Functional analysis of an intergenic non-coding sequence within mce1 operon of M.tuberculosis

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Abstract

Background: The mce operons play an important role in the entry of M. tuberculosis into macrophages and non-phagocytic cells. Their non-redundant function as well as complex regulation is implied by the phenotype of mce mutants. Recently, mce1 operon was found to extend over 13 genes, fadD5 (Rv0166) being the first gene of the operon. The presence of a non-coding sequence of 200 base pairs between Rv0166 and Rv0167 is peculiar to mce1 among the four mce operons of M.tuberculosis. We have examined the function of this region.

Results: We predicted putative promoter activity of the 200 base pairs of non-coding, intergenic region between Rv0166 and Rv0167 in silico using MEME software and designate it as intergenic promoter, IGPr. We demonstrate both promoter activity and a putative negative regulatory function of this fragment by reporter assays carried out in the surrogate host M.smegmatis. We find that the repressive elements not only control the native promoter but also repress a heterologous promoter of M.smegmatis. The higher activity of the intergenic promoter in a clinical isolate in comparison with the wild type sequence from M.tuberculosis H37Rv could be correlated with a point mutation within the negative element. We have mapped two transcription start sites for mce1 operon both of which are utilized in M.tuberculosis H37Rv as well as the clinical isolate VPCI591. Our studies show that the promoter activity in the non-coding region is relevant not only in reporter gene expression but also in the expression of mce1 operon in M. tuberculosis cells grown in synthetic medium.

Conclusion: The mce operon of M.tuberculosis H37Rv potentially can be transcribed from two promoters P1 and P2, former mapping upstream of Rv0166 and the latter in the non-coding intergenic region between Rv0166 and Rv0167. The transcription initiation from P1 results in a transcript with Rv0166 while that from P2 will be without it. The sequences between the translation start site of Rv0167 and the promoter P2 have a negative regulatory role, as point mutation within the sequence leads to enhanced activity of P2 as well as a heterologous promoter from M.smegmatis. The mutation detected in the clinical isolate VPCI591 therefore behaves like a gain-of-function mutation.

Background

Tuberculosis causes approximately two million deaths annually and it has been estimated that around two billion people are currently infected with the causative organism, Mycobacterium tuberculosis [1]. Attempts to understand the molecular basis of pathogenesis in tuberculosis include the analysis of genes involved in the entry of the bacillus following the initial identification of mammalian cell entry protein, Mce1A by Arruda et al. [2]. Subsequent whole genome analysis revealed the presence of four mce operons in M.tuberculosis H37Rv, consisting of eight genes with extensive similarity between each other [2,3]. Recently, Casali et al. [4] redefined the boundaries of mce1 making it an operon of 13 genes extending from Rv0166 to Rv0178. The importance of mce operons in virulence is illustrated by various phenotypes observed in knock-out strains and the expression profile of the operons in bacilli in culture and during infection [5-8].

The conservation of most of the mce operons in all members of the Mycobacterium tuberculosis complex, and the presence of orthologous mce genes throughout the genus Mycobacteria, including the non-pathogenic species M.smegmatis suggests their functional importance in processes besides pathogenicity [6,7,9-13]. Casali et al. [4] discovered that fadD5 gene (Rv0166) is also a
part of the mce1 operon, adding to the probable functional diversity of mce operons.

In tune with the proposed functional diversity it has been suggested that mce1 operon could be under the control of a global stress regulator or multiple negative regulators [4,14]. Rv0165c, a homologue of GntR regulator of mce1 operon and Rv1963 a TetR family regulator of mce3 operon are characterized as negative regulators of the respective operons [4,14,15]. The poor consensus of the promoter sequence of mce3 operon at -10 and -35 positions is speculated to reflect the complex regulation of the operon and its ability to interact with multiple sigma factors [4]. Given the importance of mce1 operon and evidences from knock-out studies, any alteration in the expression or genetic polymorphism in mce operons would have significant consequence on the pathogenicity and the severity of infection [6-8,16,17].

