Acetylome Analysis Reveals Diverse Functions of Lysine Acetylation in *Mycobacterium tuberculosis*[^5]

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The lysine acetylation of proteins is a reversible post-translational modification that plays a critical regulatory role in both eukaryotes and prokaryotes. *Mycobacterium tuberculosis* is a facultative intracellular pathogen and the causative agent of tuberculosis. Increasing evidence shows that lysine acetylation may play an important role in the pathogenesis of *M. tuberculosis*. However, only a few acetylated proteins of *M. tuberculosis* are known, presenting a major obstacle to understanding the functional roles of reversible lysine acetylation in this pathogen. We performed a global acetylome analysis of *M. tuberculosis* H37Ra by combining protein/peptide pre-fractionation, antibody enrichment, and LC-MS/MS. In total, we identified 226 acetylation sites in 137 proteins of *M. tuberculosis* H37Ra. The identified acetylated proteins were functionally categorized into an interaction map and shown to be involved in various biological processes. Consistent with previous reports, a large proportion of the acetylation sites were present on proteins involved in glycolysis/gluconeogenesis, the citrate cycle, and fatty acid metabolism. A NAD[^+]^−-dependent deacetylase (MRA_1161) deletion mutant of *M. tuberculosis* H37Ra was constructed and its characterization showed a different colony morphology, reduced biofilm formation, and increased tolerance of heat stress. Interestingly, lysine acetylation was found, for the first time, to block the immunogenicity of a peptide derived from a known immunogen, HspX, suggesting that lysine acetylation plays a regulatory role in immunogenicity. Our data provide the first global survey of lysine acetylation in *M. tuberculosis*. The dataset should be an important resource for the functional analysis of lysine acetylation in *M. tuberculosis* and facilitate the clarification of the entire metabolic networks of this life-threatening pathogen. *Molecular & Cellular Proteomics* 13: 10.1074/mcp.M114.041962, 3352–3366, 2014.

*Mycobacterium tuberculosis* was responsible for 1.3 million deaths and 8.6 million new cases of tuberculosis (TB[^1]) worldwide in 2012 (1). This global public health crisis remains a serious problem, with the emergence of drug-resistant *M. tuberculosis*, especially multidrug-resistant and extensively drug-resistant *M. tuberculosis*, and also the emergence of coinfections of TB and human immunodeficiency virus (2, 3). To counter the increasing threat of TB, it is critical to understand fundamental aspects of TB-related biology. Such studies will not only provide new drug targets for the design of novel therapeutic agents, but also facilitate the development of novel diagnostic tools and new vaccines.

Acetylation is one of the important protein modifications and occurs both co- and post-translationally on the ω-amino group at the N terminus of the protein, so-called “N-terminal acetylation,” or on the ω-amino group on the side chain of lysine (4). Lysine acetylation is one of the most common post-translational modifications to proteins in both eukaryotes and prokaryotes. As a dynamic and reversible process, protein acetylation plays important roles in many cellular physiological processes, including cell-cycle regulation and apoptosis, cell morphology (5), metabolic pathways (6–8), protein interactions (9), and enzymatic activity (8, 10). In recent years, great advances have been made in proteomic studies, and a large number of lysine-acetylated proteins have been identified in many eukaryotes, including human (5, 11, 12), rat (13), mouse (11), *Drosophila* (14), *Arabidopsis* (15, 16), *Saccharomyces cerevisiae* (17), and protozoans (18, 19). The global analysis of lysine acetylation has also been reported in bacteria, including *Escherichia coli* (20–22), *Erwinia amylovora* (23), *Bacillus subtilis* (24), and *Salmonella enterica* (6). These acetylome studies have generated large datasets of bacterial

[^5]: The abbreviations used are: TB, Tuberculosis; IFN-γ, interferon gamma; cAMP, cyclic adenosine monophosphate; FA, formic acid; FDR, false discovery rate; KEGG, Kyoto Encyclopedia of Genes and Genomes; GO, Gene Ontology.
proteins acetylated on lysine residues and have demonstrated the diverse cellular functions of lysine acetylation in bacteria. Increasing evidence shows that protein acetylation occurs and plays an important regulatory role in mycobacteria (8, 25–31). For example, Lange et al. reported the N-terminal acetylation of early secreted antigenic target 6 (ESAT-6) protein (31). Rv1151c is reported to be an NAD⁺-dependent protein deacetylase in M. tuberculosis that deacetylates and thus regulates the activity of acetyl-CoA synthase (25, 32). Two cyclic adenosine monophosphate (cAMP)-binding proteins in M. smegmatis and M. tuberculosis (MSMEG_5458 and Rv0998, respectively) show similarity to the GNAT family of acetyltransferases and could acetylate a universal stress protein (USP, MSMEG_4207) (30). Subsequent structural studies revealed the fine mechanisms of how cAMP regulates the protein lysine acetyltransferase in mycobacteria (27, 28). Very recently, reversible lysine acetylation was shown to regulate the activity of several fatty acyl-CoA synthetases in M. tuberculosis (8, 26), and also to regulate acetate and propionate metabolism in M. smegmatis (8, 26). However, to the best of our knowledge, only a few acetylated proteins in M. tuberculosis have been identified, presenting a major obstacle to further understanding the regulatory roles of reversible lysine acetylation in this life-threatening pathogen.

To fill this gap in our knowledge, we undertook a systematic study of the functional roles of lysine acetylation in M. tuberculosis. We performed an acetylomic analysis of M. tuberculosis H37Ra using high-accuracy MS combined with the identification of 226 unique lysine acetylation sites on 137 proteins. This set of M. tuberculosis proteins acetylated on lysine residues supports the emerging view that lysine acetylation is a general and fundamental regulatory process, and is not restricted to eukaryotes. It also opens the way for its detailed functional and evolutionary analysis of lysine acetylation in M. tuberculosis. The identified acetylated proteins that are involved in several important biological processes were functionally categorized into an interaction map. This is the first time that an interaction network of acetylated proteins in M. tuberculosis has been constructed, and should allow us to better understand the significance of acetylation in key cellular mechanisms in M. tuberculosis. To further explore the effects of lysine acetylation on the physiology of M. tuberculosis H37Ra, MRA_1161, the gene encoding the only known protein deacetylase in this bacterium, was deleted. The roles of MRA_1161 in the colony morphology, carbon source utilization, heat stress tolerance, and biofilm formation of M. tuberculosis were analyzed. The effect of lysine acetylation on the immunogenicity of a known immunogen, HspX, was also tested.

