Expression of a Subset of Heat Stress Induced Genes of *Mycobacterium tuberculosis* Is Regulated by 3',5'-Cyclic AMP

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Abstract

*Mycobacterium tuberculosis* (Mtb) secretes excess of a second messenger molecule, 3',5'-cyclic AMP (cAMP), which plays a critical role in the survival of Mtb in host macrophages. Although Mtb produces cAMP in abundance, its exact role in the physiology of mycobacteria is elusive. In this study we have analyzed the expression of 16 adenylate cyclases (ACs) and kinetics of intracellular cAMP levels in Mtb during *in vitro* growth under the regular culture conditions, and after exposure to different stress agents. We observed a distinct expression pattern of these ACs which is correlated with intracellular cAMP levels. Interestingly cAMP levels are significantly elevated in Mtb following heat stress, whereas other stress conditions such as oxidative, nitrosative or low pH do not affect intracellular cAMP pool *in vitro*. A significant increase in expression by >2-fold of five ACs namely Rv1647, Rv2212, Rv1625c, Rv2488c and Rv0386 after heat stress further suggested that cAMP plays an important role in controlling Mtb response to heat stress. In the light of these observations, effect of exogenous cAMP on global gene expression profile was examined by using microarrays. The microarray gene expression analysis demonstrated that cAMP regulates expression of a subset of heat stress-induced genes comprising of *dnaK*, *grpE*, *dnaJ*, and *Rv2025c*. Further we performed electrophoretic mobility shift assay by using cAMP-receptor protein of Mtb (CRP^M^), which demonstrated that CRP^M^ specifically recognizes a sequence –301AGCGACCCTAGCCACAG–286 in 5'-untranslated region of *dnaK*.

Introduction

*Mycobacterium tuberculosis* (Mtb), the causative agent of disease tuberculosis (TB) has evolved a clever strategy of intoxicating host macrophages by secreting a signaling molecule, 3',5'-cyclic adenosine monophosphate (cAMP) [1,2]. Cyclic AMP is continuously produced by Mtb during *in vitro* growth [3], probably due to presence of multiple adenylate cyclases (ACs). Genome sequence of Mtb reveals the presence of 16 ACs in Mtb H₃₇Rv strain, 10 of which have been biochemically characterized *in vitro* [4–7]. Fusion sequence of Cya homodimer contains six transmembrane regions and a catalytic domain, which corresponds to one half of the mammalian adenyl cyclases [10]. Other functionally characterized ACs of Mtb are: pH-sensing Rv1264 which contains an autoinhibitory N-terminal domain [11–12]; Rv1318c, Rv1319c, Rv1320c and Rv3645 containing membrane anchored HAMP (present in Histidine kinases, Adenylate cyclases, Methyl accepting proteins and Phosphatases) region [13]; Rv0386 whose adenyl cyclase catalytic domain is fused to an AAA-ATPase and a helix-turn-helix DNA-binding domain [14]; Rv1647 and Rv2212 which attain complete activity only in the presence of detergent and unsaturated fatty acids, respectively [15,16]; and Rv1900c which forms asymmetric homodimers [17]. Cyclic AMP is also secreted into host cells during infection which perturbs signaling pathways and affects bacterial persistence and killing by host macrophages [1,2].

In prokaryotes cAMP activates the function of a transcription factor known as cAMP-receptor protein or CRP which recognizes a specific sequence in the 5'-untranslated region (5'-UTR) and subsequently regulates the mRNA synthesis of candidate genes [4]. *In silico* analysis predicts 10 putative cNMP-binding proteins in Mtb [4]; two of these proteins encoded by *Rv3676* (known as cAMP-receptor protein of *M. tuberculosis*, CRP^M^) and *Rv1675c* (annotated as Cmr, for cAMP and macrophage regulator) function as cAMP-responsive transcription factors that regulate expression of multiple genes by direct binding to their promoter regions [18–24]. In addition to regulating mycobacterial pathogenesis other important biological processes are also regulated by this signaling molecule. Exogenous cAMP stimulates the expression of galactokinase in the presence of glutamate and galactose in *Mycobacterium smegmatis* (Msm), which is otherwise not induced by galactose alone [25]. In *M. bovis* BCG cAMP regulates the expression of five
proteins namely Rv1265, Rv2971, GroEL2, PE_PGRS6a, and malate dehydrogenase [26]. Very recently it is shown that cAMP plays a role in acetylation of stress proteins and acetyl-CoA synthetase [27–29], which suggests that cAMP is critical in functioning of central metabolic pathways of Mtb.

