Role of intragenic binding of cAMP responsive protein (CRP) in regulation of the succinate dehydrogenase genes Rv0249c-Rv0247c in TB complex mycobacteria

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ABSTRACT

Bacterial pathogens adapt to changing environments within their hosts, and the signaling molecule adenosine 3’, 5’-cyclic monophosphate (cAMP) facilitates this process. In this study, we characterized in vivo DNA binding and gene regulation by the cAMP-responsive protein CRP in M. bovis BCG as a model for tuberculosis (TB)-complex bacteria. Chromatin immunoprecipitation followed by deep-sequencing (ChIP-seq) showed that CRP associates with ~900 DNA binding regions, most of which occur within genes. The most highly enriched binding region was upstream of a putative copper transporter gene (ctpB), and crp-deleted bacteria showed increased sensitivity to copper toxicity. Detailed mutational analysis of four CRP binding sites upstream of the virulence-associated Rv0249c-Rv0247c succinate dehydrogenase genes demonstrated that CRP directly regulates Rv0249c-Rv0247c expression from two promoters, one of which requires sequences intragenic to Rv0250c for maximum expression. The high percentage of intragenic CRP binding sites and our demonstration that these intragenic DNA sequences significantly contribute to biologically relevant gene expression greatly expand the genome space that must be considered for gene regulatory analyses in mycobacteria. These findings also have practical implications for an important bacterial pathogen in which identification of mutations that affect expression of drug target-related genes is widely used for rapid drug resistance screening.

INTRODUCTION

Tuberculosis (TB), caused by Mycobacterium tuberculosis (Mtb), is an ancient disease that continues to cause significant morbidity and mortality worldwide. Increasing levels of drug resistance and a complex synergy with human immunodeficiency virus (HIV) complicate efforts to control this deadly pathogen (1). There is an urgent need for new therapeutics against Mtb, and development of effective new drugs requires better understanding of Mtb physiology.

Bacterial pathogens must adapt to changing environments within the host during infection, and they often use cyclic nucleotides as ‘second messengers’ to sense and respond to their external environments (2,3). Adenosine 3’, 5’-cyclic monophosphate (cAMP) is one such signaling molecule that is widely used by both microbial pathogens and their mammalian hosts (4–8). Mtb is a particularly unusual microbe in that it has ~15 biochemically distinct adenylyl cyclases (AC), which generate cAMP and allow Mtb to respond to multiple environmental cues (9,10). Mtb also encodes 10 putative cyclic nucleotide monophosphate (cNMP) binding proteins, of which three have been characterized. Rv0998 (Mt-PatA) is a cAMP-activated protein lysine acetylase that has several biological targets in mycobacteria, including Mtb acetyl-CoA synthase (11,12). Rv1675c (called Cmr for cAMP and macrophage regulator) and Rv3676 (named CRP for cAMP-responsive protein) both contain helix turn helix domains and belong to the CRP/FNR family of transcription factors (13–16).
CRP is important for Mtb pathogenesis, as Mtb mutants deleted for cpr show impaired growth in murine macrophages (15,17) and in a mouse model of TB (15). The CRP ortholog in M. bovis BCG Pasteur differs from Mtb CRP by two amino acids (L47P and E178K), and has approximately 2-fold higher affinity for DNA in vitro compared to CRP from Mtb (13). However, virulence of Mtb H37Rv cpr deletion mutants can be fully restored by expression of cpr from either BCG or Mtb, confirming that both orthologs function similarly as virulence-associated transcriptional regulators in vivo (18). For simplicity, we use CRP throughout the text to refer to protein from either Mtb or BCG except when referring to a species-specific CRP behavior.

The molecular basis of CRP’s role in virulence is not known, but it directly regulates expression of several biologically important genes. For example, CRP controls expression of rpfA, which encodes a resuscitation-promoting factor thought to play a role in the reactivation of dormant Mtb cultures (19). CRP also upregulates expression of serC, which encodes a phosphoserine aminotransferase (14,17,20), and growth of Mtb cpr mutants is slowed by a resulting defect in serine biosynthesis (17). Correction of this deficiency by serine supplementation or by constitutive expression of serC restores normal growth levels of Mtb in culture media but not within macrophages (17). More recently, CRP was shown to directly regulate expression of whiB1 (21,22), which encodes an essential transcription factor that contains a nitric oxide (NO) sensing [4Fe-4S]2+ cluster (23,24). DNaseI-footprinting showed CRP’s ability to bind cooperatively to each of two CRP sites within the whiB1 promoter region, with slightly enhanced binding in the presence of cAMP (21).

A combination of in silico and experimental studies has defined CRP as a global regulator within Mtb, but the full extent of this regulation is not known. Whole genome expression microarrays of Mtb H37Rv demonstrated potential regulation of 16 genes from 13 individual promoters, including those of iprQ, rpfA, alpC, lipQ, fadD26 and whiB1 (15). Another study (14) combined BCG DNA sequences recovered by affinity capture with previously characterized E. coli CRP binding sites to seed a computational analysis of the potential CRP regulon in TB complex bacteria. This affinity capture study predicted 114 CRP_Mt regulon members, based on conserved binding motifs, corresponding to 73 promoter regions in Mtb (14). A subsequent in silico study (25) used putative promoter sequences from the regulon of C. glutamicum GltRx, an ortholog of CRP, as seed sequences to predict 135 CRP binding sites with the potential to regulate expression of 207 genes within 121 transcriptional units.

Surprisingly little overlap was found among the regulons predicted from these prior studies, despite their identification of similar binding motifs. In this study, we characterize in vivo CRP-DNA binding and gene regulation in M. bovis BCG as a model system for TB complex bacteria. We found using Chromatin Immunoprecipitation followed by deep-sequencing (ChIP-seq) that CRP is associated with ~900 DNA binding regions in M. bovis BCG, only ~14% of which occur within linear or divergent intergenic DNA sequences. The remaining CRP binding regions were found within intragenic regions (83%) or between convergent intergenic sequences (3%). Blind deconvolution (26) of the CRP binding profile within these enriched regions revealed four CRP binding sites at the Rv0250c-Rv0249c succinate dehydrogenase locus, including one within the Rv0250c open reading frame (ORF). Deletion analyses demonstrated that CRP binding contributed to regulation of Rv0249c-Rv0247c expression from each of two promoters. An Rv0249c proximal promoter required upstream Rv0250c intragenic sequences for maximum expression, while the second promoter is located upstream of Rv0250c. These findings show that CRP regulates a critical metabolic step in central metabolism and that intragenic binding sites can significantly affect regulation of biologically important gene expression in TB complex bacteria. The extremely large number of binding regions further suggests that CRP may function as a nucleoid associated protein (NAP) in addition to its established role as a canonical transcription factor in TB complex mycobacteria.