Here we examine the function of the non-coding sequence between Rv0166 and Rv0167, which led us to detect both promoter and negative regulatory element within the sequence. A point mutation in the regulatory region abolishes the negative regulation resulting in enhanced promoter activity.

Results
Detection of a putative promoter in intergenic region of mce1 operon
ORF analysis on sequences extending from Rv0166 (nucleotide 194993-196657) across Rv0167 (nucleotide 196861-197658) revealed the expected stop codon for Rv0166 at 196655 and the initiator codon for Rv0167 at 196861. However, no initiator codon was detected in the 200 base pairs between Rv0166 and Rv0167. This region therefore appears to be non-protein coding sequence within the mce1 operon in M.tuberculosis H37Rv. We examined this sequence for probable promoter signature by in silico analysis. We retrieved 10 sequences with demonstrated promoter activity [18] in addition to the intergenic sequence of mce1 operon and aligned them with reference to the translational initiation site of the respective gene. The presence of consensus motif was analyzed using MEME http://meme.nbcr.net/meme3/meme.html. Two motifs GGTT [CG] [CG]T and TT [AT] [TC] [CT] [GA] [ACG]C were identified (p value > 1.31-e04) and both the motifs are present in the non-coding intergenic region between Rv0166 and Rv0167 of mce1 operon (Figure 1C &1D and Additional file 1). Since we detect landmarks of promoters known in M.tuberculosis within this region, we refer to it, henceforth as intergenic promoter (IGPr). We undertook the functional characterization of the predicted promoter activity of IGPr. We analyzed the effect of a point mutation in the IGPr, detected in a multi-drug resistant clinical isolate, VPCI591, under an independent analysis of genetic polymorphism in mce operons of clinical isolates of M.tuberculosis (unpublished).

Promoter Activity of IGPr
A 200 bp fragment containing IGPr sequence was amplified from M.tuberculosis H37Rv and cloned in promoter-less shuttle vector pSD5B, upstream of the lacZ as the reporter gene to generate pPrRv. Similarly 200 bp fragment from VPCI591 was cloned to produce pPr591 and both were tested for promoter activity in M.smegmatis. Different constructs used in the study are shown in Figure 2. Since a repression of mce1 operon at stationary phase was reported earlier [5], we analyzed the promoter activity of the two constructs both at log and stationary phase of growth, by ONPG assay using cell-free extracts from transformed M.smegmatis cells (Figure 3). The difference in the promoter activity of IGPr from VPCI591 (pPr591) is higher than that from M.tuberculosis H37Rv (pPrRv) by 12 fold (1025 vs 85 units of β-galactosidase activity) in log phase, which reaches 18 fold (2265 vs 130 units) in stationary phase (Figure 3). By comparing IGPr with previously characterized weak (pSD5WP) and strong (pSD5SP) promoters from M.tuberculosis [19], we find that pPrRv is a weak promoter while pPr591 acts as a strong promoter.

Deletion analysis of IGPr region
In order to delineate the region of promoter activity within the 200 base pairs of IGPr, we made a series of deletion constructs. We generated amplicons corresponding to (-50 to +1), (-100 to +1), (-150 to-50) and (-200 to -100) and cloned them in pSD5B for expression in M.smegmatis (Figure 2). The promoter activity of 200 base pairs from M.tuberculosis H37Rv (pPrRv) is very low compared to that of the same region from VPCI591 (pPr591); 130 vs 2265 units respectively. The promoter activity is highest when -100 to +1 is deleted (pPrD) both in log (2255 units) and stationary phase of growth (4961 units, Figure 3); while it is negligible, when -200 to-100 is deleted (pPrBS91; 52 and 89 units in log and stationary phase respectively). Additionally, the fragment containing only -150 to -100 (pPrC591) shows poor activity. Therefore we conclude that the promoter activity is restricted to around 50 base pairs from -200 to -150 within IGPr (Figure 3). Interestingly, significant promoter activity is detected in the construct that is deleted for -100 to +1 (pPrD). These results suggest that -100 to +1 region cloned in pPrRv has a negative effect which is lost in pPr591 derived from the clinical isolate VPCI591. We correlate this gain of expression due to loss of repression to the presence of a point mutation (G > C) at -61 in VPCI591.