Though Mtb produces significant concentration of cAMP which is also secreted into extracellular environment [30,31], direct role of cAMP in the physiology of Mtb is lacking. By using a systematic approach in this study we measured the intracellular cAMP levels and expression of ACs in Mtb during its in vitro growth in regular culture medium as well as under different stress conditions. By performing a whole genome microarray analysis, we studied the effect of cAMP on global gene expression profile of Mtb. Further, direct effect of cAMP on the expression of candidate genes was validated by performing electrophoretic mobility shift assay (EMSA). Our results demonstrate that in Mtb cAMP levels are significantly elevated after heat stress which in-turn regulates the expression of a subset of heat stress-induced genes encoding chaperones DnaK, GrpE and DnaJ respectively, by facilitating the direct binding of CRP to the promoter region of dnaK operon.

**Results**

**Analysis of intracellular cAMP levels in Mtb during in vitro growth**

Cyclic AMP is known to exert an array of regulatory functions which advocate that the cellular concentration of cAMP must itself be subject to control by culture conditions. Here we estimated intracellular cAMP levels in pathogenic Mtb grown in 7H9 culture medium supplemented with 1x OADC (oleic acid, albumin, dextrose and catalase), 0.5% glycerol and 0.02% tween-80, at different growth stages. Lysates were prepared by boiling the bacterial pellets in 0.1 M HCl to avoid degradation of cyclic nucleotides during extraction by phosphodiesterase (PDE) of Mtb, and cAMP was measured by ELISA as described in materials and methods. Our results demonstrated that Mtb exhibited maximum intracellular cAMP at day3 post-inoculation when the optical density at 600 nm (OD600) of bacterial culture was 0.4. Subsequent growth on day 4 (OD600 of 0.70) resulted in sharp decline in the intracellular cAMP pool by ~3.5 folds and this level remained constant for next four days of growth when cultures reached to stationary phase (Fig. 1A). By ELISA, it was estimated that cAMP concentrations were 5.9 nmol/gm wet weight on day 3, and 1.6–1.8 nmol/gm wet weight on days 4–8 respectively (Fig. 1A).

**Expression analysis of Mtb ACs by real-time quantitative reverse-transcription PCR (qRT-PCR)**

Cellular concentration of cAMP can be regulated at the level of expression and/or activity of AC and the PDE, or by a change in the rate of cAMP export [6,32–34]. Although intracellular cAMP levels are significantly altered in Mtb, we observed that the extracellular cAMP pool remains constant over eight days of in vitro growth (data not shown). Since it is challenging to determine intracellular enzymatic activities of multiple ACs or PDEs, we focused on studying the expression profiles of mycobacterial ACs by qRT-PCR at various OD600 (Fig. 1B). Transcript levels of each of the 16 AC-encoding genes at designated OD600 were compared with their respective expression levels at day 1 post-inoculation in wild-type Mtb when the OD600 of culture was 0.1. Figure 1B shows that majority of AC-encoding genes except Rv1647, Rv1900c and Rv2212 were overexpressed by ≥5-fold when OD600 of culture reached to 0.34 on day 3. In contrast, transcripts of Rv1647 and Rv1900c were upregulated by 3.6 and 1.9-fold, respectively, whereas Rv2212 exhibited moderate reduction at OD600 of 0.34. Interestingly, at day 3 when OD600 of Mtb culture reached to 1.28, expression of most of the genes except Rv0386, cya, Rv1900c and Rv2488c was reduced by 2–12 folds compared to their respective expression levels at OD600 of 0.34 on day 3. Contrary to these, Rv0386, cya, Rv1900c and Rv2488c either maintained or displayed moderate increase in expression at OD600 of 1.28. Further growth of mycobacteria to OD600 of 3.5 at day 8 post-inoculation resulted in moderate decrease in expression of few genes that included Rv0386, Rv1538 and Rv2435c by 3.2–, 1.97- and 2.07-folds, respectively, while others were expressing at levels similar to OD600 1.28 (Fig. 1B). In contrast, Rv2212 transcript was not detected at OD600 of 3.5. These results are in line with the profile of cellular cAMP concentrations in Mtb which suggests that intracellular cAMP levels vary in proportion to the expression of ACs in Mtb (Fig. 1).

