Lysine acetylation of DosR regulates the hypoxia response of *Mycobacterium tuberculosis*

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Abstract

Tuberculosis caused by *Mycobacterium tuberculosis* (*Mtb*) infection remains a large global public health problem. One striking characteristic of *Mtb* is its ability to adapt to hypoxia and trigger the ensuing transition to a dormant state for persistent infection, but how the hypoxia response of *Mtb* is regulated remains largely unknown. Here we performed a quantitative acetylome analysis to compare the acetylation profile of *Mtb* under aerated and hypoxic conditions, and showed that 377 acetylation sites in 269 *Mtb* proteins were significantly changed under hypoxia. In particular, deacetylation of dormancy survival regulator (DosR) at K182 promoted the hypoxia response in *Mtb* and enhanced the transcription of DosR-targeted genes. Mechanistically, recombinant DosRK182R protein demonstrated enhanced DNA-binding activity in comparison with DosRK182Q protein. Moreover, Rv0998 was identified as an acetyltransferase that mediates the acetylation of DosR at K182. Deletion of Rv0998 also promoted the adaptation of *Mtb* to hypoxia and the transcription of DosR-targeted genes. Mice infected with an *Mtb* strain containing acetylation-defective DosRK182R had much lower bacterial counts and less severe histopathological impairments compared with those infected with the wild-type strain. Our findings suggest that hypoxia induces the deacetylation of DosR, which in turn increases its DNA-binding ability to promote the transcription of target genes, allowing *Mtb* to shift to dormancy under hypoxia.

Introduction

In 2015, ~10.4 million new infections and 1.8 million deaths globally were caused by *Mycobacterium tuberculosis* (*Mtb*)1. The remarkable success of *Mtb* as a pathogen is due to its marked phenotypic drug resistance and its ability to evade the host immune system. *Mtb* can also enter into a nonreplicating state and persist in humans for extended periods without causing disease in what is known as a latent tuberculosis (TB) infection (LTBI)2. Clarification of the mechanisms underlying the LTBI state may lead to the development of useful pharmaceuticals to prevent *Mtb* infection or even cure individuals harboring *Mtb* in the dormant stage3. Adaptations to hypoxia have been implicated to play a prominent role in the dormancy of *Mtb*4. Accumulating evidence has demonstrated that oxygen tension correlates tightly with *Mtb* growth, the formation of hard, fibrous, hypoxic granulomas, and, eventually, the development of infection5. Wayne et al.6, 7 have pioneered the development of culture conditions that gradually deprive bacteria of oxygen to generate nonreplicating persistent bacilli in vitro as a model for latency. With this, Schubert et al.8 profiled the proteome of dormant *Mtb* and observed that the stress-induced dormancy survival regulator (DosR) regulon consisted of ~20% of the cellular protein content during dormancy. However, whether posttranslational modifications (PTMs) are involved in this process remains elusive.


N\textsuperscript{-}lysine acetylation is an abundant and evolutionarily conserved PTM found in prokaryotes and eukaryotes\textsuperscript{9}. As a dynamic and reversible process, lysine acetylation affects protein conformation and/or charge, thus altering DNA-binding affinity, enzymatic activity, protein–protein interactions, and protein or mRNA stability\textsuperscript{10, 11}. Increased global analyses of lysine acetylation in bacteria have been reported, including in Escherichia coli\textsuperscript{12, 13}, Erwinia amylovora\textsuperscript{14}, Bacillus subtilis\textsuperscript{15}, Salmonella enterica\textsuperscript{16}, and Mtb\textsuperscript{17, 18}, as well as in others\textsuperscript{19}. Protein acetylation is implicated in nearly all cellular processes, such as central metabolism, protein translation, and pathogen virulence\textsuperscript{20, 21}. Increasing evidence has shown that protein acetylation plays an important regulatory role in mycobacteria\textsuperscript{22, 23}. The universal stress protein (USP) was the first-characterized acetylated protein in mycobacterium\textsuperscript{24}. Recently, Bi et al.\textsuperscript{25} identified the modulation of central carbon metabolism by the acetylation of isocitrate lyase in Mtb. Given the widespread regulatory role of lysine acetylation, it is tempting to speculate that lysine acetylation is involved in the hypoxia adaption of Mtb. Hence, we examined alterations in the protein acetylation profile of Mtb in response to hypoxia by quantitative proteomics and subsequently identified 377 acetylation sites in 269 proteins that were significantly changed. Furthermore, we confirmed the lysine acetylation of DosR and its role in the response of Mtb to hypoxia.