To compare the mRNA levels from the two constructs, we isolated total RNA from M.smegmatis transformed...
with pPrRv (200 base pairs from *M. tuberculosis* H37Rv) and pPr591 (200 base pairs from VPCI591) and the transcript level was estimated by quantitative PCR with lacZ as target gene and sigA as the endogenous control in log and stationary phase. At log phase there is nearly two fold increase in lacZ transcripts in pPr591 as compared to pPrRv whereas in stationary phase it is more than four fold (Figure 4). The correlation between β-galactosidase activity and the mRNA levels of lacZ clearly indicate greater transcriptional activity in pPr591 than pPrRv. The difference in enzyme activity is much higher than the difference in mRNA levels as known in other cases [20-22].

Mapping the transcription start site in *M. tuberculosis*

We identified transcription start site of Rv0166 and Rv0167 *in vivo* in *M. tuberculosis* H37Rv and VPCI591 using fluorescence tagged primers in primer extension assay using RNA templates. The absence of DNA contamination in RNA preparation was confirmed by PCR for Rv0166 and Rv0167 in absence of reverse transcriptase (data not shown). The sizing of the products was carried out by genescan analysis and the TSS was detected at -65 position from the translation initiation site of Rv0166 and at -56 position from the translation initiation site of Rv0167 (Figure 5B-E), suggesting that there are two potential promoters for *mce1* operon gener-
ating two transcripts, one including Rv0166 and the other without it (Figure 5A). Further, this demonstrated that both promoters are active in the genomic context of *M. tuberculosis*. Considering the translation initiation site of Rv0167 as +1, we map the transcription start site within IGPr at -56 position and the mutation in VPCI591 at -61 position.

**Estimation of mce1 operon transcript levels in *M. tuberculosis***

The transcript level of Rv0167, Rv0170 and Rv0174 of *mce1* operon downstream to IGPr in *M. tuberculosis* and VPCI591 was analyzed by quantitative PCR with rpoB as the endogenous control (Figure 6A). The data reveals 1.5 fold upregulation of the *mce1* operon genes in VPCI591 as compared to *M. tuberculosis* H37Rv (Figure 6B). The difference at protein level is considerably higher than at the transcript levels in case of β-galactosidase, similar enhancement in Mce1 protein levels could also be anticipated.

**Effect of the regulatory sequence of IGPr on heterologous promoter**

To examine if the negative regulatory site, -100 to +1 region of IGPr functions independent of the associated promoter activity, we cloned it downstream of a heterolo-
gous promoter in pSdps1, driving the expression of β-galactosidase [23]. pSdps1 has 1 kb upstream region of the gene MSMEG_6467 from M. smegmatis. The promoter in pSdps1 is inducible under glucose starvation; at 0.02% glucose in Middlebrook 7H9 liquid medium in stationary phase [23]. By inclusion of +1 to -100 from IGPr of H37Rv (pDPrBRv) the promoter activity decreased by 35% relative to the control plasmid pSdps1 (895 versus 1358 units, Figure 7). When +1 to -100 from VPCI591 was cloned downstream to dps promoter (pDPrB591), the repression was reversed and the promoter activity was enhanced by 25% over that of pSdps1 (1709 versus 1358 units). This shows that negative regulation by IGPr functions in the context of a heterologous promoter also.

Discussion

The mce1 operon is different from other three mce operons in having Rv0166, a fatty acyl CoA synthetase that catalyzes the initial step in lipid degradation [4,24]. On the other hand, mce4 operon is known to be a part of the regulon involved in cholesterol metabolism, however it seems to be just one of the many possible lipid substrates. Furthermore, it is speculated that mce1 operon may not have a role in cholesterol import as the loss of Mce1 transporter system does not appear to affect the residual uptake of cholesterol in mce4-deficient strain [25].