**EXPERIMENTAL PROCEDURES**

**Construction of the M. tuberculosis H37Ra ΔMRA_1161 Mutant Strain**—Temperature-sensitive specialized transducing plasmids (33) were constructed and used for the allelic-exchange-mediated deletion of the MRA_1161 gene (also designated cobB), which encodes an NAD⁺-dependent protein deacetylase in M. tuberculosis H37Ra. DNA segments of ~1 kb flanking the gene of interest were amplified with PCR. The purified DNA segments were digested with Van911 and ligation to compatible fragments of the counter-selectable vector p0004S (a kind gift from Dr. William R. Jacobs Jr). The allelic exchange plasmids were selected and propagated following the transformation of E. coli DH5α with the hygromycin-resistance gene. The sequences of the inserted DNA fragments were verified. The allelic exchange plasmids were digested with PacI and ligated to PacI-digested phAE159 (34). The ligation products were packaged using MaxPlax™ Lambda Packaging Extracts (Epicenter Biotechnologies, Madison, WI, USA) and used to transduce E. coli HB101 for propagation as plasmids. The plasmid DNA was purified and transfected into M. smegmatis mc²155 with electroporation for phage propagation. The allelic exchange substrates were delivered to M. tuberculosis H37Ra as previously described (33). The target gene in the mutant strain was replaced with the res-hyg-sacB-res gene cassette. The γδ-resolvase-encoding vector pH532 was then used to resolve the hygromycin-resistance cassette (which was flanked by γδ resolvas recognition sequences) in the mutant strain. The mutant strain was confirmed with PCR and DNA sequencing.

**Cell Culture and Protein Extraction**—Mycobacterium tuberculosis H37Ra and the ΔMRA_1161 mutant were grown at 37 °C in Middlebrook 7H9 liquid culture medium (Difco) supplemented with 10% oleic acid, albumin, dextrose, and catalase (OADC) (Difco), 0.5% glycerol, and 0.05% Tween 80 or Middlebrook 7H10 solid culture medium (Difco) supplemented with 10% OADC (Difco) and 0.5% glycerol. To extract the proteins for MS analysis, M. tuberculosis H37Ra and the ΔMRA_1161 mutant were grown under several conditions in Middlebrook 7H9 liquid culture medium supplemented with 0.5% albumin, 0.5% glycerol, 0.085% NaCl, and 0.05% Tween 80, and harvested at different growth stages. The culture extracts were obtained in: (1) log phase (200 ml), (2) late stationary phase (200 ml), (3) log phase treated with 300 µM diethylenetriamine (D ETA)-NO overnight (200 ml), (4) log phase treated with 5 mM H₂O₂ overnight (200 ml), (5) log phase grown on 0.1% sodium acetate instead of glycerol (200 ml), (6) log phase grown on 0.5% glucose instead of glycerol (200 ml), or (7) log phase grown on 0.2% glucose with glycerol (200 ml).

The cultured cells were harvested by centrifugation at 8000 × g for 10 min at 4 °C, and the cell pellets were washed twice with ice-cold PBS (0.1 M Na₂HPO₄, 0.15 M NaCl, pH 7.5), suspended in 50 ml of chilled lysis buffer (50 mM Tris-Cl [pH 7.5], 100 mM NaCl, 5 mM DTT, 50 mM nicotinamide), and sonicated. Each lysate was centrifuged at 10,000 × g for 30 min at 4 °C and the supernatant was collected. The clarified lysate was precipitated with five volumes of ice-cold acetone, collected by centrifugation, and washed with 80% ice-cold acetone.

**In-solution Trypsin Digestion and Peptide Prefractionation**—Protein extracts (30 mg) were subjected to trypsin digestion, as described previously (35). Briefly, the protein disulfide bonds were reduced with 25 mM DTT (37 °C, 45 min), alkylated with 50 mM iodoacetamide (25 °C, 20 min in the dark), digested with sequencing-grade modified trypsin (1:100 w/w; Promega, Madison, WI) at 37 °C for 5 h, and further digested with trypsin (1:100 w/w) at 37 °C for 20 h. The trypic digestion was quenched by the addition of 0.1% TFA. The solution was clarified by centrifugation at 3000 × g. The resulting peptides were then fractionated and desalted using reversed-phase C₁₈ columns self-packed with C₁₈ material (40 µm, 60 Å pore size; Agilent Technologies, Santa Clara, CA). After the digested peptides were loaded sequentially onto the self-packed C₁₈ columns, the columns were washed five times with 2.0 ml of 0.1% TFA to desalt the
Acetylome Analysis of Mycobacterium tuberculosis

samples, and were eluted with a series of elution buffers (2.0 ml) containing 0.1% TFA and different concentrations of acetonitrile (ACN; 10%, 15%, 20%, 25%, 30%, 35%, 40%, 50%, 60%, or 100%). Each fraction was collected, dried in a SpeedVac, and stored at −20 °C until further analysis.

**In-gel Trypsin Digestion**—Protein extracts (15 mg) were separated by SDS-PAGE on 12% gels (1.5 mm thick, 80 mm wide, 70 mm long), stained with Coomassie Brilliant Blue R250, which were cut into 12 gel slices. Each slice was further diced into ~1 mm3 cubes for in-gel digestion. Each section was completely destained with 50 mM NH4HCO3 in 50% ACN at room temperature. The proteins in each fraction were disulfide reduced, alkylated, and digested with trypsin, as described above. The resulting peptides were transferred into new microcentrifuge tubes and the gels were further sonicated twice with extraction buffer (67% ACN containing 5% TFA). Finally, all peptides were combined and completely dried in a SpeedVac.