**Effect of various stresses on intracellular cAMP levels and expression of Mtb ACs**

Upon infection, virulent mycobacteria encounter stringent antimicrobial response within the host organism. However, tubercle bacilli are resistant to killing by host and persist for decades in the host tissues [35]. For their survival, mycobacterial pathogens sense and respond to exogenous stress conditions by modulating the expression of key genes. It is known that cAMP-associated transcription factors CRP and Cmr regulate the expression of several genes in Mtb which play important roles in Mtb–host interactions [18–24]. Interestingly, intracellular cAMP levels in mycobacteria increase dramatically after infection to macrophages [33]. These observations led us to hypothesize that Mtb must be equipped to modulate its cAMP pool under different stress conditions imposed by the host. Therefore we monitored the intracellular cAMP levels in Mtb after its exposure to different stress conditions that bacteria encounter during host infection.

Wild-type Mtb cells were exposed to nutrient starvation (PBS), glucose deficiency, glycerol deficiency, oxidative stress (cumene hydroperoxide, CHP), nitosative stress (DETA/NO), acid stress (pH 4.5) and heat stress (42°C), as described earlier [36]. Subsequently, cells were lysed and cAMP was measured in the lysates by ELISA. As shown in figure 2A, a moderate decrease in intracellular cAMP levels was observed after nutrient starvation and oxidative stress, whereas deficiency of any of the two carbon sources resulted in ~2-fold reduction in the intracellular cAMP concentration (Fig. 2A). In contrast, a mild increase in intracellular cAMP level was seen in the cells after nitrosative and acid stresses. Interestingly, these stresses, maximum effect was observed after heat-stress, which caused >2-fold increase in intracellular cAMP level under these culture conditions.

In order to assess whether increase in cAMP level after the heat-stress is related to the expression of ACs, we performed the qRT-PCR to measure mRNA levels of AC-encoding genes in Mtb following the heat-shock treatment. Expression level of individual gene was obtained after normalization with the level of a housekeeping gene sigA, which remained unaltered under these conditions. Figure 2B depicts that the mRNA levels of 5 out of 16 AC-encoding genes (Rv1647, Rv2212, cya, Rv2488c and Rv0386) are increased by >2-fold after heat-stress (p ≤0.05). Taken together, these observations indicate that Mtb responds to heat-shock exposure by inducing the expression of certain ACs which results in elevated cAMP-levels in the cell.

**Identification of cAMP-regulon by microarray**

To identify cAMP-regulated genes of Mtb, wild-type Mtb CDC1551 cultures were treated with di-butyl cAMP (dB-cAMP)
and the expression of genes was compared with butyric acid-treated control samples by whole genome microarrays as described in materials and methods. Before proceeding to microarray experiments, intracellular cAMP levels were measured in both the samples by ELISA, which indicated a 2-fold increase in cAMP levels after 2 hrs of incubation with 20 mM db-cAMP (Fig. 3A). In contrast, there was no change in intracellular cAMP levels after incubation with 20 mM butyric acid for 2 hrs, which suggests that db-cAMP specifically elevates intracellular cAMP pool by 2-fold (Fig. 3A). A total of three hybridization experiments were performed with three independent RNA preparations, and the results were analyzed statistically. Expression level of individual gene was measured after normalization with the level of sigA, which was not changed under these conditions. Genes that were differentially expressed by \( \geq 2.0 \)-fold \( (p \leq 0.05) \) in db-cAMP-treated samples are shown in table 1. The microarray results demonstrated that a total of 7 genes were upregulated and 5 genes were downregulated by \( \geq 2 \)-fold after treatment with db-cAMP, in comparison to butyric acid-treated samples (Table 1). The most abundant transcripts were those of a subset of heat stress-induced genes comprising of Rv2025c, dnaK, grpE and dnaJ. Further, differences in expression of these genes between db-cAMP-treated and butyric acid-treated samples by microarray were 9.8\( \pm 4.4 \), 3.7\( \pm 0.6 \), 2.9\( \pm 0.9 \), and 3.2\( \pm 0.7 \) respectively (Table 1). These differences were more prominent in qRT-PCR which exhibited 60.9\( \pm 3.2 \), 8.9\( \pm 0.4 \), 9.1\( \pm 0.1 \), and 7.3\( \pm 0.6 \)-fold induction of Rv2025c, dnaK, grpE and dnaJ respectively after db-cAMP treatment (Fig. 3B). Moreover, expression levels of these genes were significantly higher than of cAMP-regulated \( \text{whiB1} \) [20], which exhibited 1.4\( \pm 0.18 \)-fold induction under these conditions. Other genes that exhibited notable induction in expression levels were Rv0264c (2.9\( \pm 1.6 \)-fold), Rv1057 (2.3\( \pm 0.2 \)-fold), and Rv1350c.
pncB1, 2±1-fold). In contrast, five genes that include Rv0146, mmpS5, fadD9, hupB, and Rv3830c were significantly downregulated by 5.9, 2.1, 5.1, 2.0 and 8.5-folds respectively, in db-cAMP-treated Mtb samples (Table 1).