MATERIALS AND METHODS

Bacterial strains and culture

M. tuberculosis H37Rv (ATCC 25618) and M. bovis BCG (Pasteur strain, Trudeau Institute) was grown in mycomedia: Middlebrook 7H9 medium supplemented with 0.2% glycerol, 10% oleic acid-albumin-dextrose-catalase (OADC), 0.05% Tween-80 or on Middlebrook 7H10 (Difco) supplemented with 0.5% glycerol, 10% OADC and 0.01% cycloheximide. Where indicated, heat-killed M. tuberculosis H37Rv (ATCC 25618) genomic DNA was used. Fresh cultures were inoculated from frozen seed stocks for every experiment and transferred to the desired culture condition, as previously described (27). Cultures for ChIP-sequencing were grown from seed stocks in mycomedia for seven days to late log phase in shallow (2 mL) cultures 1.3% O2 + 5% CO2 in 225 cm2 flasks with gentle rocking as described (28). On day 3, dibutyryl-cAMP (dbcAMP) was added to one culture flask for a final concentration of 10mM and returned to the incubator. For starvation experiments, cultures were grown in ambient air from seed stocks in mycomedia for 10 days, pelleted and washed with Dulbecco’s phosphate buffered saline (PBS) or fresh mycomedia and returned to the incubator for 24 h. Transformations into M. bovis BCG were done by electroporation (29). E. coli was grown in Luria Broth or on Luria Broth agar plates. Kanamycin was used at 25 µg/ml as needed. All cultures were grown at 37°C.

Chromatin Immunoprecipitation and sequencing

50 ml culture volumes of M. bovis BCG Pasteur were grown to late-log phase under 1.3% O2 + 5% CO2 and cross-linked with 1% formaldehyde at room temperature for 30 min with gentle rocking. Crosslinking was quenched with glycin (250 mM final concentration) for 15 min at room temperature. The cells were harvested, washed twice with ice-cold PBS, and resuspended in 0.5–0.6 ml of Buffer I (20 mM HEPES pH 7.9, 50 mM KCl, 0.5 M DTT, 10% glycerol) with protease inhibitor cocktail (Sigma). Bacterial cells were lysed and DNA sheared by sonication for 25
min with a Covaris S2 (Covaris, Inc., Woburn, MA). The salt concentration of the cleared cell lysate was adjusted to a final concentration of 10mM Tris-HCl pH 8.0, 150 mM NaCl, 0.1% NP-40 (IPPI50 buffer). Immunoprecipitation was carried out by incubation of lysate with 10 μl of CRP antisera 4°C. 50 μl of Protein A agarose beads were rinsed with IPPI50 buffer, and added to the lysate-antiserum mixture. Protein A agarose-lysate-antisemur mixture was incubated at 4°C for 30 min and room temperature for 1.5 h. The beads were washed at least five times with IPPI50 buffer followed by two washes with TE buffer (10 mM Tris HCl pH 8.0, 1 mM EDTA). DNA from agarose beads was eluted by incubation with 150 μl elution buffer (50 mM Tris HCl pH 8.0, 10 mM EDTA, 1% SDS) at 65°C for 15 min. A second elution was carried out by incubation of pellet with 100 μl of TE + 1% SDS at 65°C for 5 min. Both elutions were pooled and treated with 1 mg/ml Proteinase K at 37°C for 2 h and 65°C overnight. DNA was purified using PCR purification kit (QIAGEN).

Sequencing was performed as described previously (30). Briefly, sequencing was performed on the Illumina platform, using a GAIIx (Boston University, sequencing core). Coverage along the genome was calculated using Bowtie2 (31) and SamTools (32). Enriched regions were called using log-normal distributions as previously described (30). The minimum region length was at least 150 nt long and had a minimum 60 nt shift between its forward and reverse peaks. Region coverage was normalized using mean coverage of an experiment, correcting for the number of reads amongst experiments. Exact binding sites were determined as described by Gomes et al. (26), using the BRACIL blind deconvolution method. BRACIL integrates ChIP-seq coverage and genome sequence to jointly identify binding sites with single nucleotide resolution and a corresponding consensus binding site motif. Regions from the BCG genome were mapped to the Mtb H37Rv genome by sequence similarity using BLAST.

**cAMP treatments**

Levels of cAMP within bacteria were increased exogenously or endogenously for some experiments. For the exogenous method, dibutyryl-cAMP (dbcAMP) was added to the culture media for four days at a final concentration of 10mM. dbcAMP is a cell permeable molecule that is cleaved to produce cAMP upon internalization (16). Alternatively, excess cAMP was endogenously produced by expression of the adenyl cyclase Rv1264 catalytic domain (Rv1264Cat) under the strong, constitutively active promoter of Rv0805 (16). This construct is cloned on the pcRII.oriM, a multicopy plasmid that contains the oriM origin of replication from pAL5000, which was shown to have a copy number of 3–10 (33). This approach was previously shown to elevate intracellular cAMP levels within bacteria by up to 40X (16).

**Functional analysis of genes**

Regulatory targets of binding sites were assigned based upon their locations relative to adjacent genes. Immediate downstream genes were assigned as targets for intra- and intergenic binding sites. Therefore, linear intergenic sites were assigned only one target while divergent intergenic sites were assigned two. Intragenic binding sites were also assigned their overlapping gene as a target, yielding a possible total of up to three targets. Non-coding RNA genes currently listed in the Tuberculist annotations were treated similarly to protein coding genes with respect to their assignment as potential regulatory targets of proximal binding sites.

We next used Tuberculist functional annotations (http://tuberculist.epfl.ch/) (34) to assign the functional classification to each target gene within the CRP ChIP-seq data set. The percentage of potentially regulated target genes was calculated for each functional classification and segregated according to the location of the binding site to compare target distributions for intragenic binding sites, intergenic binding sites and all ChIP-seq sites combined. The expectation of functional classification for the Mtb H37Rv genome was also calculated. Hypergeometric calculations with a Bonferroni correction of 11 were performed using Excel to determine the significance of differences between the genome and the type of ChIP-seq site. P < 0.05 was considered significant.