Results

Quantitative proteomic profiling of lysine acetylation in Mtb H37Rv under aeration and hypoxia

To profile the lysine acetylation of Mtb during exponential growth and hypoxia-induced dormancy, we cultured Mtb H37Rv under normal aerobic conditions and gradually decreasing oxygen levels using the Wayne model (Supplementary Figure S1)\textsuperscript{26}. We harvested cultures at day 12 during exponential growth and hypoxia. Acetylated peptides were enriched using a specific acetyl lysine antibody and analyzed by liquid chromatography–tandem mass spectrometry (LC–MS/MS) (Fig. 1a). We identified 1215 acetylation sites in 679 proteins, of which 852 acetylation sites in 497 proteins were quantified (Supplementary Table S1). Relative quantitation was divided into two categories: under hypoxia, 321 acetylation sites in 215 proteins were upregulated; and 56 acetylation sites in 54 proteins were downregulated (Supplementary Table S2). Classification results for biological processes and molecular functions both showed that the largest group of upregulated acetylated proteins under hypoxia consisted of enzymatic proteins associated with metabolism, which accounted for 31% and 56% of the upregulated acetylated proteins, respectively (Fig. 1b, c). These findings are consistent with the previous observation that most lysine-acetylated proteins are involved in metabolism.

Furthermore, we performed Gene Ontology (GO) enrichment analysis on the acetylation data. As revealed previously, hypoxia induces comprehensive changes in the bacterium, with Mtb shifting to exploit lipids as the primary nutrient source\textsuperscript{27}. Consistently, our data demonstrated that upregulated acetylated proteins under hypoxia were enriched for biological processes related to lipid metabolism (Fig. 1d). From the GO enrichment analysis, we also found that upregulated acetylated proteins under hypoxia were significantly enriched for some specific molecular functions, such as flavin adenine dinucleotide binding, oxidoreductase activity, and coenzyme binding (Fig. 1d). For the 54 acetylated proteins downregulated under hypoxia, biological processes associated with the growth of a symbiont involved in its interaction with a host were found to be enriched (data not shown). Additionally, proteins with ligase activity that form carbon–nitrogen bonds also had a higher tendency to be acetylated under normal aerobic conditions. For the Kyoto Encyclopedia of Genes and Genomes (KEGG) metabolic pathways analysis, upregulated acetylated proteins under hypoxia were highly enriched for carbon metabolism, RNA degradation, and propionate metabolism, among others (Fig. 1e), while acetylated proteins involved in the ribosome were highly enriched under aerobic conditions. Overall, these data indicate that lysine acetylation might play a comprehensive regulatory role in the adaption of Mtb to hypoxia.

Hypoxia induces deacetylation of DosR

Numerous studies have demonstrated the important role of DosR during the physiological adaptation of Mtb to hypoxia\textsuperscript{28, 29}. Three amino acids in DosR, namely, Lys179, Lys182, and Asn183, directly bind to nucleotide bases in the DNA motif and induce the expression of target genes\textsuperscript{30}. Mutation of Lys182 directly reduces the DNA-binding affinity of DosR and abrogates the induction of its regulon genes, thus highlighting the important role of K182 in DosR\textsuperscript{31}. However, the relevant mechanism is still unclear. From our quantitative acetylome profiling data, DosR was shown to be acetylated at K182 (Fig. 1f). Considering the location and positive charge of this lysine residue, we hypothesized that acetylation of K182 may be involved in regulating DosR activity, specifically the DNA-binding ability of DosR. Sequence alignment analysis demonstrated that K182, the acetylated lysine residue, is evolutionarily conserved among different mycobacteria species (Fig. 1g). Interestingly, quantitative proteomics analysis revealed that the acetylation of DosR K182 was reduced under hypoxia. To further confirm whether the acetylation levels of DosR K182 respond to hypoxic stress, we produced anti-DosR- and anti-DosR Ac-K182-specific
polyclonal antibodies (characterized as shown in Supplementary Figure S2) and applied these to examine the acetylation levels of DosR K182 in *Mtb* H₃₇₇Rv cultured under normal aerobic or hypoxic conditions. Consistently, a significant reduction of acetylation at K182 of DosR under hypoxia was observed (Fig. 1h), indicating that...
DosR may undergo deacetylation at K182 in response to hypoxia.

**Deacetylation of DosR promotes the adaption of *Mtb* to hypoxia**

Based on the quantitative acetylome data generated in this study, six of the key regulated proteins from the hypoxia regulatory network were acetylated at various levels in response to hypoxic stress, namely, DosR, PhoP, SigA, SigB, SigF, and Lsr2 (Table 1). To examine whether acetylation affects the bacterium’s adaption to hypoxia, we generated recombinant *Mycobacterium smegmatis* expressing corresponding wild-type (WT) genes, as well as counterparts with a lysine-to-arginine substitution of K182 in DosR. The effect of deacetylation on the adaption to hypoxia by mycobacteria species. The asterisk denotes the conserved lysine residue, and this result was analyzed by DNAStar software. Acetylation of DosR promoted the adaption of *Mtb* to hypoxia.