The presence of 200 base pairs of non-coding sequence between Rv0166 and Rv0167 is yet another feature peculiar to mce1 operon among the other four operons present in M. tuberculosis. In most other operons and also the other genes within mce1 operon, the intergenic distance is not more than one or two codons and often the translation initiation site of one gene is within the coding sequence of the adjacent gene [12]. In silico analysis using GeneRunner software shows the absence of any ORFs in the intergenic region between Rv0166 and Rv0167, while ribosomal binding site corresponding to the translational start site of Rv0167 is reported in Tuberculist database. Although most prokaryotes do not have introns, the intergenic region in transcripts serve as substrates for several endonucleases such as RNaseP involved in mRNA processing and hence are implicated in the regulation of gene expression [26-29].

We have characterized the promoter and negative regulatory activity in the surrogate host M. smegmatis, but the detection of two active transcription initiation sites both in M. tuberculosis H37Rv and VPCI591 suggests both promoters are functional in their native context also.

However the increased promoter strength of the regulatory region from VPCI591 in M. smegmatis is not reflected in the difference in the transcript levels for mce1 operon genes in VPCI591 as compared to M. tuberculosis H37Rv. This may have two reasons, one that both P1 and P2 promoters are active in vivo and therefore contribute to the transcript levels in both the strains, while in M. smegmatis we observe a clear upregulation of P2 when the negative regulation is lost due to point mutation and P1 is absent (since only P2 is cloned in the plasmid). Further, the difference in fold increase in β-galactosidase activity vis-à-vis its transcript levels are significantly different. Similar discordance between protein and mRNA levels is reported in Mycobacteria and S. cerevisiae [20-22]. Moreover, in vivo mce1 operon could be under the regulatory influence of several factors acting directly or indirectly [4].

We looked for concordance in the expression level of Rv0166 and 0167, as polycistronic mRNA including Rv0166 in M. tuberculosis is reported by Casali et al. [4].
For comparison, we examined the expression of pairs of adjacent genes in five different operons including Rv1964 and Rv1965 of \textit{mce} 3 operon, Rv2498c and Rv2499c of CitE-scoA operon along with that of Rv0166 and Rv0167 of \textit{mce} 1 operon. The expression data was taken from published microarray profiles of \textit{M.tuberculosis} H37Rv cells grown in culture [30]. Pearson’s correlation coefficient in the range of 0.8 to 0.58 is observed in all cases except Rv0166 and Rv0167 of \textit{mce} 1 operon [0.24; Additional file 2]. Similar difference between coefficient of correlation...
was observed when we considered the data from clinical isolates grown in Middlebrook 7H9 medium [31]. These results imply that the transcript level is lower for Rv0166 compared to Rv0167, as Rv0166 can be transcribed only from P1 while Rv0167 can be transcribed from both P1 and P2 promoters. Thus lending support to our data suggesting that both promoters of \textit{mce1} operon are active in cells in culture.

Though \textit{M. tuberculosis} system is replete with examples where the expression of an operon is driven by multiple promoters [32-34], the promoters are known to drive the expression of all the genes of the operon. A study on \textit{furA-katG} region shows differential regulation of two \textit{katG} promoters resulting in two different transcripts depending on the stage of infection of \textit{Mycobacterium tuberculosis} [35]. However the consequences of transcription from intergenic promoter could be different. It can only be speculated that two different polycistronic mRNA varying in coding capacity for a catalytic function can be produced by \textit{mce1} operon: one that includes fatty acyl-CoA synthase (Rv0166) and other lacking it, in absence of \textit{in vivo} infection data. This suggests the possible modulation of the function of \textit{mce1} operon in cell entry and lipid metabolism vis-à-vis its catalytic function. However, it remains to be examined if the intergenic promoter/regulatory region in \textit{mce1} operon could bring about differential regulation during infection.

