Cyclic AMP in Mycobacteria: Characterization and Functional Role of the Rv1647 Ortholog in *Mycobacterium smegmatis*

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Mycobacterial genomes are endowed with many eukaryote-like nucleotide cyclase genes encoding proteins that can synthesize 3',5'-cyclic AMP (cAMP). However, the roles of cAMP and the need for such redundancy in terms of adenyl cyclase genes remain unknown. We measured cAMP levels in *Mycobacterium smegmatis* during growth and under various stress conditions and report the first biochemical and functional characterization of the MSMEG_3780 adenylyl cyclase, whose orthologs in *Mycobacterium tuberculosis* (Rv1647) and *Mycobacterium leprae* (ML1399) have been recently characterized in vitro. MSMEG_3780 was important for producing cAMP levels in the logarithmic phase of growth, since the ∆MSMEG_3780 strain showed lower intracellular cAMP levels at this stage of growth. cAMP levels decreased in wild-type *M. smegmatis* under conditions of acid stress but not in the ∆MSMEG_3780 strain. This was correlated with a reduction in MSMEG_3780 promoter activity, indicating that the effect of the reduction in cAMP levels on acid stress was caused by a decrease in the transcription of MSMEG_3780. Complementation of the ∆MSMEG_3780 strain with the genomic integration of MSMEG_3780 or the Rv1647 gene could restore cAMP levels during logarithmic growth. The Rv1647 promoter was also acid sensitive, emphasizing the biochemical and functional similarities in these two adenyl cyclases. This study therefore represents the first detailed biochemical and functional analysis of an adenyl cyclase that is important for maintaining cAMP levels in mycobacteria and underscores the subtle roles that these genes may play in the physiology of the organism.

Cyclic AMP (cAMP) is an important intracellular second messenger that is involved in several signaling pathways in both bacteria and eukaryotes (2, 5). Studies have shown a role for cAMP not only in physiological processes such as catabolite repression and sporulation (16) but also in the regulation of several virulence pathways, e.g., in *Pseudomonas*, *Vibrio*, and *Candida* species (1, 17, 19, 40, 41). The synthesis of cAMP is dependent on the activity of adenyl cyclases, which can be membrane-bound or soluble proteins (7). The largely differing amino acid sequences of nucleotide cyclases allow their classification into six classes, among which the class III nucleotide cyclases have the widest phylogenetic distribution (7, 37).

Adenylyl and guanylyl cyclases of the class III family are proteins that form head-to-tail dimers and generate either two (homodimeric enzymes) or one (heterodimeric enzymes) active site at the dimer interface (20). Nucleotide cyclases catalyze the conversion of ATP or GTP to cAMP or GMP, respectively, in a metal-dependent manner, and biochemical and structural studies have identified specific sequence motifs that are involved in substrate (ATP or GTP) or metal (Mg\(^{2+}\)) binding (20). In contrast to the limited domain compositions of mammalian nucleotide cyclases, the bacterial counterparts are often fused to a variety of domains, highlighting their ability to be regulated allosterically (37). Their presumable involvement in diverse biological phenomena in bacteria is perhaps due to this remarkable biochemical and structural versatility.

Recent studies have shown the presence of a large number of class III nucleotide cyclases in the genomes of several bacteria such as actinobacteria, alphaproteobacteria, and cyanobacteria (37). The genomes of *Mycobacterium tuberculosis* strains H37Rv and CDC1551 encode 16 and 17 class III cyclases, respectively (24, 33). The presence of multiple cyclases within a genome is in striking contrast to the well-studied paradigm of the single-copy class I cyclase gene in *Escherichia coli*. Studies on the role of these mycobacterial cyclases, and indeed cAMP, in the biology of the organism are therefore critical to a comprehensive understanding of cyclic nucleotide monophosphates in more complex signaling networks. For example, the deletion of Rv3676, a homolog of the cAMP receptor protein from *M. tuberculosis*, leads to the attenuation of *M. tuberculosis* in macrophage infections and in the mouse model of tuberculosis (29), suggesting that several important genes are under cAMP regulation in *M. tuberculosis*. Furthermore, inactivating point mutations within the cAMP or DNA binding domains of Rv3676 were found in several strains of *Mycobacterium bovis* BCG, and this apparent lack of Rv3676 function was suggested to be one of the additional reasons for the attenuation of these BCG strains (42). Microarray-, proteomic-, and SELEX-based approaches have also helped in the identification of several cAMP-regulated genes (3, 11, 29). Those studies suggest that cAMP levels must be finely regulated by the adenyl cyclases, and the presence of multiple adenyl cyclases in the genome implies a certain amount of
redundancy in the proteins that are involved in cAMP production. However, given the diversity of these enzymes in terms of their domain organization, mycobacteria may have evolved a specific need for different cyclases at different times, and this can be studied only by a systematic analysis of the role of each cyclase gene, one at a time.