**Hypoxia induces deacetylation of DosR.** a Workflow of quantitative proteomics analysis of lysine acetylation in *H*/*Rv* under hypoxia. *Mtb* H37Rv was cultured under aeration or hypoxia in conical screw-capped Nephelo flasks with 20 mm side arms according to the Wayne model. Samples for aeration (Aera) and hypoxia (Hypo) cultures were harvested after 12 days. Proteins were extracted and digested by trypsin. After separate labeling with a tandem mass tag (TMT) 6-plex kit, peptides from Aera or Hypo were mixed 1:1, enriched with anti-acetyl lysine antibody beads and loaded for LC–MS/MS analysis. The resulting MS/MS data were processed for bioinformatics analysis. b–c Gene ontology functional classification of biological processes (b) and molecular functions (c) of 215 upregulated acetylated proteins under hypoxia. d Gene ontology enrichment analysis of biological processes and molecular functions of 215 upregulated acetylated proteins under hypoxia. The differently colored bars indicate the corrected *p* values for the enrichment of the annotations. e KEGG pathway enrichment analysis of 215 upregulated acetylated proteins under hypoxia. f Mass spectroscopy (MS) analysis identified K182 acetylation of DosR. g K182 in DosR is conserved among various mycobacteria species. The asterisk denotes the conserved lysine residue, and this result was analyzed by DNAStar software. h Acetylation of DosR in *H*/*Rv* WCL under hypoxia was downregulated. *H*/*Rv* were grown under aeration or hypoxia in 7H9 broth supplemented with 10% ADC for 3 days. DosR proteins were immunoprecipitated (IP) by the anti-DosR antibody. The acetylation level of DosR was determined by immunoblotting (IB) using an anti-DosR Ac-K182-specific polyclonal antibody. Western blots are representative of at least three independent experiments.

DosR controls the dormancy survival regulon that consists of ~50 genes. Since deacetylation of DosR K182 directly promotes *Mtb* adaptation to hypoxia, we hypothesized that acetylation of DosR K182 regulated the transcription of dormancy survival-related genes. Several known DosR-regulated genes, including *acr*, *dosR*, *fdxA*, *Rv1738*, *Rv2030*, *Rv2623*, *Rv2626c*, and *Rv3130c*, were selected to evaluate whether K182 acetylation affects the activity of DosR as a transcription factor. A DosR deletion mutant of *Mtb*, WT, and a deacetylation complementation mutant were cultured under hypoxic conditions as described previously. As expected, the transcription levels of eight candidate genes were all significantly upregulated for the K182R mutant under hypoxia compared with normal aerobic conditions and the WT strain (Fig. 2e–l). Furthermore, we wanted to determine whether acetylation of K182 directly hindered the DNA-binding activity of DosR. We constructed expression plasmids of WT DosR and its derivatives, K182R and K182Q (replaced with glutamine to mimic the constitutively acetylated form), using a 6His tag sequence at their C terminals.

**Rv0998 acetylates DosR and negatively regulates the adaption of *Mtb* to hypoxia**

Lysine acetylation is reversible and is performed in bacteria by members of the GCN5-related N-acetyltransferase (GNAT) superfamily, as well as by NAD+-dependent deacetylases. In *Mtb*, 20 GNATs have been predicted. The cAMP-dependent protein lysine acetyltransferase Rv0998 was recently shown to catalyze the acetylation of many important proteins. To examine whether Rv0998 catalyzes
the acetylation of DosR K182, we constructed an Rv0998 deletion mutant of Mtb H37Rv (characterized as shown in Supplementary Figures S4D and S4E). Western blotting revealed that the acetylation level of DosR at K182 in the Rv0998 deletion mutant was significantly reduced (Fig. 3a). To further confirm whether DosR is a substrate of Rv0998, we overexpressed and purified 6× His-tagged DosR proteins from E. coli, as well as Rv0998. DosR was incubated with Rv0998 in the presence of AcCoA and/or cAMP. Western blotting revealed that DosR K182 could be acetylated by Rv0998 in vitro without CAMP, although cAMP further enhanced the function of Rv0998 (Fig. 3b). We also synthesized a short peptide of DosR, from 175 to 188 amino acids in length, termed K182-pep. In vitro acetylation reactions further confirmed that Rv0998 could acetylate DosR K182-pep (Fig. 3c). Next, we used the Rv0998 deletion mutant to confirm the effect of K182 acetylation on growth under hypoxia and on target gene transcription. The Rv0998 deletion mutant was found to significantly promote the Mtb response to hypoxia (Fig. 3d) and to dramatically activate the transcription of eight DosR target genes (Fig. 3e–f).

### Protein acetylation contributes to the pathogenesis of Mtb

Earlier studies revealed that DosR plays a critical regulatory role in the adaptation and survival of Mtb within cells and tissues. We hypothesized that DosR K182 acetylation was critical for bacterial survival and proliferation in macrophages. To answer this question, primary peritoneal macrophages were infected with Mtb H37Rv, the deletion mutant of DosR, and WT or K182R complement mutants with a multiplicity of infection (MOI) of 1. After 24 h, cells were lysed, and the number of intracellular bacteria was calculated by colony-forming unit (CFU) counting. We found that the deletion mutant multiplied more efficiently than Mtb H37Rv, WT, or K182R complement mutants, in agreement with the studies of Parish. The K182R mutant exhibited lower bacterial proliferation than the WT complement mutants (Fig. 4a). We also used the deletion mutant of Rv0998 to detect the intracellular survival of Mtb in peritoneal macrophages. As expected, we observed significantly retarded growth of the Rv0998 deletion mutant, with multiplicities of infection of 1 or 2 at 24 h (Fig. 4b).