**Enrichment of Acetylated Peptides**—Anti-acetyl-lysine antibody (ImmuneChem Pharmaceuticals Inc. and Cell Signaling Technology) was immobilized on Protein A/G PLUS-Agarose Immuno precipitation Reagent (Santa Cruz Biotechnology, Inc., Santa Cruz, CA) at 4 °C for 5 h. The supernatant was removed, and the beads were washed three times with NETN buffer (50 mM Tris-HCl [pH 8.0], 100 mM NaCl, 1 mM EDTA, 0.5% Nonidet P-40). The tryptic peptides were redissolved in NETN buffer and the insoluble particles were removed by centrifugation. The agarose-conjugated anti-acetyl-lysine antibody was added to the peptide solution and incubated in Pierce Centrifuge Columns (Pierce, Rockford, IL) at 4 °C for 8 h with rotary shaking. The beads were washed three times with 1 ml of PBS and then three times with 1 ml of ice-cold water. The bound peptides were eluted with 0.1% TFA in water.

**LC-MS/MS Analysis**—The peptides were separated on the Ultimate 3000 Nano LC System (Dionex, Sunnyvale, CA) and analyzed online using an electrospray ion-trap mass spectrometer (HCTultra; Bruker Daltonics, Bremen, Germany). For each analysis, the sample was trapped on a C18 precolumn (Acclaim® PepMap; 300 μm i.d. × 5 mm; Dionex) and then on a C18 reversed-phase analytical column (Acclaim®; 75 μm i.d. × 150 mm; Dionex). The peptides were eluted from the column with a linear solvent gradient (A: 0.1% formic acid [FA] in water; B: 100% ACN/0.1% FA) for 120 min at a flow rate of 300 nL/min (6 min of 95% buffer A; 95 min gradient from 5–80% buffer B; 10 min of 80% buffer B to wash the column; 5 min gradient from 80–5% buffer B; and a final 5 min of 95% buffer A to equilibrate the column). This LC gradient was used for all mobile phase compositions. The mass spectrometer was operated in the positive ion mode at 2 kV and the capillary temperature was set to 180 °C. Mass spectra were recorded in stand-enhanced mode at a speed of 8100 m/z/s (as specified by the Bruker Esquire Control software). Tandem mass spectra were acquired in the ultra-scan operating mode at 26,000 m/z/s (Bruker Esquire Control software). After a survey scan (enhanced mode, 300–2000 m/z) consisting of four scans, the four most-intense peptides were selected automatically for CID MS/MS (m/z range 100–2000). All MS/MS spectra were acquired using the following parameters: normalized collision energy, 0.5 V; precursor selection threshold, 100,000 Abs; the selected ions were dynamically excluded for 0.25 min.

**Data Analysis and Peptide Identification**—Raw MS data files were processed using the LC/MS software DataAnalysis 4.0 (Bruker Compass™ software) and converted into the Mascot generic format (mgf) files. All MS/MS samples were analyzed with Mascot 2.3 (Matrix Science, London, U.K.) and pFind Studio 2.6 (36, 37) to improve the identification of the acetylated peptides. Both pFind and Mascot were used to search a database of forward and reversed M. tuberculosis H37Rv proteins from National Center for Biotechnology Information (NCBI; http://www.ncbi.nlm.nih.gov/) that contained 4034 protein sequences. The database search was performed with the following parameters: two missed cleavages were allowed for trypsin; carboxamidomethylation (Cys) was set as a fixed modification, whereas oxidation (M) and acetylation (K) were considered variable modifications. The maximum precursor ion mass tolerance allowed was 0.4 Da (monoisotopic) and the fragment mass tolerance was 0.6 Da (monoisotopic). Peptides shorter than six amino acids were excluded. For the search results, the peptide-spectrum matches (PSMs) were filtered based on the score threshold of a 1% false discovery rate (FDR), according to the formula: FDR = 2[(pDecoy)/(pDecoy + pTarget)] (38–40). We applied further strict criteria to the MS2 identification by requiring the peptides to have pFind E scores < 10−5 and Mascot scores > 25, resulting in estimated FDRs that were below 1%. All fragmentation spectra of the acetylated peptides assigned to the corresponding proteins were manually evaluated using a method described by Mann et al. (41, 42). To exclude isobaric trimethylation (43), all the peptides identified based on an acetylated Lys residue were also manually inspected, using a modified method, to identify the neutral loss of ion MH+−59, as described by Mann et al. (42, 44). Finally, to pinpoint the actual acetylated lysine within the identified peptide, the localization probabilities for acetylation sites were calculated from the post-translational modification score algorithm, as described previously (35). Acetylation sites that were occupied with a probability > 0.75 were identified as class I acetylation sites.

**Bioinformatic Analysis**—Biological and functional analyses of our acetylome were performed using only those identified acetylation sites with high confidence (FDR < 1% and localization p value > 0.75). For their functional annotation, the identified acetylated proteins were categorized into biological process and molecular function classes according to Gene Ontology (GO) terms, with the Blast2GO tool. The subcellular localization of the acetylated proteins was analyzed with the PSORTb program, a web-based tool that predicts bacterial protein subcellular localization. To gain information on overrepresentation, a GO enrichment analysis was performed using Cytoscape plugin BiNGO (45, 46), with the previously described parameters (9). The reference GO ontology in the Cytoscape ontology format was created using the whole M. tuberculosis H37Rv proteome, extracted from UniProt-GOA Proteome (http://www.ebi.ac.uk/GOA/proteomes).

The secondary structures of all the acetylated proteins were analyzed with NetSurfP. The mean secondary structure probabilities of the modified lysine residues were compared with the mean secondary structure probabilities of a control dataset containing all the lysine residues of all the acetylated proteins identified in this study. The p values were calculated with the Wilcoxon test.

To analyze the lysine acetylation sites, the ratio of each amino acid flanking the identified acetylation site was calculated as: frequency of residues in the acetyl-13-mers [six amino acids upstream and downstream of the acetylation site]/frequency of residue in total proteome, or frequency of residues in the acetyl-13-mers/frequency of residues in the non-acetyl-13-mers. A position-specific intensity map was generated by plotting the log10 ratios of the frequencies.