**Electrophoretic mobility shift assays (EMSA)**

DNA probes were generated by PCR using DNA primers based on the Mtb H37Rv sequences denoted in Tuberculist (34) with the addition of BamHI restriction sites for downstream cloning as needed. The PCR forward primer was labeled with [γ-32P]-ATP using T4 DNA polynucleotide kinase (New England Biolabs). DNA fragments were then labeled by PCR (30 cycles) using Mtb genomic DNA as a template, diluted 1:3 and 1 μl DNA probe was used in each 10 μl binding reaction. Samples were electrophoresed on an 8% (29:1) non-denaturing polyacrylamide gel for 2.5 h with a constant voltage of 150 v. Gels were vacuum dried, exposed on a phosphor screen, scanned with a Storm 860 PhosphorImager (Molecular Dynamics), and analyzed with ImageQuant software (Molecular Dynamics).

**Growth in copper**

Wild-type BCG, BCG Δcrp and BCG Δcrp::pMBC1029, a strain with a single-copy, integrative vector containing the crp gene with its native promoter (17), were grown in 25 cm² tissue culture flasks at 37°C in mycomedia with the indicated CuSO₄ concentrations for 14 days under ambient air conditions or in hypoxia supplemented with 5% CO₂. After testing a range of concentrations, based on results of prior studies (35,36), we chose 100 μM as our standard CuSO₄ assay concentration for media that included OADC, and 50 μM for media that lacked OADC. At the time points indicated, aliquots were removed, cells were cultured in a cup horn sonicator (Virtis Virsonic, setting 4, 10 s total time for 5 s on and 5 s off), and OD₆₂₀ was determined using a Sunrise plate reader (Tecan).

**Construction of lacZ fusions**

Lists of plasmids and primers used in this study are in Supplementary Figures S1 and S2, respectively. Promot-
ers of interest were amplified using primers that contained either BamHI or ScaI (due to an internal BamHI restriction site within Rv0250c) restriction sites flanking the ends. These PCR products were cloned into pCR2.1 or pCRII (Life Technologies) and confirmed by sequencing with either M13-20 or M13-R (Core Services, Wadsworth Center). DNA fragments were cut with the appropriate restriction enzyme, gel purified and recovered (Qiagen) and ligated into pLacInt, a transcriptional lacZ reporter fusion that contains the attachment and integration site (attP/int) of mycobacteriophage L5 and transformed into DH5α (27). Clones were sequenced verified with KM1674 or KM1029. Positive clones were transformed into M. bovis BCG Pasteur and checked for presence of insert using plasmid specific primer combinations.

DNA regions of interest were replaced using sequence overlap extension (SOE), a PCR-based overlap extension method. PCR primers complementary to the desired sequence were used to generate two partially overlapping PCR fragments, which were annealed for use as template for amplification with the outermost end primers containing the restriction sites. These products were then cloned into pCR2.1 or pCRII (Life Technologies) and sequence verified (Wadsworth Center Molecular Genetics Core). Digestion was used to subclone into either the BamHI or ScaI sites of pLacInt as described above.

β-galactosidase assays

Cells were grown as described within text and assays were done as previously described (37). Briefly, cells were de-clumped as described above, mixed with 20mM C2FDG (38) (Life Technologies) and incubated for 3 h at 37°C. The relative expression level for each sample was normalized to 10⁶ cfu based on OD₆₂₀ readings, with an OD₆₂₀ equal to 1 previously determined to have 5×10⁸ cfu by plate counts (37).

RNA and cDNA

RNA was harvested from M. bovis BCG Pasteur after being grown as described within text. Briefly, cells were treated with 5M guanidinium thiocyanate (GTC), harvested with a mini-beadbeater (BioSpec) in the presence of TriZol (Life Technologies) and processed according to manufacturer's instruction. RNA was determined to be DNA free by PCR amplification with 1 μg of RNA and gene specific primers for 23S and 16S determination of no bands present. cDNA was generated using Superscript III reverse transcriptase (Life Technologies) using random oligos as previously described (16).

RESULTS

Genome wide analysis of the CRP regulon

Hypoxia and starvation are thought to be biologically important environmental signals for Mtb during host infection, so we used ChiP-Seq to investigate the genome-wide binding of CRP during log phase growth of M. bovis BCG under low oxygen conditions (1.3% O₂ supplemented with 5% CO₂, which is referred to as baseline throughout the experiment) (28). Alternatively, cells were grown under ambient air conditions to stationary phase and exposed to PBS for 24 h to induce a starvation response. The union of all experiments generated over 900 enriched CRP binding regions in vivo, and the BRACIL blind deconvolution method (26) was used to predict ~2000 individual binding sites within these binding regions. The binding sites were distributed evenly across the genome when normalized peak heights were plotted against their genomic coordinates (Figure 1A) and represented 1365 potential gene regulatory targets (see below and Methods).

Expected peaks were observed for each of 10 regions previously shown to bind CRP by using electrophoretic mobility shift assays (EMSA) (13,15,17). These DNA sequences include the orthologous promoter regions of serC-Rv0885, fredA, Rv0950c-sucC, Rv0145, Rv1386, Rv1158-Rv1159, Rv3857c and Rv1230c, and showed a wide range of normalized peak heights in vivo. For example, the peak between BCG-1290c and BCG-1291c (corresponding to the Rv1230c-Rv1231c intergenic region in Mtb) had a height six times above the experimental mean coverage (also referred to as normalized coverage), while the region between BCG-1004c and sucC (orthologous to Mtb intergenic region of Rv0950c-sucC) had a normalized height of 68. By comparison, the highest normalized signal observed was 196.7, which corresponded to an intergenic site between BCG-0136c-BCG-0137 (orthologous to Rv0103c-Rv0104 in Mtb (Table 1)).

Binding of CRP to DNA is enhanced by cAMP in vitro (13,21–22), so we also examined CRP's in vivo DNA binding profile in BCG exposed to elevated levels of cAMP. Excess cAMP was generated endogenously for these experiments by over-expression of the constitutively active Rv1264 adenyl cyclase catalytic domain (Rv1264Cat) using the Mtb Rv0805 promoter on the multicopy plasmid pMBC621 (16). This approach was previously shown to elevate intracellular cAMP levels by up to 40X (16). No significant differences were observed for in vivo CRP binding in the presence of excess versus baseline cAMP levels with the exception of two strong peaks in the Rv1264Cat samples (Figure 1B). One of these two peaks mapped to a previously identified CRP binding site upstream of Rv0805 that is present within the pMBC621 expression plasmid (14), while the other peak represents a newly identified site. The observed enrichment of these two sites (normalized coverage of 15.7 and 16) correlates well with the previously determined estimate of 3–10 copies/cell for the plasmid’s origin of replication (33). These binding peaks in the Rv1264Cat samples thus serve as a useful watermark for the binding of CRP to the additional plasmid-based copies of these two sites within the Rv0805 promoter region (Figure 1B, Supplementary Figure S1D, F). Dibutylryl-cAMP (dbcAMP) was also added exogenously for some experiments as an alternative means of elevating bacterial cAMP levels, as previously described (39). The cAMP supplementation results were similar to those of the Rv1264Cat experiments, as no CRP binding differences were observed in the cAMP-supplemented versus untreated baseline samples (Supplemental Figure S1). Additionally, no significant differences in CRP binding occurred in rich media compared to the starvation condition (Supplemental Figure S1F).
A high percentage of intragenic binding sites