To further understand the functional role of DosR K182 acetylation in the pathogenesis of TB, we performed intraperitoneal (i.p.) infections with Mtb in mice. We found that even though the average injected dose for the four groups was similar (Fig. 4c), the numbers of bacteria in the lungs and spleen were both elevated in mice infected with the DosR deletion mutant for 28 days (Fig. 4d, e). The number of DosR complement mutant bacilli in the lungs and spleen was significantly decreased compared with the deletion mutant, while that of the K182R complement mutant was even lower (Fig. 4d, e). From lung histopathological analysis, larger lesions and the pathological involvement of a greater proportion of the lung were observed for mice infected with the DosR deletion mutant relative to mice infected with DosR and K182R complement mutant bacilli, although damage by the K182R mutant was milder (Fig. 4f). In conclusion, the deacetylation of K182 caused significantly lower Mtb bacterial loads as well as attenuated pulmonary inflammation in a mouse model.

### Discussion

As is well known, hypoxia stress limits the in vivo growth of Mtb. However, Mtb can adapt to limited oxygen by entering into a metabolically altered state while awaiting the opportunity to reactivate. Defining the strategies employed by Mtb to shape its adaptation to hypoxia is of intense interest. Many “omics” analyses have been used to comprehensively evaluate the response of Mtb to hypoxia. The utilization of specific antibodies to acetylated peptides in immunoprecipitation has significantly improved the enrichment and identification of acetylated lysine residues with proteomic profiling. Although an analysis of the acetylome of Mtb has recently been reported, the impact of lysine acetylation on the adaption of Mtb to hypoxia has not yet been addressed. Understanding how Mtb modulates lysine acetylation in response to oxygen limitation may lead to new strategies for its control.

We implemented a universally applicable, tandem mass tag (TMT)-labeled approach to estimate variations in lysine acetylation in Mtb during exponential growth and hypoxia-induced dormancy. We identified 1215 acetylation sites on 679 proteins, many more than found by Bi et al. The 852 acetylation sites on 497 proteins were identified under both aerated and hypoxic conditions and were quantified. In addition, 363 residual unquantifiable

Table 1  Acetylation of six regulated proteins from Mtb hypoxia regulatory network

| Protein     | Protein description                  | Position | Hypo-vs-Aera |
|-------------|-------------------------------------|----------|--------------|
| P71814      | OmpR family two-component system response regulator | 197      | 3.47         |
| P9WG13      | RNA polymerase sigma factor SigF     | 121      | 2.45         |
| P9WG15      | RNA polymerase sigma factor SigB     | 308      | 2.21         |
| P9WG11      | RNA polymerase sigma factor SigA     | 334      | 1.57         |
| P9WMM9      | Transcriptional regulatory protein DevR (DosR) | 179      | 1.53         |
| P9WP7       | Nucleoid-associated protein Lsr2      | 43       | 0.53         |

| Protein     | Protein description                  | Position | Hypo-vs-Aera |
|-------------|-------------------------------------|----------|--------------|
| P9WMM9      | Transcriptional regulatory protein DevR (DosR) | 182      | 0.65         |
Fig. 2 (See legend on next page.)
acetylation sites may have also been derived from dramatic variations in this modification under different growth conditions, or they could have been due to the poor replicability of the high-throughput analysers. Among the quantified sites, we found 377 acetylation sites on 269 proteins that were significantly changed. These findings strongly indicate that protein acetylation is important for the regulation of protein function and the subsequent adaption to hypoxia. Unfortunately, the quantification experiment was performed only once (two samples and one biological replicate for each experimental condition), so there was neither a description of reproducibility nor any statistical significance. Although we lacked biological replicates, we produced many functional experiments to verify the reliability of the proteomics data. After screening by proteomics data, we selected DosR as our research object. Taking into account the reliability of the data, we produced an anti-DosR antibody to lysine K182. This antibody was used to detect the acetylation of the target protein in the quantitative experiment (data not shown). From experiments using the deletion mutant and an in vitro acetylation assay (Fig. 3a), we postulated that DosR was a substrate of Rv0998. The residual acetylation of DosR in the Rv0998 deletion mutant indicated that another as yet to be identified acetyltransferase(s) is also involved in the regulation of the acetylation of DosR. The phenotype obtained from the Rv0998 deletion mutant further supported the notion that deacetylation of DosR promoted target gene transcription and had a profound effect on the growth of Mtb under hypoxia (Fig. 3d–l).