A pathway analysis was performed using the Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways database (http://www.genome.jp/kegg) and the DAVID bioinformatics tools. DAVID provided the corresponding information on the p value, count, percentage (%), and fold enrichment for each KEGG pathway. In our data, KEGG pathways with p values < 0.05 were considered significantly enriched pathways. A functional interaction network analysis was performed using interaction data from the STRING database (score > 0.4) and was visualized with Cytoscape v 2.8.3 (http://www.cytoscape.org) (46).

An evolutionary conservation analysis was performed with multi-sequence alignments. The proteins homologous to M. tuberculosis...
H37Ra proteins across 231 species, from Archaea to human, were determined with two-directional BLASTP (47). Eighteen archaea, 152 bacterial, and 61 eukaryotic protein databases were retrieved from Uniprot (http://www.uniprot.org/uniprot), NCBI (http://www.ncbi.nlm.nih.gov), and EnsemBL (http://www.ensembl.org), respectively. The two-directional BLAST alignments were classified as significant if the corresponding BLAST E values were lower than 1E-5.

**Immunoblotting with Anti-acetyl-lysine Antibody—Mycobacterium tuberculosis**

H37Ra and M. tuberculosis H37Ra ΔMRA_1161 were grown, harvested, and lysed as described above. The protein concentrations were determined with the Bradford assay (Bio-Rad). The extracted proteins were separated by 10% SDS-PAGE and transferred to a polyvinylidene difluoride (PVDF) membrane. The membrane was blocked overnight at 4 °C or 37 °C for 2 h with TBST (25 mM Tris-HCl [pH 8.0], 150 mM NaCl, 0.1% Tween 20) containing 5% nonfat dried milk. A primary rabbit anti-acetyl-lysine polyclonal antibody (Cell Signaling Technology) was diluted (1:1000) in TBST/1% nonfat dried milk and incubated with the membrane at 37 °C for 2 h. After the membrane was washed with TBST, it was incubated with alkaline-phosphatase-conjugated goat anti-rabbit antibody (1:5000 dilution) for 1 h at 37 °C, and then detected according to the manufacturer’s instructions.

**Biofilm Formation Assay—Mycobacterium tuberculosis**

H37Ra and M. tuberculosis H37Ra DMRA_1161 were grown in liquid culture medium to mid-log phase, and the cell density of each culture was adjusted to OD600 = 0.8. Then both strains were subcultured (1:100) in fresh 7H9 medium (10% OADC, 0.5% glycerol, 0.05% Tween 80) in 25 cm² cell culture flasks (NEST Biotechnology) and incubated at 37 °C for 3–5 weeks without shaking. All experiments were performed in triplicate.

**Carbon Source Utilization Experiments—Mycobacterium tuberculosis**

H37Ra and M. tuberculosis H37Ra DMRA_1161 were grown to mid-log phase in liquid culture medium, collected by centrifugation at 8000 × g for 10 min, and washed three times with 7H9 medium + 0.05% Tween 80 (7H9T) to remove the remaining medium, before they were resuspended in 7H9T and the OD600 adjusted to 0.5. Aliquots (100 μl) of the suspension culture were used to inoculate 10 ml of the following media: (1) 7H9 liquid culture medium supplemented with 10% OADC, and 0.5% glucose, 0.05% Tween 80; (2) 7H9 liquid culture medium supplemented with 0.5% albumin, 0.05% Tween 80, and different concentrations of acetate (5 mM, 10 mM, 30 mM, or 50 mM).

**Heat Shock Assay—Mycobacterium tuberculosis**

H37Ra and M. tuberculosis H37Ra DMRA_1161 were grown in 7H9 medium to mid-log phase, and the cell density of each culture was adjusted to OD600 = 0.5. The cultures were then heat treated in a water bath at 53 °C. Aliquots were taken and serial dilutions were made before plating (48). The bacterial colony-forming units (cfu) were determined after 25 d at 37 °C. All the experiments were performed in triplicate.

**Enzyme-linked Immunospot (ELISPOT) Tests of Two Synthetic Peptides Derived from HspX of M. tuberculosis**

H37Ra—Two peptides (6291 and A-6291) were synthesized (GL Biochem Ltd, Shanghai, China) based on the amino acid sequence of HspX from M. tuberculosis H37Ra. The two peptides shares the same sequence, spanning residues 62–91 of HspX (PDFKDVIIMVRRDQGLTKAERTEQKDFGR), except that all three lysines in A-6291 are acetylated, according to our MS data. The ELISPOT test (T-SPOT TB 96 assay, catalog no. TB.200; Oxford Immunotec) was performed according to the instructions of the manufacturer. Twenty-six tuberculosis patients were recruited from Foshan Fourth People’s Hospital, and the data for the patients enrolled in this experiment are listed in supplemental Table S1. The study was approved by and performed under the guidelines of the Ethical Committee of Wuhan Institute of Virology (Chinese Academy of Sciences). Written informed consent was obtained directly from each participant. Whole blood (5 ml) was drawn from each individual with venipuncture and stored at room temperature. After isolation by Ficoll-Hypaque (MD Pacific, Tianjin, China) density-gradient centrifugation, the cloudy peripheral blood mononuclear cells (PBMCs) harvested were washed twice with prewarmed RPMI 1640 medium (MD Pacific) or AIM-V medium (Invitrogen, Karlsruhe, Germany). Adequate AIM-V medium was used to resuspend the PBMCs and blood cell counting plates were used to count the cells. Four wells in a 96-well microtiter plate, precoated with monoclonal antibody directed against interferon gamma (IFN-γ; provided with the kit), were seeded with the previously prepared PBMC suspension. The positive control was stimulated with phytohemagglutinin (provided with the kit). One well was a no-peptide control, containing PBS and medium as stimulants, and was included to detect contamination. Identical concentrations (10 μg ml⁻¹) of the acetylated peptide A-6291 or the normal peptide 6291, dissolved in PBS, were added to the other two wells. The plate was then incubated at 37 °C under 5% CO₂ for 16–20 h, washed thoroughly several times with PBS, and incubated at 4 °C for 1 h with 50 μl of alkaline-phosphatase-conjugated anti-IFN-γ monoclonal antibody. The plate was then washed thoroughly again with PBS and 50 μl of substrate solution was added to each well. After incubation for 7 min at room temperature, the reaction was stopped with tap water and the plate was dried to visualize the spots. The total numbers of spots were counted and the spot-forming cells in each well was analyzed with a Bioreader 5000 (Dakewe Biotech Company, Shenzhen, China).