CRP binding sites from all conditions were classified according to whether they were located within intragenic (within annotated open reading frames (ORFs)) or intergenic (between ORFs) sequences (Figure 1C), and compared with the distribution of total genomic nucleotides in intragenic (91%) versus intergenic (9%) regions. A majority (∼83%) of the binding sites we identified in these ChIP-seq analyses were intragenic and approximately two-thirds of these intragenic sites were located within the middle-coding region of the ORF (Figure 1C). The remaining third fell within the first or last 10% of an ORF. The high percentage of intragenic CRP binding sites was surprising, because transcription factors have been thought to function...
Table 1. Top CRP$_{\text{BCG}}$ binding sites and their potential regulatory targets

| Peak | Baseline Height | Genome coordinates (BCG) | Genome Coordinates (Mt H37Rv) | Location | Potential Targets | Functional description |
|------|----------------|--------------------------|-----------------------------|----------|-------------------|-----------------------|
| 1    | 196.7          | 152045–152088            | 122303–122346               | Intergenic | Rv0103c/crpB      | Cation-transporter P-type ATPase B |
| 2    | 154.1          | 152114–152154            | 122372–122412               | Intragenic | Rv0104*           | cNMP binding protein |
| 3    | 122.3          | 2968736–2968778          | 3015051–3015093             | Intergenic | Rv2699c/Rv2700*   | CHP                   |
| 4    | 76.5           | 410455–410597            | 4131442–4131484             | Intragenic | Rv3688c/Rv3689*   | Secreted essential protein CHP |
| 5    | 66.9           | 305864–305930            | 277834–277900               | Intergenic | Rv0231/fadE4      | Acyl-CoA dehydrogenase FadE4 |
| 6    | 66.4           | 13448–13490              | 13449–13491                 | Intragenic | Rv008/Rv0010c*   | Conserved Membrane Protein |
| 7    | 66.1           | 1092254–1092333          | 106177–1061856              | Intergenic | Rv0990c/Rv0951*  | Succinyl-CoA synthetase |
| 8    | 65.9           | 43115–43157              | 13449–13491                 | Intragenic | BCG_0038*/BCG_0040** | Unknown |
| 9    | 62.3           | 2717743–2717807          | 2752573–2752637             | Intragenic | Rv2451*           | Hypothetical proline and serine rich protein |
| 10   | 61.8           | 44779–44823              | 15113–15157                 | Intragenic | BCG_0041c*       | Conserved Membrane Protein |

+binding site is within repeat region of BCG.
*gene to which binding site is intragenic.

primarily through binding of intergenic sequences (40–42). While only 14% of the CRP binding sites occurred within canonical intergenic sequences (divergent or linear), an additional 3% occurred in convergent intergenic sequences. Combined, we observed ~2-fold enrichment for CRP binding sites among intergenic sequences relative to percent of total genomic sequences. Approximately 55% of these intergenic sites were located upstream of ORFs in a linear orientation (lin, Figure 1C), while 32% were located between two divergent ORFs (div, Figure 1C). Similar distributions of intragenic versus intergenic binding sites were found in the baseline and Rv1264Cat samples (data not shown).

Broad functional roles of putative CRP regulated genes

We separately analyzed intragenic versus intergenic CRP binding sites in relation to their potential gene regulatory targets (as described in Methods), relative to all sites combined, using the orthologous Mt $b$ DNA sequences for each binding site identified by ChIP-seq. Genes that contained or were immediately downstream of a CRP binding site were counted as potential regulatory targets of the CRP that bound at that locus. The distribution of all genes within the genome was calculated using the Tuberculist (43) assignment and the distribution for each functional classification for the genome was compared to the target gene calculations above (Figure 1D). CRP’s potential regulatory targets were broadly distributed among functional classes, with fairly little bias. However, there was an overall enrichment among potential CRP regulatory targets for genes involved in cell wall processes (Category 3) and intermediary metabolism (Category 7), with intragenic binding sites driving these increases. Fewer than expected intergenic sites were associated with intermediary metabolism targets, but stable RNAs (Category 4), including rRNAs and ncRNAs, were enriched for intergenic binding sites (Figure 1D). Genes that encode insertion sequences (Category 5) were also underrepresented as potential CRP regulatory targets when compared to their total representation in the genome.

Characterization of in vivo binding sites

ChIP-seq may detect indirect as well as direct TF binding, so we used EMSA to further characterize binding of
Figure 2. in vitro binding of CRP<sub>BCG</sub> and CRP<sub>Mt</sub> by electrophoretic mobility shift assays. ChIP-seq sites are shown as red circles. The binding motif is shown for the identified binding sites. Probes used in each shift are labeled with small lines. EMSAs were performed with 100ng CRP<sub>BCG</sub> (first column), 100ng CRP<sub>Mt</sub> (second column) or 300ng CRP<sub>Mt</sub> (third column). Excess cold double stranded specific (S) and nonspecific (ns) probes used as competitors are labeled as such.

Comparison of CRP regulon studies

A CRP binding motif was generated using FIMO (Find Individual Motif Occurrences), which is part of the MEME suite (49–51) for all in vivo binding sites identified in the ChIP-seq analyses (Figure 4A). The <i>M. bovis</i> BCG Pasteur genome was then searched for potential binding sites using this motif. Only ~30% of the strong CRP binding motifs instances (defined by FIMO as a <i>P</i> < 0.0001 in which <i>p</i> measures the similarity between the motif and motif instance) in the genome were enriched for binding in the ChIP-seq assays (Supplemental Figure S2), suggesting a high degree of selectivity for the sites that were bound and recovered in these experiments.
Figure 3. Effect of copper exposure on growth in BCG Δcrp strains. Ambient shaking cultures were monitored for growth in (A) untreated, (B) 100uM CuSO₄ and (C) 250uM CuSO₄ for 14 days. Black line, BCG wild-type, red dashed line, BCG Δcrp, blue line, BCG Δcrp single complement, encoded on pMBC1029. The standard deviations are shown from three independent experiments. (*, P < 0.05 as determined by a student’s t-test).