Next, we examined whether expression of these genes is altered in mycobacteria due to direct effect of cAMP on their promoter activity. A 450bp long DNA sequence corresponding to 5'-UTR of dnaK was cloned upstream to the lacZ in promoter probe plasmid pSD5B [20] and the recombinant plasmid pSD5B-dnaK was transformed into Msm. Expression of lacZ was subsequently monitored in the presence or absence of db-cAMP in Msm::pSD5B-dnaK. As shown in figure 3C, in the absence of db-cAMP (+cAMP), Msm::pSD5B-dnaK exhibited significant β-gal activity in comparison to Msm containing empty plasmid. Interestingly, addition of db-cAMP to bacterial cultures (+cAMP) resulted in ~40-fold increase in activity of dnaK promoter, whereas there was no effect of db-cAMP on the basal expression levels of lacZ in empty plasmid containing control strain (Fig. 3C).

These results thus clearly demonstrate that cAMP acts as a regulator of the expression of multiple genes including a subset of heat stress-induced genes of Mtb.

Identification of CRPM-recognition sequence in 5'-UTR of cAMP-regulon

The CRP of E. coli binds to a 16bp sequence TGTGA-N6-TCACA in the 5'-UTR of cAMP-regulon. Similar to E. coli the CRP of Mtb (CRPM) was also shown to regulate expression of key Mtb genes that bear the similar sequences in their 5'-UTR [18–20,22,23]. In order to understand whether cAMP-mediated expression of genes in our study is due to direct binding of CRPM to their respective 5'-UTRs, we analyzed 350bp sequences upstream to the start codon of each of the 10 open reading frames (ORFs) by using the TubercuList database (http://genolist.pasteur.fr/TubercuList/) to identify the putative CRPM consensus sequences bearing two mismatches in either of the left (TGTGA) or right (TCACA) arms. Since dnaK is transcribed in operon with

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Figure 2. Effect of various stresses on intracellular cAMP levels and expression of ACs in Mtb. A) Estimation of cAMP in Mtb during in vitro growth under different stresses. Wild-type culture suspension in Middlebrook 7H9 broth at OD600 of 0.4 was exposed to different stresses as described earlier [36] and intracellular cAMP levels were determined by ELISA. Bar graph represents cAMP level in each of the stressed samples as percentage of un-treated control. Data are the averages of three independent experiments and the mean values ± standard deviations are shown. B) Expression analysis of the AC-encoding genes following thermal stress in Mtb. The fold-expression, as measured by qRT-PCR, indicates the ratio of sigA-normalized gene expression levels in Mtb exposed to 42 °C relative to those in the Mtb grown at 37 °C. The graph represents an average of three experiments. Error bars indicate standard deviations and asterisk denotes p ≤ 0.05.

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and dnaJ [37], we omitted the upstream sequences to grpE and dnaJ in our analysis. As shown in table 2 except hupB, all other ORFs exhibited putative CRPM-recognition sequences in their 5'-UTR. These observations suggest that cAMP may regulate the expression of these genes by facilitating the binding of cAMP-activated CRPM to their promoter region.