**RESULTS AND DISCUSSION**

**Identification of Lysine-acetylated Proteins in M. tuberculosis**

H37Ra—Reversible lysine acetylation is emerging as a major regulatory mechanism in metabolism and is an important post-translational modification in both prokaryotes and eukaryotes (7, 49). To evaluate the diversity and relative abundance of acetylated proteins in M. tuberculosis H37Ra, a ΔMRA_1161 (encoding lysine deacetylase) mutant was constructed (map of the MRA_1161 gene knockout and PCR verification of the knockout strain are shown in Fig. 1A and 1B) and immunoblots of the bacterial cell lysates were probed with an anti-acetyl-lysine antibody. An immunoblotting analysis of protein extracts from the wild-type and ΔMRA_1161 mutant strains detected multiple protein bands, spanning a wide mass range, with the antibody in both extracts, and many substrates showed increased acetylation levels in the ΔMRA_1161 strain (Fig. 1C and 1D). These results suggest that the antibody used is specific and that many lysine-acetylated proteins remain to be identified in M. tuberculosis. After primary verification, the protein extracts were digested both in solution and in gel with sequential rounds of trypsin treatment. To collect low-abundance modification sites, the digested peptides were fractionated and desalted on self-packed reversed-phase C₁₈ columns. The lysine-acetylated peptides were immune-enriched with the anti-acetyl-lysine antibody and identified with nano-LC/MS/MS in an electrospray ion-trap mass spectrometer (HCTultra), as summarized in Fig. 1E. The raw data were processed with two different search algorithms (MASCOT and pFind). All spectra containing acetylation modifications were manually inspected to check for the diagnostic b-type or y-type ion series, as described by Mann.
et al. (see criteria under “Experimental Procedures”). All the spectra were also checked for the neutral loss of ion “MH−59,” which can be used as a unique marker for trimethylation (43), and peptides containing this marker were removed from the analysis. In total, 226 lysine acetylation sites were identified in 137 proteins at an FDR < 0.01 for peptides with high confidence. This constitutes the first acetylation dataset obtained for the mycobacteria. Details of the modification-specific peptides, identified lysine acetylation sites and proteins are shown in supplemental Table S2. The annotated spectra of all the acetylated peptides identified are presented in supplemental Fig. S1. To understand the scope of acetylation in bacteria, we compared the M. tuberculosis H37Ra acetylsome with the bacterial acetylosomes reported by other groups. The results are presented in Table I. Among the highly acetylated proteins were the chaperone protein GroEL, antigen Hsp20.

Fig. 1. Synopsis of lysine acetylation proteomics. A, Map of the MRA_1161 gene knockout, map of the MRA_1161 gene region in the genome and its corresponding region in the conditional mutant strain. MRA_1160 and MRA_1162 are the upstream and downstream regions of the MRA_1161 gene, respectively; res, γ resolvase site; hyg, hygromycin-resistance gene; sacB, sucrose counterselectable gene. The primers used for PCR amplification are indicated by arrows and the expected sizes of the PCR products are indicated by straight lines. B, PCR verification of the MRA_1161 gene knockout strain. Agarose gel electrophoresis was used to verify the PCR products amplified from the genomic DNA of different mycobacterial strains. M, DNA marker; lane 1, M. tuberculosis H37Ra; lane 2, M. tuberculosis H37Ra ΔMRA_1161; lane 3, M. tuberculosis H37Ra ΔMRA_1161 (hyg-sacB). PCR products are indicated by white arrows. SDS-PAGE C, and immunoblotting D, analyses of protein lysates from the M. tuberculosis H37Ra (lane 1) and MRA_1161 knockout (lane 2) strains; M, protein marker. E, Schematic representation of the sequential steps used for the global profiling of lysine acetylation in M. tuberculosis.
(HspX), and many enzymes involved in diverse metabolic pathways (supplemental Table S2). Of the acetylation sites identified in this study, 46.02% could be successfully predicted with PAIL (prediction of acetylation on internal lysines), a predictor developed from the 249 experimentally verified acetylation sites of 92 distinct proteins in the scientific literature (50) (supplemental Table S2). The raw data (including a converted MASCOT generic format file) have been uploaded onto the public database PeptideAtlas (http://www.peptideatlas.org) (51, 52) and can be accessed through the link http://www.peptideatlas.org/PASS/PASS00238.

**Table I**

| Organism          | Strain     | Genome size (Mb) | Genes | Proteins | No. ac-proteins | No. ac-peptides | No. ac-sites | ref   |
|-------------------|------------|------------------|-------|----------|-----------------|-----------------|--------------|-------|
| *Escherichia coli*| MG1655     | 4.64             | 4496  | 4146     | 91              | 138             | 138          | (21)  |
| *Escherichia coli*| DH10       | 4.69             | 4352  | 4124     | 349             | 1034            | 1070         | (22)  |
| *Salmonella enterica*| G2466     | 4.95             | 4732  | 4525     | 191             | 235             | 261          | (6)   |
| *Erwinia amylovora*| Ea1189/Ea273 | 3.91             | 3712  | 3565     | 96              | 167             | 141          | (23)  |
| *Bacillus subtilis*| 168        | 4.22             | 4422  | 4176     | 185             | 332             | 332          | (24)  |
| *Thermus thermophilus*| HB8        | 2.12             | 2245  | 2238     | 128             | 198             | 201          | (78)  |
| *Geobacillus kaustophilus*| HTA426    | 3.59             | 3655  | 3653     | 114             | 253             | 253          | (79)  |
| *Mycobacterium tuberculosis*| H37Ra     | 4.42             | 4084  | 4034     | 137             | 230             | 226          | This work |

| Acetylome Analysis of *Mycobacterium tuberculosis* |

Comparison of *Mycobacterium tuberculosis* H37Ra acetylome with other published bacterial acetylomes