The \textit{mce1} and \textit{mce2} operons are known to be negatively regulated by divergently transcribed genes mapping immediately upstream of the operon [4,36]. Though Mce1R, the product of Rv0166c is characterized as a negative regulator of \textit{mce1} operon, its binding site is not deciphered so far. The results of Casali et al. [4] suggest that the site of interaction of Mce1R is in a region upstream of Rv0166, while the negative regulatory element we have identified is downstream to Rv0166. Further we failed to detect direct binding of intergenic promoter with purified His-tagged Rv0166c cloned in pET-28a in gel-shift assays even at high molar ratio of protein to DNA (2000:1). Therefore, it appears that \textit{mce1} operon has more than one negative regulator. However, it is interesting to note that a heterologous promoter in pSdps1 is also down regulated by the regulatory region of -100 to +1 fragment of IGPr, thus demonstrating that the 100 bp fragment is necessary and sufficient for repressive activity.

Casali et al. [4] also observed that \textit{mce1} operon can be repressed independent of Mce1R by incubation in DMEM medium and suggest that \textit{mce1} operon may be under multiple negative regulators. Based on their study on lipid degradation operon Kendall et al. [24] observed that operon regulation may be more complex than one would expect for a prokaryotic system and may not be guided by just a single regulator.

**Conclusions**

Our data strongly supports the presence of two functional promoters for \textit{mce1} operon in \textit{M. tuberculosis} that could potentially segregate different functions of a single operon. Our results demarcating the regulatory sequences in the intergenic region of \textit{mce1} operon provide a handle for identifying interacting factors and studying the implications of derepression in the clinical isolate.
Methods
In silico analysis
The non-coding sequence was detected through ORF analysis of mce1 operon using Gene Runner Version 3.01 available at http://www.generunner.net. To identify promoter-like sequences in the intergenic region, the 200 base pair sequence between Rv0166 and Rv0167 was aligned with validated promoter sequences given by Bashyam et al. [18]. The presence of a consensus motif was analysed using the MEME program http://meme.nbcr.net/meme3/meme.html available in the public domain [37].

Enzymes and Chemicals
Restriction enzymes, T4 DNA ligase, RNase free DNaseI were purchased from MBI Fermentas. Kanamycin was from Himedia laboratories Pvt. Ltd., India. The reagents for competent cell preparation, transformation, reporter assays were obtained from Sigma laboratories, USA. [γ-32 P] ATP was from Board of Radiation and Isotope Technology, India.

Bacterial strains and culture conditions
All the strains and plasmid constructs used in the present study are described in Additional file 3. M. smegmatis mc²155 (ATCC 700084) was obtained from Dr. Anil Tyagi, South Campus, University of Delhi and Mycobacterium tuberculosis H37Rv were obtained from Central Jimal Institute for leprosy, Agra, India; Mycobacterium tuberculosis VPCI591 is a clinical isolate from Vallabhbhai Patel Chest Institute; Delhi. M. tuberculosis strains were grown in Middlebrook 7H9 broth supplemented with OADC (Oleic acid, Bovine albumin fraction V, dextrose-catalase) from Difco laboratories, USA and 0.05% Tween 80 (Sigma). M. smegmatis was grown either in Middlebrook 7H9 supplemented with glycerol or on Middlebrook 7H11 plates. Middlebrook 7H9 medium was supplemented with appropriate concentration of glucose whenever M. smegmatis clones with dps promoter were grown, as specified in the results section. Cloning was carried out in Escherichia coli DH5α (Stratagene) grown in Luria-Bertani medium (Difco laboratories, USA). Kanamycin (20 μg/ml) was included for maintenance of plasmids. Transformation in E. coli DH5α was carried out using heat shock method [14] and in M. smegmatis mc²155 by electroporation [19] using Gene Pulser (Bio Rad Laboratories Inc. Richmond, California) at 2.5 kV, 25 μF and 1000 Ω in 0.2 cm gap electroporation cuvettes.