EMSA to validate direct binding of CRPM to 5'-UTR of dnaK operon

Analysis of 5'-UTR of dnaK operon indicates the presence of two putative CRPM-consensus sequences. The first sequence CRP-1 (5'- CGTTAGCATGCTCAGT-3') is located between positions -155 and -120 from translation start codon, whereas the second sequence CRP-2 (5'- AGCGACCGTCA GCACG-3') is situated further upstream between positions –301 and –286 from translation start codon (Fig. 4A). A careful observation demonstrates that CRP-2 is located 127bp upstream to SigH-binding site which is positioned between –158 to –130 from translation stop codon (5'- GGGACAAAGACCAGCAGCCAGACGGTTA-3'), whereas CRP-1 overlaps with SigH-recognition sequence in 5'-UTR of dnaK (Fig. 4A). Since dnaK, grpE and dnaJ were amongst the most abundant transcripts in db-cAMP-treated Mtb, we sought to determine if their expression is controlled by direct binding of CRPM to the promoter sequence in the 5'-UTR. CRPM was purified as earlier [20] and subjected to binding with DNA fragment comprising of sequence between –1 to –346 from translation start codon of dnaK (dnaK-346). As shown in figure 4B, CRPM makes a specific complex with this sequence. Further deletion of 43bp from 5' end (dnaK-303) indicated that CRPM continues to bind this sequence, albeit with ~2-fold reduced affinity as assessed by the intensity of CRPM-DNA complex. On the other hand deletion of additional 22bp (dnaK-281) which resulted in complete loss of CRP-2 site located between –301 and –286, completely abolished the complex formation (Fig. 4B). These results thus establish that: i) regulation of dnaK expression is governed by direct binding of CRPM to its 5'-UTR, ii) CRPM binds at CRP-2 site and not at CRP-1 site in the dnaK promoter, and iii) binding of CRPM to 5'-UTR of dnaK and its subsequent effect on mRNA expression is independent of SigH binding.

Presence of cAMP enhances binding of CRPM to corresponding promoter DNA sequences [20,23]. Hence, we analyzed if association of CRPM to dnaK promoter is also influenced by cAMP. Binding reactions were carried out in the absence (apo-CRPM) or the presence of 1 mM cAMP, and the CRPM-DNA complexes were resolved on polyacrylamide gel as described in materials and methods. Figure 4C shows that presence of cAMP enhances the complex formation between CRPM and dnaK-346 promoter fragment by 3-fold compared to that with apo-CRPM. These results thus clearly indicate that cAMP regulates the...
Role of cAMP in Mycobacteria

Microbial pathogens including Mtb are adapted to survive under diverse stress conditions. Effects of cAMP on bacterial responses to cold-shock [44], expression of stress proteins [45] and RpoS, a late stationary sigma factor which also regulates oxidative stress response [46,47] indicate that cAMP is an important determinant of the bacterial response to variety of stresses other than carbon metabolism in many pathogenic bacteria. Recently it was shown that cAMP regulates the response of uropathogenic E. coli to nitrosative, oxidative and acid stresses, and affects its virulence [48]. In contrast to these organisms, intracellular cAMP in Mtb was significantly stimulated after the heat stress whereas other stresses such as oxidative, nitrosative or acid exhibited a milder effect (Fig. 2A). Response to heat stress is an adaptive response to a sudden increase in ambient temperature, which involves a group of proteins such as chaperons, proteases, and regulatory factors commonly known as heat shock proteins [37]. Transcriptionally heat shock proteins are expressed in Mtb by involving an extracellular function sigma protein, SigH [49].

Upregulated expression of 5 ACs after the heat stress and presence of SigH-consensus sequences in the upstream promoter region of their respective ORFs (Fig. 2B and Table 3) together suggest that apparent induction of cAMP pool under these conditions may be due to differential transcription of mycobacterial AC-encoding genes by SigH. Parallel to these observations, induction of genes encoding primary heat shock proteins DnaK, GrpE, DnaJ and Rv2025c by exogenous cAMP indicates that a subset of heat shock proteins are expressed in Mtb under the effect of cAMP (Table 1 and Fig. 3B-C). Absence of CRP/Cmr-recognition sequences in sigH promoter and lack of its induction by cAMP together suggest that cAMP-driven expression of heat shock genes may not be directly controlled by SigH. Alternatively our results propose that cAMP regulates the expression of these genes by facilitating the binding of CRP\(^{\text{M}}\) to their promoter.