Of note, earlier studies revealed a contradictory role for DosR in Mtb virulence. Parish et al. reported that the DosR mutant of Mtb is hypervirulent in both SCID and C57BL/6 mice, while studies in guinea pigs, rabbits, and even macaques revealed that mutants of DosR exhibited reduced infectivity. Here we showed that the deletion mutant of DosR promoted the intracellular survival of Mtb in macrophages, while a K182R mutant exhibited lower proliferation than WT complement
mutants (Fig. 4a) as well as an Rv0998 deletion mutant (Fig. 4b). In infected mice, deacetylation of K182 also caused significantly lower bacterial loads as well as attenuated lung damage (Fig. 4c–f). Although isolated macrophages and a C57Bl/6 mouse infection model may not represent the real hypoxic situation in vivo, these
infection results suggest that deacetylation of K182 can promote the regulatory effect of DosR on Mtb virulence under different conditions. Studies in more effective infection models should be performed to further confirm our findings.

In conclusion, we propose that hypoxia may induce the deacetylation of DosR, which in turn increases its DNA-binding ability to promote the transcription of DosR downstream target genes, allowing Mtb to rapidly adapt to hypoxia (Supplementary Figure S5).

### Materials and methods

#### Bacterial strains, plasmids, primers, and media

All strains, plasmids, and primers used in this study have been described in Supplementary Table S3 and S4. Middlebrook 7H9 broth (Difco/Becton Dickson, Franklin Lakes, NJ) supplemented with 10% ADC (5% bovine serum albumin (BSA), 2% dextrose, and 5% catalase) was used as an enriched medium. Middlebrook 7H10 agar (Difco) plates contained 10% ADC, 1.5% (w/v) agar, and antibiotic supplements as required. The antibiotics used...
were 100 μg/mL ampicillin, 50 μg/mL or 75 μg/mL hygromycin B, and 50 μg/mL kanamycin.

**Bacterial cultures**

Experiments performed by Wayne in in Mtb H37Rv have been described previously. In brief, conical screw-capped Nephelo flasks with 20 mm side arms and flat bases (Wheaton Scientific Products, Millville, NJ) were used to culture bacteria. Mtb H37Rv was grown in 7H9 broth supplemented with 10% ADC and 0.05% Tween 80 at 37 °C with shaking to mid-log phase (OD₅₉₀ ≈ 0.4, ∼2.5 × 10⁸ CFU/mL). For aerobic, fully replicating control cultures, 200 mL of medium was inoculated with 2 mL of the culture (OD₅₉₀ of 0.4) in the flask and incubated at 37 °C on a magnetic stirrer set to rotate at 180 r.p.m. Simultaneously, 400 mL of medium was inoculated with 4 mL of identical culture in a tightly capped flask, which left 200 mL of air space, corresponding to a 0.5 head space ratio. The tightly capped flask culture was then placed on a tissue culture magnetic stirrer set at 70 r.p.m. and incubated at 37 °C to avoid perturbation of the surface. Cultures of the two flasks were gently tipped into the side arm to read OD₅₉₀ daily. Samples for the quantitative proteomics assay were taken after 12 days of vigorous aerated culture (Aera), which then entered the platform stage. Similarly, after 12 days of hypoxic culture (Hypo), samples entered anaerobic nonreplicating persistent stage 2. Using 20 mm screw-capped culture tubes, small volumes of cultures under aeration or hypoxia were collected as samples for western blot or real-time (RT)–polymerase chain reaction (PCR) detection.

**Protein extraction, trypsin digestion, TMT labeling, and enrichment of acetylated lysine peptides**

Cultured cells were harvested and washed twice with cold phosphate-buffered saline (PBS); lysed in 8 M urea supplemented with 1 mM dithiothreitol (DTT), 2 mM ethylenediaminetetraacetic acid (EDTA), 1% Protease Inhibitor Cocktail Set III (Calbiochem, San Diego, CA), and 30 mM nicotinamide; and then disrupted by sonication. Soluble proteins were collected after centrifugation at 20 000 × g for 10 min at 4 °C. Protein concentrations were determined with a BCA protein assay kit (Thermo Fisher Scientific, Waltham, MA) according to the manufacturer’s instructions. For trypsin digestion, the protein sample was diluted by adding 100 mM triethylammonium bicarbonate buffer (TEAB; Sigma-Aldrich, St. Louis, MI) to a urea concentration <2 M; trypsin was then added at a 1:50 trypsin-to-protein mass ratio for the first digestion overnight and at a 1:100 trypsin-to-protein mass ratio for a second 4 h digestion. After trypsin digestion, peptides were desalted using a Strata X C18 SPE column (Phenomenex, Torrance, CA) and vacuum-dried. Peptides were reconstituted in 0.5 M TEAB and labeled according to the manufacturer’s protocol for a TMT 6-plex kit (Thermo Fisher Scientific). To enrich Kac peptides, tryptic peptides dissolved in NETN buffer (100 mM NaCl, 1 mM EDTA, 50 mM Tris-HCl, and 0.5% NP-40, pH 8.0) were incubated with pre-washed antibody beads (PTM Biolabs, Hangzhou, China) at 4 °C overnight with gentle shaking. The beads were washed four times with NETN buffer and twice with ddH₂O. Bound peptides were eluted from beads with 0.1% trifluoroacetic acid. The eluted fractions were combined and vacuum-dried. The resulting peptides were cleaned with C18 ZipTips (Millipore, Billerica, MA) according to the manufacturer’s instructions.