The Rv1647 cyclase from M. tuberculosis is conserved in Mycobacterium leprae (ML1399), even though 12 other cyclases found in M. tuberculosis are deleted or are pseudogenes in M. leprae (35). In addition, Rv1647 orthologs can be found in M. bovis, Mycobacterium avium, Mycobacterium smegmatis, as well as Mycobacterium marinum (38). We therefore hypothesized that Rv1647 and its orthologs serve a housekeeping role in mycobacteria and maintain cAMP levels in the organism. In this study, we investigated the role of MSMEG_3780, the Rv1647 ortholog, in the biology of mycobacteria. Our results provide evidence to suggest that MSMEG_3780 maintains basal levels of cAMP in M. smegmatis during the logarithmic phase of growth. Certain stress conditions alter intracellular levels of cAMP in M. smegmatis, and the reduction in intracellular levels of cAMP seen upon the acidification of cultures could also be correlated with a reduced transcription of the MSMEG_3780 gene. This reduction in intracellular cAMP levels was restored upon complementation of the knockout strain with the MSMEG_3780 or the Rv1647 gene, suggesting that M. smegmatis could be used to understand fundamental aspects of cAMP and cAMP signaling in mycobacteria.

MATERIALS AND METHODS

Cloning, expression, and purification of MSMEG_3780. The MSMEG_3780 gene was annotated as tigrN by TIGR was cloned using primers MS3780F-LFwd (5'-ACA GGTCAGGTGTTCCAGACG-3') and MS3780rws (5'-CGCGCGGCCGC GCCACCACATCGACTATGGCTATGACG-3'). PCR was carried out on the genomic DNA of M. smegmatis mc²155 (35), and the product was digested with Xhol and NotI and cloned into the compatible SalI-NotI sites of pPROEx-HT-C to generate plasmid pPRO-MSMEG_3780FL. The catalytic domain of the protein, from amino acids 149 to 392, was obtained by cloning the EcoRI-NotI fragment from pPRO-MSMEG_3780FL into pPROEx-HT-A to generate plasmid pPRO-MSMEG_3780CD. The protein was expressed and purified from E. coli BL21(DE3) cells as described previously for the Rv1647 protein (35). Adenyl cyclase assays were performed as described previously, and cAMP was measured by radioimmunoassay (36).

GST pull-down assays. The EcoRI-NotI fragment from pPRO-MSMEG_3780-3780CD was cloned into similar digested vector pGEX-6P-1 to generate plasmid pGEX-MSMEG_3780CD. Plasmid pGEX-Rv1647CD was generated by substituting the Xhol-NotI fragment from pPRO-1647-528 (35) into similar cut vector pGEX-6P-1. Glutathione S-transferase (GST), GST-MSMEG_3780, and GST-Rv1647 proteins were purified according to a procedure described previously (18). GST or GST fusion proteins (3 μg) on glutathione beads were interacted independently with 5 μg of His-tagged prey protein in 100 mM buffer at pH 9.0 (25) containing 5 mM CHAPS (3-[3-cholamidopropyl]-dimethylammonio)-1-propanesulfonate), 5 mM 2-mercaptoethanol, 12 mM M, and 10% glycerol at 4°C for 1 h, and unbound proteins were removed by washing the beads five times with 1 ml of interaction buffer. Bound proteins were assessed by immunoblotting with a hexahistidine-specific monoclonal antibody (GE Health Sciences).