To further investigate, the distributions of the acetylated proteins were detected with both methods (Fig. 2C). 105 acetylation sites and 72 acetylated proteins were identified with SDS-PAGE, 187 acetylation sites and 115 acetylated proteins were identified with LC, and 66 acetylation sites and 50 acetylated proteins were shown in a Venn diagram (Fig. 2C). In the GO molecular function category, the acetylated proteins identified were assigned to several groups, including catalytic activity (46.12%), binding (39.66%), and unknown (3.88%). The remaining proteins were assigned to specific functions, such as structural molecules (3.45%), electron carrier activity (2.59), transporter activity (1.72%), antioxidant activity (1.29%), molecular transducer activity (0.85%), and nucleic-acid-binding transcription factors (0.43%) (Fig. 2E). To investigate the distribution of the acetylated proteins across the cellular compartments, we evaluated them using the PSORTb program according to the confidence values for each localization site. Our results show that the majority of acetylated proteins occurred most frequently in the cytoplasm (106); many fewer occurred on the cytoplasmic membrane (13) or the cell wall (1) (Fig. 2F; supplemental Table S3). A GO enrichment analysis was also performed to identify the biological processes and molecular functions in which the acetylated proteins were involved, and to examine the preferential subcellular localizations of the acetylated proteins. Like previous findings (6, 21), our data suggest that the acetylated proteins of *M. tuberculosis* H37Ra were significantly enriched for several GO biological processes, including the generation of precursor metabolites and energy, acetyl-CoA catabolic processes, metabolic processes, the citrate cycle, acetyl-CoA metabolic processes, cofactor catabolic processes, coenzyme catabolic processes, aerobic respiration, cellular respiration, energy derivation by the oxidation of organic compounds, cellular protein metabolic processes, translation, protein folding, and protein metabolic processes. From the GO enrichment analysis, we also found that the lysine-acetylated proteins were significantly enriched for two specific functions, such as ligase activity ($\rho = 1.06 \times 10^{-4}$) and transferase activity, the transfer of acyl groups, the conversion of acyl groups to alkyl groups on transfer ($\rho = 9.11 \times 10^{-5}$). In the GO cellular component categories, we found that a considerable proportion of the acetylated proteins identified were localized intracellularly ($\rho = 3.21 \times 10^{-10}$); cytoplasm ($\rho = 1.63 \times 10^{-9}$) and intracellular ($\rho = 0.01 \times 10^{-7}$) (Fig. 2G; supplemental Table S4). The identified acetyl proteins were also assigned to KEGG metabolic pathways, which indicated that lysine acetylation occurs on many enzymes involved in energy metabolism (Fig. 2G; supplemental Table S5).

**Analysis of Lysine Acetylation Sites**—The magnitude of our dataset allowed us to identify site-specific acetylation motifs. We compared the acetylated sites to the nonacetylated lysine...
Fig. 2. **Overview of the acetylated proteins identified in *M. tuberculosis* H37Ra.** A, A pie chart illustrates the number of lysine acetylation sites identified per protein. B, Numbers of acetylated proteins and C, acetylation sites identified with SDS-PAGE and LC and their overlap are shown in the individual circles of the Venn diagram. D-F, Pie chart representations of the distributions of the acetylated proteins identified according to their biological processes, molecular functions, and cellular localization. E, Enrichment analysis. The differently colored bars indicate the corrected p values for the enrichment of the annotations.
residues in this dataset to identify any bias that may occur in the amino acids neighboring the acetylation sites. Thus, we checked the 12 residues flanking the acetylated sites for the overrepresentation of specific amino acids relative to their expression in the proteome background distribution (Fig. 3A).

An intensity map was generated as described previously (53). This analysis showed preferences for glutamic acid at the +1 position, phenylalanine at the +1 position, and lysine at the +3 position. This strong bias for a specific acetylation-site motif is quite different from the situation in other representative model organisms. No significant overall sequence recognition motif was detected among the acetylation sites of *S. enterica* (6) or *Erwinia amylovora* (23). A strong bias for the flanking amino acid residue at the +1 position was observed in other bacteria, such as the histidine or tyrosine amino acid residue preferred by *E. coli* (21) and the glutamic acid, aspar-
Acetylome Analysis of Mycobacterium tuberculosis

tic acid, lysine, or proline preferred by B. subtilis (24). In contrast to the bacterial acetylome, there are consensus motifs in the Drosophila and human acetylomes. For example, glycine and glutamic acid are the preferred amino acids beyond the -1 position, whereas tyrosine, phenylalanine, and proline occur frequently at the +1 position in these organisms. Lysine residues at the -5, -6, +3, +4, +5, and +6 positions have also been observed (14). The differences in the preferred amino acid residues flanking the acetylation sites in the M. tuberculosis H37Ra acetylome proteins suggest that this modification is catalyzed by a new subset of acetyltransferases, with unique substrate preferences. In contrast to phosphorylation sites, acetylation sites frequently occur in regions with ordered secondary structures, as previously reported (54). For a more detailed investigation of the local secondary structures of the protein sequences surrounding acetylation sites, we compared the secondary structures of acetylated lysines to those of all lysines. In general, the acetylated lysines were more frequently found in structured regions ($\rho = 1.08 \times 10^{-2}$ for $\alpha$-helices and $\rho = 4.09 \times 10^{-2}$ for $\beta$-strands) and less frequently in unstructured regions ($\rho = 8.52 \times 10^{-4}$ for coils) (Fig. 3B). We also evaluated the acetylated lysine sites for solvent accessibility; 84.3% of identified acetylated lysine sites were exposed to the protein surface, compared with 78.7% of all lysine residues ($\rho = 3.08 \times 10^{-2}$). This result suggests that the highly conserved acetylated sites have local structural preferences, which are quite different from those of phosphorylation sites.