**LC–MS/MS analysis, database search, and bioinformatics analysis**

The tryptic peptides were dissolved in 0.1% formic acid (solvent A), directly loaded onto a homemade reversed-phase analytical column (15 cm length and 75 μm inner diameter). The gradient consisted of an increase from 6 to 23% solvent B (0.1% formic acid in 98% acetonitrile) over 26 min, 23 to 35% over 8 min, an increase to 80% over 3 min, and then holding at 80% for the last 3 min, all at a constant flow rate of 400 nL/min on an EASY-nLC 1000 UPLC system. The peptides were subjected to an NSI source followed by MS/MS in Q ExactiveTM Plus (Thermo Fisher Scientific) coupled online to the UPLC. The electrospray voltage applied was 2.0 kV. The m/z scan range was 350–1800 for a full scan, and intact peptides were detected in the Orbitrap at a resolution of 70 000. Peptides were then selected for MS/MS using an NCE setting of 28, and the fragments were detected in the Orbitrap at a resolution of 17 500. A data-dependent procedure alternated between one MS scan followed by MS/MS in Q ExactiveTM Plus and 20 MS/MS scans with a 15.0 s dynamic exclusion. Automatic gain control was set at 5E4. The fixed first mass was set as 100 m/z.

The resulting MS/MS data were processed using the Maxquant search engine (v.1.5.2.8). Tandem mass spectra were searched against an Mtb database concatenated with a reverse decoy database. Trypsin/P was specified as the cleavage enzyme, allowing up to two missing cleavages. The first search range was set to 5 p.p.m. for precursor ions; the main search range was set to 5 p.p.m. and 0.02 Da for fragment ions. Carbamidomethyl on Cys was specified as a fixed modification, and oxidation on Met was specified as a variable modification. False discovery rate was adjusted to <1%, and the minimum score for peptides was set to >40. For the quantification method, TMT 6-plex was selected. After checking the mass error and length of all identified peptides that fit both requirements, the relative quantitation was divided into two categories: a quantitative ratio of more than 1.5 was considered to be upregulated, while a quantitative ratio of <0.67 was considered to be downregulated for the
bioinformatics analysis. GO annotation of the proteome was performed using the UniProt-GOA Database (http://www.ebi.ac.uk/GOA/). Proteins were categorized into biological processes and molecular functions according to the GO annotation. The KEGG was utilized to annotate pathways. For functional enrichment analysis, we used Fisher’s exact test to assess the enrichment or depletion (two-tailed test) of specific annotation terms among members of the resulting protein clusters. Any terms with adjusted p values below 0.05 in any of the clusters were treated as significant.

Purification of DosR and its derivative mutants

For the purification of DosR, pET28a-dosR, pET28a-dosR<sup>R182K</sup>, and pET28a-dosR<sup>182KΔ</sup> were constructed and verified by sequencing. All confirmed plasmids were transformed individually into <i>E. coli</i> BL21, and recombinant strains were grown in Luria Bertani medium containing 100 μg/mL ampicillin at 37 °C. Isopropyl β-D-1-thiogalactopyranoside was added to the culture at a final concentration of 0.1 mM when the absorbance at 600 nm reached 0.6–0.8. The culture was continuously incubated for 16 h at 18 °C. Afterwards, cells were harvested by centrifugation and stored at −80 °C. Thawed bacteria were resuspended in lysis buffer (50 mM Tris-HCl and 0.5 M NaCl, pH 8.0) and lysed by sonication. Insoluble material was removed by centrifugation for 30 min at 19 000 × g. The soluble extract was purified by Ni<sup>2+</sup> affinity chromatography (Qiagen, Hilden Germany) according to the manufacturer’s instructions. Protein purity was monitored by 12% sodium dodecyl sulfate (SDS)–polyacrylamide gel electrophoresis (PAGE) and Coomassie blue staining. The concentration was determined using a BCA protein assay kit (Thermo Fisher Scientific) with BSA as the standard.

Anti-DosR- and anti-DosR Ac-K182-specific polyclonal antibody preparation

The 6×His-tagged DosR was purified as previously described<sup>46</sup> and used to immunize rabbits for the production of anti-DosR polyclonal antibody. Simultaneously, the acetylated Ac-K182 peptide FLAEKTVK(ace) NYVSRL, a 175–188 amino acid of DosR, was coupled with keyhole limpet hemocyanin and used as an antigen to immunize rabbits. The antiserum was collected and used to immunize rabbits. The antiserum was collected and used as an antigen.

