtbΔmms vaccinated animals exhibited moderate inflammation in lungs similar to BCG vaccinated animals. On comparing the pathological changes in liver (Figure 9B), sham immunized animals exhibited effacement of a large proportion of hepatic parenchyma due to multiple coalescing foci of necrotic granulomas. Immunization with BCG resulted in a significant reduction in hepatic damage with only a negligible granulomatous infiltration. However, in the case of MtbΔmms immunization, the hepatic lobules displayed an extensive granulomatous infiltration (Figure 9B).

**Discussion**

The development and widespread administration of the BCG vaccine since the early 1920s was originally hailed as a major breakthrough with the promise to eradicate the scourge of TB from the world. However, the early promise was not realized and with the growing incidence of TB cases and inconsistent protective efficacy of BCG, it became evident that the BCG vaccine, in its existing form is of limited use in controlling the disease particularly in the elderly [36]. The availability of complete *M. tuberculosis* genome sequence and an increased understanding of the genes involved in *M. tuberculosis* virulence and immune responses has led to a renewed optimism that it should be possible to develop more efficient TB vaccines than the existing BCG [37,38].

In this study, we have developed a multigene mutant of *M. tuberculosis*, having deletions in three genes namely, *mptpA, mptpB* and *sapM* that are involved in host-pathogen interaction and signal transduction. *M. tuberculosis* Erdman and *M. tuberculosis* H37Rv have been commonly used as basis for generating attenuated strains of *M. tuberculosis* [39-44]. However, to ensure that their virulence was not diminished on account of repeated *in vitro* subculturing, the bacilli recovered from the organs of infected animals were subcultured only once.

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**Figure 7. Determination of bacillary load in guinea pig organs after immunization with various strains.**

(A) Experimental protocol for evaluating the protective efficacy of MtbΔmms against *M. tuberculosis* infection. (B, C) The bacillary load in the lungs and spleens of guinea pigs (n=8) vaccinated with BCG, MtbΔmms or saline at 4 weeks after *M. tuberculosis* challenge. (D, E) The bacillary load in the lungs and spleens of guinea pigs (n=8) vaccinated with BCG, MtbΔmms or saline at 12 weeks after challenge. MtbΔmms imparted a significant protection as compared to the sham immunized animals in the lungs of vaccinated animals. Each data point represents the Log_{10} CFU value for an individual animal and the bar depicts mean (±SE) for each group. *p<0.05 and ***p<0.001 (One-way ANOVA).

Figure 9. Histopathological sections of lung and liver from vaccinated and control guinea pigs. (A) Necrosis and scattered areas of inflammation with discrete granulomas along with necrotic centre in lungs. BCG vaccinated animals exhibited minimal involvement of hepatic tissues; however, MtbΔmms immunized guinea pigs exhibited moderate involvement with granulomatous infiltration. BCG immunized animals exhibited the presence of scattered areas of inflammation with discrete granulomas along with necrotic centre in lungs. MtbΔmms vaccinated animals exhibited moderate inflammation in lungs similar to BCG vaccinated animals. On comparing the pathological changes in liver (Figure 9B), sham immunized animals exhibited effacement of a large proportion of hepatic parenchyma due to multiple coalescing foci of necrotic granulomas. Immunization with BCG resulted in a significant reduction in hepatic damage with only a negligible granulomatous infiltration. However, in the case of MtbΔmms immunization, the hepatic lobules displayed an extensive granulomatous infiltration (Figure 9B).
Figure 8. Gross pathology and histopathology of vaccinated guinea pig organs at 4 weeks after challenge. (A) Representative photographs of lungs, liver and spleen of individual animal (n=8) vaccinated with either BCG, MtbΔmms or saline and euthanized at 4 weeks after challenge. The graphical representation of gross scores of various organs is also shown alongside. The bar depicts median for each group. ∗p<0.05; ∗∗p<0.01 and ∗∗∗p<0.001 (Kruskal Wallis test). (B) The representative photomicrographs of the lung and liver tissues stained with Haematoxylin-Eosin staining (H&E). The figure depicts 20x magnification of representative photomicrographs of organs of the animals vaccinated with either BCG, MtbΔmms or saline and euthanized at 4 weeks after challenge. The scale bars depict 500 μm for lung as well as liver sections. Unvaccinated animals show severe pathology in lungs characterized by the presence of numerous large and small sized tubercles. However, hepatic and splenic tissues show moderate involvement. In the case of immunization with MtbΔmms, the animals show moderately inflamed lungs with minimal hepatic tissue destruction similar to that observed in the case of BCG vaccination. However, spleens of MtbΔmms vaccinated animals exhibit pathological damage similar to sham immunized animals.

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Figure 9. Gross pathology and histopathology of vaccinated guinea pig organs at 12 weeks after challenge. (A) Representative photographs of lungs, liver and spleen of individual animal (n=8) vaccinated with either BCG, MtbΔmms or saline and euthanized at 12 weeks after challenge. The graphical representation of gross scores of various organs is also shown alongside. The bar depicts median for each group. ∗p<0.05 and ∗∗p<0.01 (Kruskal Wallis test). (B) The representative photomicrographs of the lung and liver tissues stained with Haematoxylin-Eosin staining (H&E). The figure depicts 20x magnification of representative photomicrographs of organs of the animals vaccinated with either BCG, MtbΔmms or saline and euthanized at 12 weeks after challenge. The scale bars depict 500 μm for lung as well as liver sections. Unvaccinated animals show characteristic presence of multiple coalescing granulomas with necrosis in both lungs and liver. In addition, the animals show a marked discoloration of spleen with numerous large and small sized tubercles. MtbΔmms immunized animals, on the other hand show reduced granulomatous infiltration with only a few small and discrete granulomas in lungs. However, liver and spleen of MtbΔmms vaccinated animals exhibit pathological damage similar to sham immunized animals.