The in vivo binding model in this study closely resembles motifs from two previous studies, each of which used different approaches for prediction of CRPₘₐₜ binding sites (14,25) (Figure 4A). Bai et al. (14) used a combination of E. coli CRP (CRPₑₑ) binding sites and M. tuberculosis DNA sequences recovered by affinity capture using CRPₘₐₜ as a seed their computational analysis. In contrast, Krawczyk et al. (25) approached their analysis from the perspective of the transcription factor, by using the regulon of the C. glutamicum CRP orthologue GlxR to predict orthologous CRPₘₐₜ binding sequences. The strong agreement between binding models in all three studies suggests the presence of a robust core CRP binding site, which is supported by in vitro binding studies (13,14). All three binding models are comprised of a palindromic sequence with two 5 bp half sites separated by a 6 bp spacer with highly conserved GT in the 5’ half and an AC dinucleotide in the 3’ half-sequence. Nonetheless, there was limited overlap (40%) among the intergenic binding sites predicted in the Bai et al. (14) and Krawczyk et al. (25) studies despite their nearly identical binding motif models (Figure 4B).

We examined the concordance of 415 intergenic binding sites identified in this ChIP-seq analysis with predictions made in each of two previous studies to determine whether the CRP binding site discovery algorithm also affected binding site predictions (Figure 4B, top). This comparison was limited to intergenic sequences because intragenic sequences were excluded from the Bai et al. study (14). Eighteen of the total predicted binding sites from Bai et al. (37%) and Krawczyk et al. (29%) were common to all three studies (Figure 4C), but 57% of the Bai et al. sites and 68% of the Krawczyk et al. binding sites were also found in this ChIP-seq study (Figure 4B and C). Krawczyk et al. also identified 47 intragenic binding sites of which 24 (51%) were among our ChIP sites (Figure 4B, bottom). These results suggest that the discovery algorithms biased the specific regulon predictions in the prior studies, but the different approaches were similarly effective for predicting in vivo binding sites overall.

CRP binds upstream of the fumarate reductase and succinate dehydrogenase genes

We examined CRP’s regulatory activity in more detail by focusing on the Rv0250c-Rv0249c locus, which contained four CRP binding sites in the ChIP-seq analysis. Rv0249c-Rv0247c is predicted to encode one of three fumarate reductase/succinate dehydrogenase enzyme complexes in Mtb; the remaining two complexes are encoded by sdhCDAB (Rv3316–3319) and frdABC (Rv1552–1555) (34). An established CRP binding site upstream of frdA (Rv1552) (13) was confirmed in this ChIP-seq study, but a predicted CRP binding site upstream of sdhCD (Rv3316/Rv3317) (25) was not. CRPₘₐₜ also failed to bind to the upstream region of sdhCD in vitro (Figure 5C), even at low pH, although CRPₐₜ was positive for this DNA fragment by EMSA (data not shown).

Only three of the four CRP sites identified by ChIP-seq in the Rv0250c-Rv0249c region were previously predicted (Figure 5A and B). Krawczyk et al. (25) predicted binding sites 2 and 3, while Bai et al. (14) identified site 4. Sites 2, 3 and 4 were also identified in another recent ChIP-seq study in Mtb (52). EMSA was used to investigate the in vitro binding of CRP₈₉₉ and CRPₚₚₙₐₜ to the upstream regions of Rv0249c and Rv0250c, with the frdAbinding site serving as a positive control (Figure 5C). Both CRPₘₐₜ and CRP₈₉₉ proteins bound to the upstream regions of Rv0250c and
Figure 4. Comparison of in silico prediction and in vivo binding of CRP in Mycobacterium. (A) CRP binding motifs predicted from this study, Krawczyk et al. (25) [Figure copyright (c) 2009, Oxford University Press, Nucleic Acids Research, Vol 37, 2009, e97] and Bai et al. (14) [Figure copyright (c) 2005, American Society for Microbiology, Journal of Bacteriology, Vol. 187, 2005, pp 7795–7804. doi:10.1128/JB.187.22.7795-7804.2005]. (B) Top: comparison of the three studies, examining intergenic binding: this study, black, Krawczyk, red, Bai purple. Bottom: comparison of this study and Krawczyk et al. intragenic binding sequences. (C) Core intergenic binding sites from all three studies with overlapping Mtb H37Rv coordinates and corresponding sequences.

Rv0249c, and competition with excess cold specific or non-specific DNA fragments demonstrated binding specificity for both DNA probes (Figure 5D).

Role of intragenic binding in CRP mediated regulation of Rv0250c and Rv0249c

Having established direct binding of CRP upstream of each coding region, we took advantage of binding site 3 within the Rv0250c ORF to test the role of intragenic DNA sequences in the regulation of gene expression in mycobacteria. The high density of CRP binding sites at this locus also provided an ideal opportunity to better understand CRP activity at a potentially complex virulence-associated regulatory region, as relatively few mycobacterial promoters have been carefully analyzed to date. Transcriptional reporter fusions were used to evaluate the role of CRP in regulation of Rv0250c or Rv0249c expression. PCR products A (Rv0250c) or D (large Rv0249c) (Figure 6A), which also corresponds to the EMSA probes shown in Figure 5, were cloned into the BamHI site of pLacInt, an integrative shuttle plasmid that contains a promoterless lacZ gene, to generate transcriptional promoter: reporter fusions. A smaller upstream region of Rv0249c (PCR product E Figure 6A) was also initially tested for promoter activity.

β-galactosidase expression levels were measured for each of these reporter constructs in wild-type or Δcrp M. bovis BCG grown shaking to late log phase in ambient air conditions (17) (Figure 6B). All three DNA fragments were able to drive expression of lacZ, demonstrating the presence of functional promoter elements upstream of each gene. However, expression of the Rv0249c promoters was decreased, while that of Rv0250c was increased, in Δcrp BCG compared to wild-type BCG. Therefore, CRP behaved as an ac-
Figure 5. Binding of CRP to the predicted fumarate reductase and succinate dehydrogenase complexes in Mycobacteria. (A) Locus organization of frdABCD, sdhCDAB and Rv0250c-Rv0247c. (B) Source of CRP binding sites with sequences in relation to the upstream region of Rv0250c and Rv0249c. (C) Electrophoretic mobility shift assays of CRP$_{BCG}$ and CRP$_{Mt}$ to the upstream promoter regions of indicated genes. Black bars above each of the suggested CRP sites in panel (A) indicate probe used for EMSA. (D) Competition assays with excess cold specific (S) or nonspecific (NS) double stranded DNA probes.