Discussion

This study was designed to estimate the intracellular cAMP levels over a period of \textit{in vitro} growth and its role in gene expression of Mtb. Cyclic AMP in bacteria was first reported half a century ago when the cyclic nucleotide was observed in the culture filtrate of \textit{Brucella abortus} [38] and later in \textit{E. coli} [39]. Subsequently occurrence of cAMP was reported in other bacteria including mycobacteria [3]. Later after discovery of cAMP in bacteria, it was studied that cAMP plays an important role in assimilation of sugar molecules by regulating a process called catabolite repression [40,41]. Levels of cAMP keep changing as a function of glucose level in the cell [41]. Although in mycobacteria cAMP is not involved in carbon metabolism, variation in intracellular cAMP levels over a period of \textit{in vitro} growth (Fig. 1A) indicates that cAMP levels are dynamic. A constant extracellular cAMP concentration rules out the prospect of differential export of cAMP to outside culture medium. Although effects of differential activities of ACs, PDEs and other associated factors on intracellular cAMP concentrations is not ruled out, a similar pattern of expression of 16 ACs and intracellular cAMP levels during \textit{in vitro} growth of Mtb suggests that the relative abundance of ACs could be an important factor governing the intracellular cAMP levels in Mtb (Fig. 1A and B). While, the trigger(s) of sudden changes in expression of ACs and subsequent cellular cAMP levels at late growth stages of Mtb is yet to identify, the intracellular polyphosphates may be an important determinant of cellular cAMP levels [42]. Polyphosphates, generated by the activity of an enzyme known as polyphosphate kinase, is highly accumulated at the stationary phase as well as under different stress conditions in Mtb [43], and are known to inhibit the activity of ACs [42]. These information warrant further studies to analyze the effect of polyphosphates on intracellular cAMP levels.

Table 1. Differentially expressed genes in Mtb treated with db-cAMP relative to butyric acid treated bacteria.

| Gene product\(^{a}\) | Locus | Change in expression (fold) | SD (fold) | \(P\) value |
|---------------------|-------|-----------------------------|-----------|------------|
| CHP                 | Rv0146| -5.9                        | 0.0       | 0.01       |
| CHP                 | Rv0264c| 2.9                         | 1.6       | 0.02       |
| DnaK                | Rv0350| 3.7                         | 0.6       | 0.02       |
| GrpE                | Rv0351| 2.9                         | 0.9       | 0.05       |
| DnaJ                | Rv0352| 3.2                         | 0.7       | 0.03       |
| MmpS5               | Rv0677c| -2.1                        | 0.2       | 0.00       |
| CHP                 | Rv1057| 2.3                         | 0.2       | 0.04       |
| CHP                 | Rv1330c| 2.0                         | 1.0       | 0.03       |
| CMP                 | Rv2025c| 9.8                         | 4.4       | 0.02       |
| FadD9               | Rv2590| -5.1                        | 0.1       | 0.04       |
| IupB                | Rv2986c| -2.0                        | 0.0       | 0.02       |
| TRP                 | Rv3830c| -8.5                        | 0.0       | 0.03       |

Expression of 3924 genes of Mtb was compared between the cells treated with db-cAMP and butyric acid. An average of data from three independent experiments (\(p\leq0.05\)) is included in the table. Genes that exhibit an \(\geq2\) fold difference in expression levels by whole-genome microarray are shown. SD: standard deviation. \(^{a}\)CHP, CMP and TRP denote conserved hypothetical protein, conserved membrane protein and transcription regulatory protein, respectively.

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expression of \textit{dualk} operon by facilitating the binding of CRP\(^{M}\) to its promoter.

Materials and Methods

Bacterial strains, culture conditions and plasmid

For culturing of Mtb CDC1551, Middlebrook 7H9 broth supplemented with 1x OADC, 0.02% tween-80 and 0.5% glycerol, and Middlebrook 7H10 agar supplemented with 1x OADC and 0.5% glycerol were used. Msm mc\(^{\text{L}}\)155 was cultured in 7H9 broth containing 0.02% tween-80 and 0.5% glycerol. Bacteria were grown at 37°C with (in 7H9 broth) or without (on 7H10-agar) shaking at 200 rotations per minute. For culturing in glucose-free medium, synthetic supplement containing oleic acid, albumin and saline equivalent to 1x concentration were added to
Table 2. Analysis of putative CRP-recognition sequences in 5’-UTRs of cAMP-regulon of Mtb.