Measurement of cAMP levels in M. smegmatis. Mycobacterial strains were grown in 7H9TY (Middlebrook 7H9 broth supplemented with 0.2% glycerol and 0.05% Tween 80) at 37°C, with shaking at 250 rpm, or on Middlebrook 7H10 agar. Whenever necessary, hygromycin (150 μg/ml for E. coli and 50 μg/ml for mycobacteria) was used.

Cells were centrifuged at 13,000 × g, cells washed with a solution containing 10 mM Tris-HCl (pH 7.5), 0.89% NaCl, and 0.05% Tween 80 (TBST); the pellet was resuspended in 0.1 N HCl; and the suspension was boiled for 10 min. Samples were then flash-frozen and lyzed by bead beating for 1 min. Aliquots of the lysate were taken for estimation of cAMP by radioimmunoassay (35). Spent medium (supernatant of the 13,000 × g spin) was acidified by twofold dilution in 0.2 N HCl and boiled for 10 min, and suitable aliquots were monitored for cAMP directly by radioimmunoassay.

For measurement of cAMP levels under stress conditions, 2-day-old cultures started from single colonies were diluted 1:100 into fresh medium and grown for 30 h. Cells were then centrifuged, washed in chilled sterile TBST, and resuspended in either fresh control medium or medium containing 50 mM MES [2(N-morpholino)ethanesulfonic acid] (pH 5.5), 5 mM H₂O₂, and 1 mM sodium nitroprusside, 10 mM sodium phosphate (pH 7.2) containing 0.9% NaCl (phosphate-buffered saline [PBS]), 0.02% glycerol, or 0.05% sodium dodecyl sulfate (SDS) (22). Cultures were grown for 1.5 or 5 h at 37°C and/or at 42°C for 1.5 h. Cells were then centrifuged, and the pellet was taken for measurement of cAMP levels.

Generation of the ΔMSMEG_3780 strain. A strain with a deletion of the MSMEG_3780 gene (ΔMSMEG_3780) was generated using the suicide vector approach (27). The 5' knockout fragment consisting of 1,013 bases upstream of MSMEG_3780 was amplified by PCR using primers MS3780-5'KO-HindIII-fwd (5'-ACAGTGGAACCTCGCCTGCATGCACTG-3') and MS3780-5'KO-XbaI-rvs (5'-GGGCTTTTCAGACCGCGCGTGCTGCTGAGC-3'). A fragment of 1,101 bases downstream of the catalytic domain of MSMEG_3780 was amplified using primers MS3780-3'KO-XbaI-fwd (5'-ACTCCCTAGATTTGACACG-3') and MS3780-3'KO-NotI-rvs (5'-TCGCGCGCGCGCGCGTTCCGCGTTTGC-3'). The 5' fragment was cloned into vector pBluescript II(−) using the HindIII and XbaI sites to obtain plasmid pBluescript-MSMEG_3780-5'KO, and the 3' fragment was cloned into pPROEx-HT-B using the XbaI and NotI sites to generate plasmid pBluescript-MSMEG_3780-3'KO. DNA sequencing confirmed the identity of the cloned sequences compared to that reported in the genome. The HindIII-XbaI and XbaI-NotI fragments from pBluescript-MSMEG_3780-5'KO and pBluescript-MSMEG_3780-3'KO, respectively, were ligated into HindIII- and XbaI-cut vector p2NIL to generate plasmid p2NIL-ΔMSMEG_3780-5'3'KO-PacI. Plasmid p2NIL-ΔMSMEG_3780-5'3'KO-PacI (1 μg) was electroporated into M. smegmatis mc²155, and single crossovers and double crossovers were obtained essentially as described previously (27). Double crossovers were further tested by genomic PCR and Southern blotting as described below to obtain the ΔMSMEG_3780 strain.