We next examined the potential roles of individual CRP binding sites in the regulation of Rv0250c and Rv0249c. CRP$_{Mt}$ binding was tested for each site by performing EMSAs with DNA probes in which one of the binding sites was replaced with an altered sequence. Results indicated that CRP$_{Mt}$ bound at all four wild-type sites, with sites 2 and 3 showing the most robust interactions in vitro (Figure 6C). DNA fragments containing mutated sequences at sites 2 (W1M2) or 3 (M3W4) were then tested for promoter activity with the lacZ reporter system. The importance for promoter activity of each binding site was determined by comparing expression of the wild type versus the mutated promoter sequences in a wild type BCG background. Consistent with the crp knockout results (Figure 6B), mutation of site 2 increased β-galactosidase expression, while modification of site 3 decreased Rv0249c promoter activity (Figure 6D). Together, these results indicate that CRP regulates expression of both Rv0250c and Rv0249c through its binding of specific DNA sequences upstream of each gene.

We used RT-PCR to determine whether Rv0249c can be expressed from the Rv0250c promoter as part of an operon as well as from its own internal promoter. RT-PCR products D, E and F are consistent with co-transcription of Rv0249c and Rv0250c (Figure 6E). Together, these results indicate that Rv0249c can be expressed from the Rv0250c promoter as well as from the internal promoter that requires a portion of the Rv0250c coding sequences for maximal activity (Figure 6B–D). Amplification of junction fragments H and K (Figure 6E) also confirm earlier predictions that Rv0249c, Rv0248c and Rv0247c form an operon (34).

Expression of Rv0249c from the Rv0250c promoter

We next assessed the effects of individual binding sites on expression of lacZ when both the Rv0250c and Rv0249c promoters were present. We constructed a promoter: lacZ fusion using a DNA fragment that spanned both promoters and included all four CRP binding sites (designated ‘WWWW’ for the wild-type sequence of each of the four CRP binding sites). The expression of this construct was compared to that of each individual, wild type promoter (Figure 7A). The expression of the combined wild type construct was at least 10 times higher than either the Rv0250c or Rv0249c promoter fragment and ∼2–3-fold greater than that of the large Rv0249c promoter fragment.

Additional reporter constructs with altered CRP binding sequences were also generated and tested in wild-type BCG grown to late-log or stationary phase in ambient air (Figure 7B) or hypoxia (1.3% O$_2$ + 5% CO$_2$) (Figure 7C). Expression from the WWMW construct, which contained a mutated binding site 3, was significantly reduced compared to the WWWW in all conditions tested (Figure 7B and C). These observations correlate well with those obtained with the individual promoter fusions in which decreased expression was observed in the small and large Rv0249c promoter fragments that had CRP binding site 3 replacements (Figure 6D). However, when expression from the WWWW construct was measured, activity was similar to that of the WWWW construct. This observation suggests that site 4...
Figure 6. CRP binds and regulates at the Rv0250c locus. (A) Organization of the region. ~60nt of the 3’ end of hsp are shown on the far left. The top bar shows approximate distance in nt. Gray arrows represent the anti-sense ncRNAs IG200 (ncRv10250) and IG199 (ncRv10249), identified by Arnvig et al. (72,75). (B) β-galactosidase assay of the individual promoter fusion constructs in M. bovis BCG wild type (black bar) and crp background (hashed bars). (C) Purified CRPMt’s ability to bind to either site 1 or 2 in the upstream region of Rv0250c or sites 3 or 4 in the upstream region of Rv0249c. (D) Contribution of sites 2 and 3 to expression. Black bars, wild type; hashed bars, replaced promoter fusion construct at site 2 or site 3 as indicated by x-axis. (E) RT-PCR using cDNA generated in the absence (−) or presence (+) of reverse transcriptase. Heat killed genomic DNA from M. tuberculosis H37Rv (H). Corresponding products are indicated in panel (A). (F) Summary of data. Small arrows above sites 2 and 4 indicate the positions of the binding sites relative to the transcriptional start sites (tss) as published (71). Binding site 1 is centered 64 nt from the +1 of Rv0250c, while the center of binding site 3 is 67.5 nt from the +1 of Rv0249c. Sites 2 and 4 overlap tss’s for Rv0250c and Rv0249c promoters, respectively. Small arrows under each locus indicate promoter fragments used in these studies. NT not tested. Shown are the means with standard deviations from three independent experiments. (*, P < 0.05; **, P < 0.005 as determined by a student’s t-test).
Figure 7. Contribution of binding sites to regulation of Rv0250c and Rv0249c. (A) The expression of the WWWW construct was compared to each of the individual promoters using β-galactosidase assays. Significance is shown for individual promoters relative to the WWWW promoter. (B and C) Activity of the mutated dual promoter fragments was assessed in late-log (solid bars) or stationary (striped bars) cultures grown in (B), ambient oxygen or (C) hypoxic (1.3% oxygen + 5% CO2) conditions. The means with standard deviations are shown from three independent experiments. (*P < 0.05, **P < 0.005 as determined by a two-tailed students t-test).

has repressing activity that is suppressed by CRP binding to site 3.

Expression from the WWWW fragment was significantly increased in stationary versus log phase bacteria subjected to hypoxia. This trend was also observed for WWMW and WWMM constructs, but not for the MMWW reporter (Figure 7C). This result suggests that the Rv0250c, but not the Rv0249c, promoter is responsive to growth phase during hypoxia, an observation that warrants further investigation. Together, these data indicate that DNA sequences within the Rv0250c coding region are required for maximal activity of Rv0249c promoter, and that CRP binding to intragenic site 3 within Rv0250c plays a significant role in the regulation of Rv0249c-Rv0247c expression in TB complex bacteria.