The primers used are listed in Additional file 4. The intergenic region of Rv0166-Rv0167 was PCR amplified using primers Mce1AF and Mce1AR from genomic DNA of Mycobacterium tuberculosis H37Rv and the clinical isolate VPCI591, cloned in Xbal-SphI sites of pSD5B [Additional file 4, [38]]. Deletion constructs were created by PCR amplification of selected region with specific primers followed by cloning in XbaI-SphI sites of pSD5B. Fragment corresponding to +1 to -100 region of intergenic promoter region (IGPr) was amplified from both M. tuberculosis H37Rv and VPCI591 strain, cloned in the vector pSDps1 downstream of glucose regulated dps promoter [23,39] to generate pDPrBRv and pDPrB591 respectively at VspI-PstI site and electroporated into M. smegmatis mc²155. pSDps1 has 1 kb upstream region of dps gene (MSMEG_6467, DNA binding protein from starved cells) from M. smegmatis. The transformants were screened by PCR, confirmed by restriction digestion and sequencing.

The expression of β-galactosidase was assayed both in the log (O.D.600 0.8) and stationary phase (O.D.600 2.0) cultures of the transformants using modified protocol of Miller et al. [40]. Promoter activity is expressed as units of β-galactosidase activity in terms of ONPG converted to o-nitrophenol (nmol per min per mg of protein).

Mapping transcription start site
The transcription start site was mapped using the strategy described by Lloyd et al. [41]. Primer extension was carried out on DNA free RNA with fluorescence labeled primers HEX-tsp1 and FAM-tsp2 mapping 100 nucleotides downstream of the translation initiation site of Rv0166 and Rv0167 respectively [Additional file 4]. The DNA sequence analysis and Genescan analysis was carried out at the commercial facility of The Centre for Genomic Application, Okhla, New Delhi and Labindia, Udyog Vihar, Gurgaon, India respectively. The Genescan analysis was carried out on 3130×l Genetic Analyzer from Applied Biosystems with GSLIZ 500 as marker set. The data was analyzed using GeneMapper V4.0.

Quantitative RT-PCR
The transcriptional activity in log and stationary phase, was estimated by quantitative PCR using cDNA samples. 15 ml cultures of M. tuberculosis H37Rv and VPCI591 from log (day10) and stationary phase (day 20) were harvested at 4°C. RNA isolation was performed using RNeasy Mini Kit (Qiagen) and treated with DNaseI (MBI Fermentas). Absence of amplicons in PCR without reverse transcriptase confirmed the absence of DNA contamination. 500 ng of DNase I treated total RNA samples extracted were retrotranscribed using cDNA synthesis kit (MBI Fermentas) with random hexamer primers. Real Time PCR was performed using SYBR Green PCR master mix (Applied Biosystems, USA); sigA or rpoB was used as endogenous control. The relative expression of mce1 operon genes (Rv0167, Rv0170 and Rv0178) in M. tuberculosis H37Rv and VPCI591 and lacZ expression from the clones pPrRv and pPr591 in M. smegmatis was deter-
mined, using similar protocol. The experiments were repeated three times and the data was analyzed using the ΔΔCt method [42].

Additional material

Additional file 1 Detection of putative promoter motif. Output consensus sequences of MEME mapped (bold upper case) on validated promoter sequences. The input sequences are from T6 to PA [gyr]. IGPr is the query sequence. Translation start site (ATG/GTG) of the gene driven by each promoter used as the reference for alignment is shown in capital.

Additional file 2 Comparison of expression level of adjacent genes in different operons. Pearson's correlation coefficient of the first two genes of mce1 operon is compared to that of neighbouring genes in five different operons. Operons predicted by Rebask et al [44] and Moreno-Hagelsieb et al [44] are used; * represents the operons extending from Rv1460 to Rv1466 (operon A) and Rv3083-3089 (operon B). Least correlation is observed between Rv0166 and Rv0167. Expression data of Fu and Fu-Liu [30] was taken for analysis.

Additional file 3 Strains and plasmids used in the present study.

Additional file 4 List of primers.

Authors' contributions

V8 and MB conceived the project. RP helped with M tuberculosis culturing. SB and MI contributed equally to the experiments. V8, MB, SB and MI participated in experimental design and data interpretation and manuscript preparation. All authors read and approved the manuscript.

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