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for their use for generating mutants or for challenging the animals.

We have evaluated the vaccine efficacy of the resulting mutant against *M. tuberculosis* challenge in guinea pigs. To ensure that the mutated genes were essential only for the growth of the pathogen in the host and not in the broth culture, we selected the genes implicated in the host-pathogen interaction. As expected, MtbΔmms grew in broth culture similar to the parental strain, however, it displayed a significantly reduced ability to infect and grow inside the human THP-1 macrophages emphasizing that the phosphatases are vital for the growth of the pathogen in macrophages.

In addition, our studies in guinea pigs also provide evidence that MtbΔmms is highly attenuated for its growth and ability to cause pathology in the host. On inoculation of guinea pigs with MtbΔmms, no bacilli were recovered from the lungs of the animals at any time point of the study. From the spleens of these animals, bacilli were recovered during early phases after the inoculation (4 weeks) but no bacilli were recovered after this initial period. This demonstrated that MtbΔmms could survive only for a short while; therefore, at 10 weeks post inoculation, the organs of MtbΔmms as well as BCG inoculated animals appeared to be similar with no apparent damage indicating that the mutant was safe to be used as a vaccine candidate.

For the evaluation of protective efficacy, we employed guinea pig model of experimental tuberculosis. The guinea pig model of low dose aerogenic infection with virulent *M. tuberculosis* has been preferentially used to elucidate the events in the pathogenesis of pulmonary tuberculosis [45]. Guinea pigs are more susceptible to tuberculosis infection and have the advantage over mice in that the pathology of the disease in this model is closer to human tuberculosis. Thus, it serves as an effective model to evaluate vaccine efficacy. When guinea pigs are infected with less than 10 CFU of virulent *M. tuberculosis*, it has been observed that the pathogen disseminates from lungs to the pulmonary lymph nodes via hematogenous spread and then appears in spleens within ~3 weeks post infection [46,47]. Bacilli reseed the lung by ~4 weeks to form secondary granulomas. The protective efficacy of MtbΔmms was evaluated based on its ability to reduce the bacillary load in lungs and spleens of guinea pigs post *M. tuberculosis* infection as well as to control the pathological damage.

Our study shows that MtbΔmms vaccination was able to restrict bacterial multiplication at the primary site of infection leading to reduction in the pulmonary bacillary load and this bacillary load reduction by MtbΔmms as well as by BCG was comparable during early phase (4 weeks) after infection. Unlike in the case of BCG vaccination, however, MtbΔmms was not able to control hematogenous spread. A number of studies have reported the examples of vaccines which fail to provide consistent protection in all the organs uniformly [12,48,49]. For example, it has been reported that a recombinant BCG expressing ESAT-6 provided significant protection in both mice and guinea pigs against dissemination at extra-pulmonary site but failed to protect against pulmonary form of the disease [49]. On the other hand, vaccination of guinea pigs with DNA encoding the mycobacterial antigen MPB83 influenced the pulmonary pathology but not the hematogenous spread following the aerogenic infection with *Mycobacterium bovis* [48]. Also, in the case of vaccination of guinea pigs with the ΔphoP mutant, a significant reduction in the bacillary load in lungs but not in the spleens was observed as compared to the unvaccinated animals [12].

On extending the period between the *M. tuberculosis* challenge and euthanasia to 12 weeks, although BCG appeared to lose the control of bacillary multiplication in the pulmonary tissue, MtbΔmms was still very effective in controlling the lung infection. At this time point, however, neither BCG nor MtbΔmms exhibited any significant control over the hematogenous spread. The pathological damage in the animals from various groups corroborated the CFU data. From this, we could infer that MtbΔmms imparted as much or better control of the disease than BCG at the pulmonary site. However, immunization with the phosphatase mutant did not show any superior control over the bacillary multiplication in spleens, when compared with the sham immunized animals.

As phosphatases play an important role in the lipid metabolism, we evaluated the lipid profiles of *M. tuberculosis* and MtbΔmms to ascertain whether disruption of phosphatase genes might result in the altered lipid profile in MtbΔmms. However, our observations demonstrated that the lipid profiles of both the strains were identical inspite of disruption of phosphatase genes in MtbΔmms indicating thereby that the phenotype of the mutant was ascribed to the loss of phosphatase genes and the influence was not related to any alteration in the lipid composition.

To summarize, we demonstrate that mutation of genes encoding the signal transduction associated phosphatases of *M. tuberculosis* provides optimism for the generation of novel potential vaccine candidates against tuberculosis. The MtbΔmms was not only significantly attenuated for growth in macrophages and guinea pigs, it also imparted an enhanced protection against pulmonary TB. However, further modifications would be required in order for MtbΔmms to elicit more appropriate immune responses for imparting superior protection including the control of hematogenous spread. Moreover, due to increasing concern about the emergence of antibiotic resistance in human pathogens, use of antibiotic-resistant genes in recombinant vaccines meant for use in humans is not permissible [50]. Hence, the antibiotic resistance genes from MtbΔmms would have to be removed before any possibility of its use in human clinical trials. Our future efforts would focus on addressing these issues.

**Supporting Information**

**Table S1. Solvent systems employed for lipid analyses.**

**DOC**

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