DISCUSSION

Regulation of succinate dehydrogenase expression by CRP is direct, but complex

ChIP-seq is a powerful technique for detecting global TF binding under biologically relevant conditions and it has been especially useful for identifying new biological roles for TFs within mycobacteria (30,53). In the present study, we combined ChIP-seq analysis with examination of the regulatory roles of individual CRP binding sites at the Rv0250c-Rv0247c locus. Rv0249c is essential for in vitro growth of H37Rv and an Rv0249c transposon mutant is attenuated in infected mice (45,46). Previous studies have also shown that crp is needed for Mtb virulence in a murine infection model (15), and it is likely that reduced expression of Rv0249c-Rv0247c expression in the crp mutant contributes to this attenuated virulence phenotype. Our results establish a significant role for CRP in regulating expression of this putative succinate dehydrogenase complex encoded by Rv0249c-Rv0247c, along with the adjacent core conserved mycobacterial gene Rv0250c (43). Importantly, an intragenic CRP binding site, along with additional Rv0250c sequences, was required for maximal expression of Rv0249c. The presence at this locus of multiple CRP binding sites with differing regulatory properties, along with the likelihood of additional regulatory sequences adjacent to some of these CRP binding sites, could provide opportunities for integration of multiple signal inputs. Similarly, the presence of two promoters allows modulation of Rv0249c-Rv0247c expression in the presence or absence of Rv0250c expression depending on the environment Mtb is experiencing at any given time.

Succinate dehydrogenases complexes are the only enzymes that are part of both the respiratory complex and the TCA cycle, serving as a bridge between oxidative phosphorylation and central metabolism. Succinate dehydrogenase oxidizes succinate to fumarate in a reaction coupled to the reduction of FAD, while fumarate reductase catalyzes the reverse reaction (54). Accumulation and secretion of succinate is essential for Mtb's adaptation to hypoxia (55), and it has been suggested that succinate secretion maintains the membrane potential for energy generation (56). The second half of the TCA cycle in Mtb operates in a reductive direction, which requires conversion of fumarate to succinate by fumarate reductase (56). Our previous observation that CRP also binds to the upstream region of frdA, a putative fumarate reductase complex (13) suggests that CRP is integrally involved in regulating succinate metabolism in Mtb. We find CRP's role in regulating the expression of multiple enzymes involved in the reductive conversion of fumarate to succinate intriguing, and think the importance of CRP for Mtb's adaptation to hypoxic stress during infection especially warrants further investigation. It has been postulated that most mycobacteria harbor two succinate dehydrogenases because of their need to adapt to a range of environments (57). Recent studies by Pecsi et al. (58) examined the roles of the orthologous Rv0249c-Rv0247c (MSMEG_0416-MSMEG_0420 or sdh1) and sdhCDAB (MSMEG_1672-MSMEG_1669 or sdh2) complexes within M. smegmatis, which lacks a frdA or-
Intragenic CRP binding contributes to gene expression

Our observation that intragenic coding sequences from Rv0250c that include CRP binding site 3 significantly increased expression of Rv0249c (Figures 6 and 7A) demonstrates the importance of intragenic DNA sequences for gene regulation in mycobacteria (Figures 6 and 7B–C). Expression of the combined Rv0250c-Rv0249c promoter regions was ~2x higher than the cumulative expression from the individual promoters, suggesting a coordinated regulatory effect between the promoters (Figure 7A). However, it is possible that the inclusion of additional contiguous sequences from Rv0250c coding region, beyond what are included in the largest individual Rv0249c promoter reporter, rather than synergy between the promoters, provides this enhancing effect.

While TF binding within genes does not fit the canonical view of transcriptional regulation in prokaryotes (40–42), sigma factors have been shown to bind at non-canonical promoters located within intragenic regions (61,62). Specifically, SigF, one of 13 sigma factors identified in Mtb, was associated with promoters located within the intragenic regions of Rv1358 and Rv1870c (63). Intragenic binding of CRP orthologs in other organisms has also been reported, but its significance has not been addressed. For example, 43% of the total GlxR binding sites in C. glutamicum detected by ChIP-seq are intragenic, and 30% of these sites are upstream of a divergently transcribed gene (64). Similarly, 54% of the total CRP_Mt binding sites in a recent study, including several of the most enriched peaks (e.g. Rv0010c, Rv3689, Rv2245 and Rv2585c), are located within intragenic regions (52). Intragenic TF binding within prokaryotes is not limited to CRP orthologs, as E. coli RpoH is reported to bind many intragenic sites (65) and over half of the FliA and LeuO binding sites identified in recent ChIP-seq studies are intragenic in E. coli and Salmonella enterica serovar Typhimurium, respectively (66,67).

TF binding within intragenic regions also plays a critical role in eukaryotic transcription (68,69), and a recent study by Stergachis et al. (70) highlighted the co-dependence of intragenic TF binding sites and the protein coding sequences in which they reside. These sequences, termed duons, are intragenic coding sequences that must co-evolve with the TF binding sites to accommodate both codon usage and TF binding requirements. Thus, intragenic TF binding in eukaryotes not only plays a role in regulating downstream gene expression, but also puts evolutionary pressure on the codon usage of the genes within which the TF binds.

The position of CRP binding site 3 centered at −67.5 relative to the internal Rv0249c tss (71) is most consistent with CRP functioning as a canonical transcription factor at this site, and its intragenic location may reflect an example of a bacterial duon. However, the possibility that CRP regulates Rv0249c expression by altering chromosome structure cannot be ruled out. Small antisense RNAs that overlap the intergenic regions between Rv0251c/Rv0250c and Rv0250c/Rv0249c have also been reported (72), but the position of CRP binding site 3 is not consistent with indirect regulation of Rv0249c expression through either of these sRNAs. Nonetheless, the presence of these sRNAs likely contributes to the overall regulatory complexity at this locus, and CRP-mediated regulation of sRNA expression from other intragenic, as well as intergenic, sites seems likely, given the abundance of sRNAs in mycobacteria (72–75).

The very high percentage of intragenic binding sites for CRP identified in this study has important implications for understanding gene expression and developing molecular diagnostics in Mtb. One area of particular relevance is the wide use of single-nucleotide polymorphisms (SNPs) to predict drug resistance in Mtb, because SNPs that do not cause amino acid changes in a drug target gene or its upstream intergenic promoter are not typically considered as strong candidates for drug resistance mutations. However, our results indicate that the relevant genome space considered for resistance-associated SNPs should be expanded to include coding regions upstream of known drug target genes. A recent study (76) investigating the mechanism of ethambutol resistance supports the use of this wider net, as
the investigators identified a correlation between an SNP within the Rv3792 (aff,4) gene and ethambutol drug resistance in Mtb. Rv3792 is immediately upstream of embC, an established target gene for ethambutol resistance. The SNP within Rv3792 is associated with elevated expression of embC, and it is hypothesized that this SNP is located within a intragenic promoter for embC. ChIP-seq data from the TB consortium show potential binding of four TFs within the Rv3792 ORF (30), consistent with the presence of gene regulatory sequences within the Rv3792 ORF.