To measure acetylome conservation, we aligned the best-scoring homologous proteins from every species with all the identified acetylated proteins and all the proteomes. The orthologs that shared the highest homology with each other in both directions were counted and reported as the percentage of the total number of proteins in all the species analyzed. The percentages of M. tuberculosis H37Ra homologues of the acetylated and nonacetylated proteins in diverse organisms are shown in Fig. 3C. The acetylated proteins of M. tuberculosis H37Ra showed significantly higher conservation across the evolutionary tree, and especially among bacteria, than the nonacetylated proteins. More than half the M. tuberculosis H37Ra acetylated proteins were conserved in other bacterial species. Interestingly, 76 acetylated proteins of M. tuberculosis H37Ra were found to be conserved in other bacterial acetylomes (supplemental Table S6). Because lysine acetylation is an ancient post-translational modification and is also highly conserved throughout all species, we speculate that the functions of the detected acetylated proteins could be essential the mycobacteria.

Functional Interaction Networks of Acetylated Proteins in M. tuberculosis H37Ra—Protein interaction networks were established to identify the physical and functional interactions among the identified acetylated proteins using protein interaction information from the STRING database. As shown in Fig. 4, 137 proteins in our data set were mapped to the protein interaction database of M. tuberculosis (supplemental Table S7). The interaction network presents an overview of how acetylated proteins perform various types of functions in M. tuberculosis. Five highly interconnected clusters of acetylated proteins were retrieved using the MCODE algorithm (supplemental Table S8). Interestingly, the first high-ranking complexes we identified (Cluster I) consist of ribosome-associated proteins (Fig. 4B), whereas Clusters II-IV (Fig. 4C–4E) consist of metabolism-associated proteins, especially proteins involved in the citrate cycle. These findings show that protein translation is probably regulated by lysine acetylation in M. tuberculosis, and that most of the acetylated proteins identified are involved in central metabolic pathways responsible for the control of key metabolic processes. Interestingly, one of the known immunogens of M. tuberculosis, Acr (HspX), interacts tightly with other function-associated proteins (Fig. 4F). Together with other acetylation data, these protein clusters will help to decipher the physiological roles of lysine acetylation in M. tuberculosis.

Acetylation of Enzymes Involved in the Central Metabolism and Lipid Metabolism Pathways of M. tuberculosis—It is well established that most enzymes involved in the central metabolic pathways are lysine acetylated in various organisms (6, 20–24). In this study, we found that many of the acetylated proteins identified in M. tuberculosis H37Ra are involved in central metabolic pathways. As shown in Fig. 5A, a large proportion of enzymes that are involved in glycolysis/glucogenesis and the citrate cycle are lysine acetylated, which is consistent with our previous understanding of lysine acetylation (6, 20–24). Therefore, lysine acetylation may regulate the central metabolic pathways at multiple sites in M. tuberculosis. Among the enzymes shown to be acetylated in the central metabolic pathways, eight are components of the citrate cycle (12, 55, 56) and two enzymes, isocitrate lyase (ICL-1) and malate synthase G (GlcB), are involved in the glyoxylate pathway. In M. tuberculosis, both copies of the isocitrate-lyase-encoding genes are required for the survival of the bacterium in vivo (57, 58). A structural analysis of ICL showed that the protein has an active-site loop region (residues 185–196), which contains the ICL signature sequence (K189KCGH193) that controls its enzymatic activity (59). The identified acetylated lysine residue (K189) occurs in this region, so the acetylation of this residue may lead to a conformational change in the protein and therefore affect its enzymatic activity. GlcB is a secreted protein and enhances the adherence of the pathogen to lung epithelial cells, so it is also related to the bacterium-host interaction (60).

Apart from the enzymes in the citrate cycle and the glyoxylate pathway, many enzymes related to fatty acid metabolism were shown to be acetylated in M. tuberculosis, including several enzymes involved in the $\beta$-oxidation of fatty acids and several involved in fatty acid biosynthesis (Fig. 5A and 5B). The enzymes involved in fatty acid metabolism that were shown to be acetylated are significantly enriched for ligase
Protein interaction network of the *M. tuberculosis* H37Ra acetylome. A, The complete acetylation interaction network obtained with STRING (9.0) at confidence scores ≥ 0.4. B–F, The five most highly ranked and tightly connected network clusters obtained with an MCODE analysis are color-coded and then rendered as separate modules.

**Fig. 4.** Protein interaction network of the *M. tuberculosis* H37Ra acetylome.
activity and transferase activity (Fig. 2E; supplemental Table S4). In M. tuberculosis, fatty acids and their metabolic derivatives are important components of the cell wall complex, which is related to the pathogenicity of the bacterium (61, 62). In fact, it has been reported that reversible acetylation can modulate the activity of several fatty-acid-CoA ligases (FadD2, FadD5, and FadD13) in M. tuberculosis (7), which play important roles in the fatty acid metabolism of the bacterium. Therefore, our proteomic data suggest that acetylation plays an important role in the fatty acid metabolism of M. tuberculosis.

Deletion of MRA_1161 affects Colony Morphology and Biofilm Formation, and Increases Tolerance of Heat Shock Stress in M. tuberculosis H37Ra—Previous studies have shown that the lipid metabolism of mycobacteria correlates with colony morphology and biofilm formation (63–67). About 250 distinct enzymes are involved in the fatty acid metabolism of M. tuberculosis (68), and 25 of these were shown to be acetylated in this study, including two enzymes (FabG1 and HadB) essential to the type II fatty acid synthase (FAS II) system. Very recently, Nambi et al. found that acetylation affects the activity of several fatty-acid-CoA ligases in M. tuberculosis (8). Therefore, we speculate that the lysine acetylation of proteins may affect the colony morphology and biofilm formation of M. tuberculosis H37Ra. Therefore, we deleted the only known copy of the protein-deacetylase-encoding gene, MRA_1161, and showed that its deletion affected the colony morphology of M. tuberculosis H37Ra on solid 7H10 plates. The ΔMRA_1161 mutant appeared to be more granular than its parental strain (Fig. 6A and 6B) and showed defective biofilm formation after 4 weeks of growth in 7H9 medium (Fig. 6C and 6D). However, during the first 2 weeks of growth, its biofilm formation did not differ from that of the control. Our results suggest that lysine acetylation regulates fatty acid metabolism during the late growth stage, which is consistent with a previous report that the lysine acetylation of proteins is more abundant during the stationary phase of E. coli (20). Our results suggest that lysine acetylation affects the colony morphology and biofilm formation by regulating the activities of enzymes involved in the fatty acid metabolism of M. tuberculosis H37Ra.