Construction of plasmids and recombinant <i>M. smegmatis</i>

The dosR, poh, sigA, sigB, sigF, and lsr2 genes of <i>Mtb</i> were amplified using <i>Mtb</i> H<sub>37</sub>ΔRv genomic DNA as a template and gene-specific primers with Ndel and HindIII enzyme-cutting sites (Supplementary Table S4) were cloned into the <i>E. coli</i> mycobacterium shuttle plasmid, pVV16, and the K-R mutant. Recombinant plasmids were electrically transformed into <i>M. smegmatis</i>, and transformants were selected on 7H10 agar containing 10% ADC, 50 μg/mL hygromycin B, and 50 μg/mL kanamycin. Recombinant mycobacteria were further cultured in 7H9 broth supplemented with 10% ADC enrichment, 0.05% Tween 80, 50 μg/mL hygromycin B, and 50 μg/mL kanamycin. <i>M. smegmatis</i> harboring the vector backbone, pVV16, was included as a control.

Rv0998 and DosR mutant constructions in <i>Mtb</i>

Gene knockouts of Rv0998 and DosR were generated using allelic exchange and a specialized transducing phage, phAE87 (provided by Dr. LD Yu, Shanghai Institutes for Biological Sciences)°<sup>47</sup>. Constructs for allelic exchange were generated by amplifying the upstream and downstream flanking regions of each of the two genes using the primer pairs listed in Supplementary Table S4. Upstream and downstream flanking regions were cloned into the delivery vector pYUB854. Allelic exchange constructs were incorporated into the phAE87 phage. Phagemid DNA was electroporated into electrocompetent <i>M. smegmatis</i> cells, and these were then plated on 7H10 agar plates to obtain plaques at the permissive temperature of 30 °C. Specialized transducing phages were picked and amplified at 30 °C to generate high titer mycobacteriophages. The desired phage was transduced into the exponential phase of <i>Mtb</i> H<sub>37</sub>ΔRv to delete genes of interest by specialized transduction. The transductants were plated on selective medium consisting of 7H10 agar containing 10% ADC enrichment and 75 μg/mL hygromycin and then cultured at 37 °C. Hygromycin-resistant clones were isolated and analyzed by PCR and RT–PCR to confirm the deletion of target genes. Western blotting was used to confirm the deletion of DosR with anti-DosR polyclonal antibody; anti-SigA (Biolegend, San Diego, CA) monoclonal antibody was used as the reference antibody. The complementation plasmid for the dosR and K182R mutant was generated by ligating the PCR product into the shuttle vector, pVV16. The resulting plasmid was transformed into the Δ dosR strain and plated on a 7H10 plate containing kanamycin at 50 μg/mL. Positive integrants carrying the required insert were screened by colony PCR and validated by RT–PCR.

Growth curve

Mycobacteria were grown to mid-log phase in 7H9 broth with 10% ADC, 0.05% Tween 80, and antibiotics, as required. Growth curves for each strain were determined using a Bioscreen C Growth Curve Instrument (Labsystems Oy, Helsinki, Finland) and a honeycomb plate with
100 wells (Labystems Oy)\textsuperscript{48}. Briefly, 200 μL of each bacterial suspension, adjusted to a similar density, was added to each well and cultured with shaking at 37 °C. The optical density was measured at an absorbance of 590 nm every 2 h. Hypoxic conditions were established by covering each culture with 50 μL of paraffin oil. Cultures were incubated at 37 °C for 7 days for \textit{M. smegmatis}, and 14 days for \textit{Mtb}. Two independent experiments were performed, each in triplicate.

**Quantitative RT–PCR assay**

Bacteria were cultured as previously described, harvested, and disrupted by a bead homogenizer. RNA was isolated using Trizol (Invitrogen). Primers for quantitative PCR are listed in Table S4. Samples were run in triplicate and amplified using real-time SYBR Green quantitative reagents (Toyobo, Osaka, Japan). The relative transcriptional level was determined by the $2^{-ΔΔCt}$ method. The reference gene was 16S rRNA.

**Electrophoretic mobility shift assays**

DNA fragments used for EMSAs were amplified by PCR using \textit{Mtb} H37Rv genomic DNA as a template. The promoter region of \textit{acr} was amplified using the primers \textit{acr}-F and \textit{acr}-R, which are listed in Supplementary Table S4. PCR amplification yielded fragments of 148 bp. Approximately 0.25 μg of DNA in a 20 μL volume was incubated at room temperature for 10 min with 5 μg of purified DosR or derivative mutant protein. The binding buffer used for protein–DNA incubations contained 25 mM Tris–HCl (pH 8.0), 20 mM KCl, 0.5 mM EDTA, 6 mM MgCl\textsubscript{2}, 1 mM DTT, and 1 μg/mL poly I:C. Samples were separated on a 6.5% non-denaturing Tris-glycine polyacrylamide gel and detected by silver staining, as described previously\textsuperscript{49}.

**Purification of Rv0998**

Rv0998 were overexpressed and purified using similar procedure as that for DosR. A plasmid, pET28a-Rv0998/BL21, was constructed and used to overexpress proteins. The 6× His-tagged Rv0998 was purified by Ni\textsuperscript{2+} affinity chromatography, and the amount was estimated by SDS–PAGE and BCA protein assay.