An expanded view of the CRP regulon

Our 10 most enriched CRP binding sites include both intragenic and intergenic binding sites, only two of which were previously predicted (site 1 within Rv0104, and site 7 between Rv0950c and Rv0951 (sucC), Table 1) (14,25). Site 1 within Rv0104 was also one of the most enriched binding sites in Mtb identified in another ChIP-seq study (52). Functions are not known for more than half the potential regulatory targets of these top 10 binding sites, suggesting that the majority of CRP’s regulatory impact in Mtb remains to be elucidated. We previously demonstrated binding of CRP binding upstream of sucCD (14), whose expression is regulated by CRP under starvation conditions (13). The in vivo binding of CRP at this locus in the present study provides additional support for direct sucCD regulation by CRP and another link between CRP and sucC metabolism. Similarly, the increased sensitivity of the cpr mutant to copper provides the first evidence for CRP-mediated regulation of copper homeostasis in mycobacteria, which could contribute to Mtb survival within macrophages. As noted, two of the most enriched binding sites from this study are located upstream of the ctpB gene, which encodes a putative copper transporter. Copper is toxic to Mtb, so its levels within the cell must be tightly controlled (35,36). The possibility that CRP controls copper resistance through regulation of ctpB expression in Mtb provides another potential role for CRP during host infection that should be explored.

CRP binding in the DNA region upstream of serC is consistent with our previous studies demonstrating CRP’s role in the regulation of serC expression (17), and several other highly enriched binding sites have the potential to regulate cAMP-associated genes. In addition to binding the putative cNMP encoded by the Mtb ortholog Rv0104, CRP directly bound upstream of AC genes Rv1647 (77) and Rv3645, suggesting multiple levels of regulation within the cAMP signaling network in mycobacteria (9). We were interested to note that elevated levels of cAMP did not affect the CRP’s in vivo binding profile (Figure 1B and Supplementary Figure S1), despite evidence that cAMP enhances interactions in vitro (Figure 2, (13–14,22,78)). It is possible that the baseline levels of cAMP under the conditions examined are already sufficient to fully bind CRP (16,79).

There was general agreement between our ChIP results and those of another recent ChIP study in Mtb (52), although we found nearly 10x as many binding sites in our study. The percentage (~83%) of intragenic sites in our study was also greater than the proportion (~50%) in the Kahramanoglu et al. (52) study. Overall, we identified orthologous sequences for ~90% of the Mtb CRP sites recovered by Kahramanoglu et al. in Mtb (52), supporting our use of BCG as a model for identification of in vivo CRP binding in TB complex bacteria. Both studies identified the intragenic Rv0104 binding site as the most enriched site, and our third, fourth and sixth most enriched peaks were also among the top five in the Kahramanoglu et al. study (52). Binding sites in our study with peak heights as low as 4.5 were also found in the Mtb ChIP experiments, further validating CRP binding of these sites in Mtb.

The reason for the much larger number of binding sites in our study is not clear, but we think a higher level of sensitivity in our ChIP analysis is a major contributor to this difference. We used BCG as a model for these studies in part because the more stringent fixation conditions required for Mtb to ensure its biological inactivation led to inefficient chromosomal shearing and reduced the robustness of our ChIP assays. Biological differences could also contribute to the disparity in number of binding sites between the different studies. CRP<sub>B</sub> has a 2x higher affinity than CRP<sub>Mt</sub>, in vitro (13). In vitro binding affinity does not directly translate to in vivo binding, and we previously noted only subtle differences between BCG and Mtb in CRP binding by ChIP-PCR (13). Nonetheless, higher affinity could increase in vivo CRP binding and/or contribute to more efficient recovery of bound DNA fragments by immunoprecipitation. While growth conditions also varied between studies, the relative lack of variation in CRP binding profiles across multiple conditions in our study (Supplemental Figure S1) argues against this being a significant variable.

Significance of low affinity binding sites

We observed a wide range of binding site enrichment, with roughly half of the binding sites having a normalized peak height less than 5. We retained all observed binding sites following background normalization because we think that the many of the less enriched sites have potential to be biologically relevant despite the likelihood of some false positives. Independent experiments across a range of environmental conditions, including the biological repeats of baseline versus Rv1264Cat, generated highly reproducible CRP binding profiles (Figure 1B and Supplemental Figure S1). In addition, identification of binding sites within the duplicated region of BCG<sub>001</sub>-BCG<sub>0040</sub>, served as an internal control of sorts. For example, an enriched binding region was found between 9534–9574 (BCG coordinates) and at the equivalent position in the duplicated region, between 39201–39242 (BCG coordinates). Both regions were similarly enriched, with peak heights of 2.64 and 2.27, respectively. The high number of strong binding motifs that were not bound by CRP further suggests that even low level binding is specific and should be considered in the analysis (Supplemental Figure S2). Moreover, highly enriched binding sites were often associated with closely linked sites that were less enriched, including three sites downstream of the two highest normalized peaks in this study that are intragenic to Rv0104c. These observations fit well with previous ChIP-seq studies in which the binding of eight different Mtb transcription factors showed highly reproducible weak binding profiles (30). In addition, thousands of weak CRP
binding sites were identified within the *E. coli* genome by using ChIP-chip analysis (80), and over 2000 binding sites were found using ChIP-seq with NtcA, a CRP ortholog in cyanobacteria (81).

This theme of extensive weak binding by some transcription factors emerging from global ChIP analyses suggests that cooperative binding of transcription factors at multiple weak sites is a more common mechanism for modulation of promoter activity than has been previously appreciated. Nucleoid associated proteins (NAPs) can also contribute to gene regulation by modifying chromatin structure rather than through the types of direct interaction with RNA polymerase associated with canonical transcription factors. In *Mtb*, Lsr2 is a NAP that binds DNA at AT rich regions (82–84), while HupB is an HU ortholog that interacts with CRP binding regions, such as one associated with Rv1230c (85,86). EspR, was first described as a transcription factor that modulates expression of the virulence-associated espACD operon in *Mtb* (87). However, subsequent biophysical studies demonstrated that EspR regulates transcription over long distances by physically looping the DNA (53,88). CRP in *TB*-complex mycobacteria may represent an example of a ‘dual acting’ transcription factors with NAP activity, as has been suggested for CRP orthologs in *E. coli* and cyanobacteria. The localization of binding sites at positions associated with direct polymerase interaction reported previously (17,21,89) and in this study, including the four binding sites at the Rv0250c-Rv0249c locus, is consistent previously (17,21,89) and in this study, including the four bindings sites in those of a NAP, and are currently addressing this possibility.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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