Previous research has shown that the deletion of cobB in E. coli affects the stress responses of the bacterium, including its responses to oxidative stress and heat stress (69). Therefore, we tested the effects of MRA_1161 deletion on heat stress in M. tuberculosis H37Ra. In contrast to the wild-type strain, the ΔMRA_1161 mutant showed marked resistance to heat stress (53 °C) (Fig. 6E), similar to the previous observations in E. coli, suggesting that lysine acetylation affects the stress responses of bacteria. Although some stress proteins, including Dnak and GroEL, were shown to be acetylated in M. tuberculosis H37Ra, the mechanism underlying the effect of...
protein acetylation on the heat stress response remains unclear, and warrants further investigation.

Deletion of MRA_1161 affects Carbon Source Utilization by M. tuberculosis H37Ra—As described above, many enzymes involved in glycolysis/gluconeogenesis, the citrate cycle, and the lipid metabolism pathways of M. tuberculosis H37Ra are lysine acetylated, which is consistent with previous reports of E. coli and S. enterica (6, 20–22). In E. coli, S. enterica, M. bovis BCG, and M. smegmatis, the counterpart of MRA_1161, Pat/CobB, is responsible for the reversible acetylation of those metabolic enzymes, so the deletion of cobB and pat affects the carbon source utilization of these bacteria (6, 8, 26, 70, 71). Therefore, the effect of MRA_1161 deletion on the carbon source utilization of M. tuberculosis H37Ra was analyzed. Our results show that the ΔMRA_1161 mutant strain grew more slowly than the wild-type strain when glucose was used as the sole carbon source (supplemental Fig. S2A), which differs from previous observations of S. enterica (6). This result suggests that, although carbon source utilization is commonly regulated by reversible protein lysine acetylation, the fine mechanisms of this regulation may differ among different species. Further studies are required to determine how reversible protein lysine acetylation regulates carbon source utilization in M. tuberculosis. The effect of MRA_1161 deletion on acetate utilization also differed from that in other bacteria, including S. enterica and E. coli (70, 71). When acetate was used as the sole carbon source, the ΔMRA_1161 mutant showed no marked changes in its growth rate with different concentrations of acetate (5, 10, 30, or 50 mM) (supplemental Fig. S2B–S2E), although previous studies have shown that Rv1151c, an orthologue of MRA_1161, modulates the activity of the acetyl-CoA synthetase of M. tuberculosis in vitro by deacetylation (32). This implies either the existence of other protein deacetylases or that a different mechanism regulates acetate utilization in M. tuberculosis H37Ra.

**Fig. 6.** Effects of MRA_1161 deletion in M. tuberculosis H37Ra. A, Colony morphology of M. tuberculosis H37Ra and ΔMRA_1161 mutant. Bacterial cells were plated to generate single colonies on solid medium and incubated for 4 weeks at 37°C. C, Biofilm formation of M. tuberculosis H37Ra and ΔMRA_1161 mutant. Bacterial cells were grown in 7H9 medium (10% OADC + 0.5% glycerol + 0.05% Tween 80). D, Effects of MRA_1161 deletion on the heat shock response of M. tuberculosis.

**Fig. 7.** Results of ELISPOT tests. A, Images of the PVDF microfiltration membrane in each well. Each patient was assigned four reaction wells: row a, the no-peptide negative control; row b, peptide 6291; row c, A-6291; row d, the positive control. B, Total number of the spot-forming cells in each well according to A. Subjects 3, 13, and 25 (indicated in red boxes) showed different responses to peptides 6291 and A-6291, indicating the different immunogenicity of the two peptides.
Effect of Acetylation on the Immunogenicity of a Peptide Derived from HspX—Interestingly, the acetylome data generated in this study showed that 45 secretory proteins are acetylated in *M. tuberculosis* (supplemental Table S9). In many situations, these secretory proteins can stimulate the immune responses of the host (72). Therefore, we examined whether acetylation affects the immunogenicity of secretory proteins. Among the 45 secretory proteins listed in supplemental Table S9, *M. tuberculosis* heat-shock protein X (HspX) was shown to be acetylated at multiple sites. As a molecular chaperone, HspX is linked to the latent infection of *M. tuberculosis*, because its deletion results in increased bacterial growth in both mice and macrophages (73). Moreover, there is evidence that HspX of *M. tuberculosis* induces an effective immune response in its host, and the immunization of mice with recombinant HspX protein resulted in enhanced levels of IFN-γ. HspX also increases the expression of IFN-γ in TB patients (74–77). Therefore, HspX was selected for further investigation. Because it is difficult to prepare acetylated and deacetylated full-length HspX in vitro, two 30-mer peptides with the same sequence (spanning residues 62–91 of HspX) were synthesized, one with all three lysine residue acetylated (peptide A-6291) and the other with no acetylation (peptide 6291).

In total, 26 blood samples were collected from patients with active TB or patients with suspected TB, and ELISpot tests were used to compare the immunogenicity of 6291 and A-6291. As shown in Fig. 7, among the 26 samples tested, three (samples 3, 13, and 25) responded positively to peptide A-6291. As shown in Table S1, these results suggest that acetylation affects the immunogenicity of the secretory proteins of *M. tuberculosis*, which warrants further investigation.

**CONCLUSION**

In conclusion, our results provide the first extensive data on lysine acetylation in *M. tuberculosis*. We identified a total of 230 unique acetylated peptides from 137 *M. tuberculosis* proteins. Further functional studies revealed that the reversible lysine acetylation of proteins is involved in the carbon source utilization, colony morphology, biofilm formation, and stress response of *M. tuberculosis*, and may also affect the immunogenicity of secretory proteins. Therefore, our results provide novel insight into the range of functions regulated by lysine acetylation in *M. tuberculosis*. The dataset provided should help to determine the physiological functions affected by lysine acetylation and facilitate the clarification of the entire metabolic networks of *M. tuberculosis*.

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**This article contains supplemental Tables S1 to S9 and Figs. S1 and S2.**

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