**Immunoprecipitation of DosR**

Previously produced polyclonal antibodies to DosR were used to precipitate specific DosR proteins. For preclari

**Western or dot blots**

Standard western and dot blot procedures were used. Purified recombinant protein (2 μg) or cell extracts (50 μg) were separated by 12% SDS–PAGE and then transferred to a polyvinylidene fluoride membrane. The concentrations of the primary antibodies used in the corresponding blot were anti-DosR (1:1000), anti-DosR Ac-K182 (1:1000), and anti-SigA (1:1000). Horseradish peroxidase-conjugated goat anti-rabbit polyclonal antibody was used as the secondary antibody at a 1:5000 dilution.

**In vitro acetylation**

Assays were performed in a 20 μL total reaction volume containing 50 mM Tris–Cl (pH 8.0), 100 mM NaCl, 100 μM acetyl-CoA, 2 μg Rv0998, and 2 μg of DosR or K182 peptide as the substrate in the presence or absence of 200 μM cAMP. Reactions occurred at 25 °C for 10 min and were stopped by boiling in SDS sample buffer. They were then analyzed by western blot with anti-DosR Ac-K182 antibody.

**Intracellular CFUs in peritoneal macrophages**

Primary peritoneal macrophages were harvested as described previously\textsuperscript{50}. Briefly, C57Bl/6 mice were intra-peritoneally injected with 4% Brewer thioglycollate medium (Sigma-Aldrich). After 3 days, the mice were sacrificed by cervical dislocation, and the cells were isolated by flushing the peritoneal cavity with 10 mL of RPMI 1640 per mouse. Cells were seeded in 24-well dishes, and non-adherent cells were removed by extensive washing with RPMI 1640. The adherent peritoneal macrophages were used for subsequent experiments. Bacteria were diluted to achieve a MOI of 1 or 2 and then incubated with cells for 3 h at 37 °C in 5% CO\textsubscript{2}. Then, uninfected cells were washed twice, and infected cells were incubated in fresh tissue culture medium containing 50 μg/mL amikacin for 2 h post infection. Infected cells were lysed at 24 h with 0.1% (v/v) Triton X-100 in PBS. The number of viable intracellular bacteria (CFU) was determined by serial dilutions and plating out.

**Mouse infection**

Female C57Bl/6 mice (6–8 weeks old) were challenged by i.p. injection with $5 \times 10^6$ CFUs per mouse of the indicated bacterial strains as described previously\textsuperscript{50}. Mice were housed in a specific pathogen-free animal laboratory before being moved into a biosafety level (BSL)-3 laboratory for \textit{Mtb} infection. Four weeks after infection, the mice were sacrificed, and the bacterial loads in the lungs and spleens were determined by CFU counts.
CFU assay
The left portions of lung tissues from infected mice and the whole spleen were homogenized separately in 1 mL of PBS. Homogenates in 10-fold serial dilutions were plated on 7H10 agar supplemented with 10% ADC enrichment and incubated at 37°C. Colonies were counted after 4 weeks.

Histopathology analysis
Half of each lung was fixed in 4% neutral-buffered paraformaldehyde solution for 24 h. Lung tissue was then embedded within paraffin. A series of sections with a thickness of 4–7 μm were then cut and stained with hematoxylin and eosin by standard methods. The pathology was evaluated by pathologists in a blinded manner.

Statistical analysis
Statistical significance was determined with an unpaired two-tailed Student’s t-test at a P < 0.05 level of significance using GraphPad Prism 5.0 software.

Data availability
The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD008587. Material requests for bacterial experiments should be addressed to X.C.H. (huangxiao-chen.1001@163.com) or J.W. (wjtw@163.com). Material requests for proteomics studies should be addressed to H. Y. (yanghua97065@163.com) or W.S. (shfksw@126.com). Material requests for purified proteins or antibodies should be addressed to Z.H.L. (nlzh@126.com) or F.L. (liufeng@sibs.ac.cn). All other material requests should be addressed to H.Y. (yanghua97065@163.com) or B.X.G. (baoxue_ge@tongji.edu.cn).

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Authors’ contributions
B.X.G. and H.Y. designed the study. H.Y., W.S., Z.H.L., R.T.Q., L.H.Q., Z.L.C., J.X.C., X.C.H., and J.W. performed the experiments. B.X.G., H.Y., H.P.L., F.L., R.J.Z., and Y.H. F. analyzed the data. B.X.G., H.Y., W.S., Z.H.L., H.P.L., F.L., X.C.H., and J.W. contributed reagents/materials/analysis tools. B.X.G., H.Y., H.P.L., and R.J.Z. drafted the manuscript. All authors contributed to the final version of the manuscript.

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Conflict of interest
The authors declare that they have no conflict of interest.

Ethical approval
Animal procedures were approved by the Shanghai Pulmonary Hospital, and this study was performed in strict accordance with the National Research Council Guide for Care and Use of Laboratory Animals (SYXK2009-0085). All surgeries were performed under sodium pentobarbital anesthesia, and all efforts were made to minimize suffering. The mouse infection study was performed in an ABSL-3 laboratory of the Second Military Medical University. Animal study protocols were also reviewed and approved by the Institutional Animal Care and Use Committee of the Second Military Medical University (Permit Number: ABSL3-003).

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