| Locus (Rv/ annotation) | Sequence* | Position from Start codon |
|------------------------|-----------|---------------------------|
| Rv0146                 | tgtcagggctttacc | −325 |
| Rv0146                 | ttgcacatgaacc | −316 |
| Rv0146                 | tpagccgcttcgacg | −45 |
| Rv0146                 | agpcccgttggtaga | −266 |
| Rv0146                 | ttcgccggtaccgac | −244 |
| Rv0146                 | cgacgaccgctccgacc | −103 |
| Rv0264c                | tgcgtatgatgaccg | −300 |
| Rv0350/dnaK            | tgcgtatgatgaccg | −301 |
| Rv0350/dnaK            | ccctctagaggtgaatttgg | −135 |
| Rv0677c/mmpS5          | ttcactgacttggga | −81 |
| Rv0677c/mmpS5          | tgtcgcagacttcaca | −149 |
| Rv0677c/mmpS5          | tctgacacttgtagcac | −70 |
| Rv1057                 | cgccagctggtaacc | −248 |
| Rv1057                 | tccagatttgctgctgtc | −208 |
| Rv1330c                | tggcagcccgtgccgaccg | −21 |
| Rv1330c                | agtcatagctcacttgcag | −233 |
| Rv1330c                | tgcgcagccgttgcgctgtc | −276 |
| Rv2025c                | tgcgccctggtgctgtc | −246 |
| Rv2025c                | tgcctgctgctgctgct | −82 |
| Rv2590/fadD9           | tgcctgctgctgctgcttgc | −206 |
| Rv2590/fadD9           | cggcagctggtaacc | −285 |
| Rv2590/fadD9           | cggcagctggtaacc | −133 |
| Rv3830c                | tgcgccgctgtgccgctgtc | −72 |
| Rv3830c                | tgcgccgctgtgccgctgtc | −42 |
| Rv3830c                | ttttcgccgctgtgccgctgtc | −44 |

Three hundred fifty base pair long DNA sequences upstream to the respective ORFs were analyzed to identify CRP<sup>+</sup>-recognition sequence. *The underlined sequences represent putative CRP<sup>+</sup>-recognition sequences (TGTTGA-N<sub>6</sub>-TCACA, with two mismatches in any of the arms) in the 5’-UTR of the respective genes.

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7H9 medium containing 0.02% tween-80 and 0.5% glycerol. Bacteria were treated with different stress agents as described earlier [36]. The promoter probe vector pSD5B [20] was kindly provided by Dr. Anil Tyagi, University of Delhi South Campus, New Delhi, India. To determine the in vivo activity of dnaK promoter in mycobacteria, the 450bp region upstream to translational start site of dnaK ORF was amplified using primer pairs dnaK-450F (5’-GGGTCTAGAGCACCGTTGGCCCGTGTTGATG-3’) and dnaK-1R (5’-CCCTCTAGAGGGATCCCTCTAGAATCTCATGTAGTGAC-3’), and cloned at XbaI site in pSD5B. The pSD5B-dnaK plasmid harboring dnaK promoter in sense orientation was subsequently electroporated in Msm and the recombinant Msm::pSD5B-dnaK strains were selected on Middlebrook 7H10 agar containing 0.5% glycerol and 25 mg/L kanamycin.

Estimation of cAMP by ELISA

Intracellular cAMP determination in Mtb was performed with clarified cell lysate after heat lysis of cell pellets in 0.1 M HCL. Cyclic AMP was estimated by ELISAs using Enzyme Immunoassay Kits (Assay Designs Inc., Ann Arbor, MI) according to the manufacturer’s instructions. The intracellular cAMP levels in Mtb were estimated as nmol/gm wet weight.

RNA extraction

Total RNA was isolated from Mtb by using the TRIZol suspension according to manufacturer’s instructions (Invitrogen Corporation, Carlsbad, CA).

Real-time quantitative reverse transcription PCR

RNA isolated from the bacterial cultures was subjected to treatment with RNase-free DNase I (Ambion) to remove traces of contaminating DNA. Subsequently, absence of DNA in the RNA preparations was verified by 30 cycles of PCR followed by ethidium-bromide-stained agarose gel analysis before proceeding with reverse transcription of the RNAs. Complementary DNA (cDNA) synthesized from total RNA was subjected to real-time quantitative reverse transcription PCR typically as described earlier [36].

Microarray analysis

To analyze the effect of cAMP on the expression of mycobacterial genes, Mtb was cultured to OD<sub>600</sub> of 0.6 and pelleted down by centrifugation at 6000 x g for 10min at 4°C. Culture pellets were washed twice with 1 x phosphate buffered saline, pH 7.4 (PBS) and suspended in 1/10<sup>th</sup> volume of 7H9 medium containing either of the 20 mM butyric acid (control) or di-butyryl cAMP (db-cAMP, test). After 2 hrs of incubation at 37°C in roller bottles, cells were pelleted down, washed 3 x with PBS and RNA was isolated as described above. For probe preparation, cDNA was synthesized from 3 µg of total RNA from the test and control strains of Mtb and labeled with Cy3 or Cy5 (GE Healthcare). Microarray slides were prepared by using 70-mer oligos encompassing the entire genome of Mtb (GE accession number GSE54289) and hybridization of labeled cDNAs was performed typically as described earlier [36]. RNA samples were prepared from 3 biological replicates. Slide scanning and data analysis was performed as previously described [36].

β-Galactosidase assay

Promoter activity of dnaK was measured in Msm::pSD5B-dnaK by fluorescent based detection of lacZ expression, as described earlier [51]. Briefly, Msm::pSD5B-dnaK cultures were grown in 7H9 broth medium containing kanamycin (25 mg/L) to an OD<sub>600</sub> of 1.0, washed twice with 7H9 medium and resuspended in 7H9 medium. The OD<sub>600</sub> of culture suspension was adjusted to 1.0, and the cultures were incubated with either 10 mM db-cAMP or 10 mM butyric acid for 2 hrs. Hundred microliter of culture suspensions were taken into 3 separate wells of a 96-well black fluoroplate (Greiner Bio-One). Fluorescent β-gal substrate, 5- acetylamino-FDG (C2FDG) (Life Technologies) was subsequently added to a final concentration of 33 µM and incubation using a spectrofluorometer (Biotek) with an excitation of 485±20 nm and emission of 528±20 nm. A similar experiment was performed with Msm containing empty plasmid, pSD5B as control.

EMSA

DNA probes for EMSA analysis were synthesized by PCR amplification of the desired regions of the dnaK promoter, using biotin-labeled oligonucleotides as the primers (Fig. 4A). The amplicons were purified from agarose gels and used for gel-shift.
Figure 4. Identification of CRP-binding motif in promoter region of dnaK operon by EMSA. A) Sequence of the 5'-UTR of dnaK operon. The putative ribosome-binding site (RBS), and the translation start site (TSS) are boldfaced and underlined. The putative CRP-binding sites are shaded, whereas the SigH-recognition sequence is shown in italics. Positions and directions of the primers that were used for PCR amplification of DNA fragments shown in (B) are marked by horizontal arrows. B) Interaction of CRPM with the 5'-UTR of dnaK operon. Biotin-labeled dnaK promoter fragments of various lengths, as shown, were incubated with CRPM in the presence of 1 mM cAMP before separation of DNA-protein complexes by gel electrophoresis. Reaction mixtures containing promoter fragments but lacking the CRPM were resolved in adjacent lanes as controls. C) Effect of cAMP on complex formation between CRPM and dnaK promoter. Purified CRPM pre-incubated with or without 1 mM cAMP for 30 min on ice, was subject to binding with full length dnaK promoter fragment (dnaK-346) for 15 min at 37°C before separation of CRPM–dnaK complexes by gel electrophoresis. Complex, CRPM–dnaK complex; Free, unbound DNA fragment.

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Table 3. Identification of putative SigH-recognition sequences in the 5'-UTRs of AC-encoding genes of Mtb.

| Locus (Rv/annotation) | Sequence*  | Position from Start codon |
|-----------------------|------------|---------------------------|
| Rv0386                | cggaaatccacgcgtccgraytgcggtgcggttc | −98 |
| Rv1625c/cya            | cggacaatctcgcagctgtgcgtgcggttc | −335 |
| Rv1647                | cggacaatctcgcagctgtgcgtgcggttc | −311 |
| Rv2212                | cggacaatctcgcagctgtgcgtgcggttc | −114 |
| Rv2488c               | cggacaatctcgcagctgtgcgtgcggttc | −152 |

Three hundred fifty base pair long DNA sequences upstream to the AC-encoding genes were analyzed to identify SigH-recognition sequences. *The underlined sequences represent putative SigH-recognition sequences (c/gGGAAc-N17–21-c/gGTTc/g) in the 5'-UTR of the respective genes.

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The EMSA experiments were performed typically as described earlier [20].

Author Contributions
Conceived and designed the experiments: NA. Performed the experiments: EC NA. Analyzed the data: EC NA WB. Contributed reagents/materials/analysis tools: NA WB. Wrote the paper: EC NA WB.

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