To construct the ΔMSMEG_3780 strain complemented with the MSMEG_3780 gene driven by its own promoter, a Stul-XbaI fragment from pPRO-MSMEG_3780FL was cloned into compatible HindII-XbaI sites of vector pBluescriptII(−) to generate plasmid pBluescript-MSMEG_3780FL. The insert was removed from this plasmid as a KpnI-XbaI fragment and cloned into similarly digested integration vector pH949 (36) harboring CmR of W. Bishai’s laboratory. The 5' fragment was cloned into vector pH949 to generate plasmid pH949-ΔMSMEG_3780, which was electroporated into the ΔMSMEG_3780 strain. Positive integrants (ΔMSMEG_3780C) carrying the required insert were screened by colony PCR and validated by Southern hybridization. PCR were carried out with primers MS3780FL-Fwd and MS3780rws on genomic DNA prepared from the wild-type strain and the ΔMSMEG_3780 and ΔMSMEG_3780C strains to confirm the presence of the deletion in the ΔMSMEG_3780 strain and the complementation of MSMEG_3780. Southern blotting was carried out by electrophoretic separation of 2 μg of genomic DNA of wild-type, knockout, and complement strains digested with SacI and by transferring the DNA to Hybond nylon membranes (GE Healthcare). The probe was prepared with a random hexamer labeling kit (Promega) using an EcoRI-Smal fragment present upstream of the catalytic domain of MSMEG_3780. The blot was probed at 60°C for 16 h and washed once in 2× SSC (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate) for 15 min at 60°C prior to exposure to a phosphorimagener.

Preparation of RNA from M. smegmatis. Cells (5 × 10⁸ CFU) were lysed by bead beating in 200 μl of Tri reagent (Sigma-Aldrich) prewarmed to 65°C. The lysate was mixed by pipetting, followed by centrifugation at 12,000 × g for 10 min at 4°C. The supernatant was treated with chloroform (40 μl) followed by centrifugation at 12,000 × g for 15 min at 4°C. The RNA was precipitated with 100 μl of isopropanol alcohol, and the RNA pellet was treated with RNase-free DNase (1 to 3 units). RNA (5 μg) was used for reverse transcription using 200 μl of Moloney murine leukemia virus reverse transcriptase enzyme in the presence of 10 units of RNase inhibitor using random primers. PCR was carried out using 1 μl of the reverse transcriptase reaction mixture and primers MS3779RTF (5'-GACCTTTTCTGCGGTACG-3') and MS3779RTR (5'-CTGGCGGCGCTGTTAACGTAC-3'). The sig4 transcript levels were assessed using primers MSsigART2426F (5'-ACG
AGGAAAGGTCCGAGGCCTTG-3') and MStsigART356R (5'-GTCCTGACCTTCTTCGGCTGTTAG-3').

Fractionation of M. smegmatis cells for Western blot analysis. Cultures of M. smegmatis were grown at 37°C and centrifuged at 5,000 × g for 15 min at 4°C, and the cell pellet was washed three times with wash buffer (10 mM Tris-HCl [pH 7.5], 250 mM sucrose, and 0.05% Tween 80). Frozen in liquid nitrogen, and stored at −70°C until processing. The frozen pellet was resuspended in a solution containing 3 ml of 20 mM Tris-HCl (pH 7.5), 5 mM 2-mercaptoethanol, 2 mM sodium dihydrogen phosphate, 2 mM sodium orthovanadate, 2 mM phenylmethylsulfonyl fluoride, and 1 mM benzamidine; sonicated; and centrifuged at 100,000 × g for 4 h at 4°C (L8-80 ultracentrifuge; Beckman). The supernatant was separated as the cytosol, and the pellet was kept as the membrane fraction. The pellet fraction was resuspended in lysis buffer, and both fractions were stored at −70°C until use. Western blot analysis was performed using an antibody raised against the GST-MSMEG_3780 catalytic domain at a dilution of 1/5,000 by using enhanced chemiluminescence.

Cloning and characterization of promoters. The genomic region of MSMEG_3780 containing the putative promoter was ampliﬁed by PCR using primers MS3780promScaSpe (5’-TCTGATTCAGTCGCTGAGCAGGA-3’) and MS3780promXbaStu (5’-ACTATGCACTTCGGCTGGCACC-3’). The product was digested with XbaI and SpeI and cloned into plasmid pBKS-PMSMEG_3780, and the sequence of the insert was veriﬁed. The Vibrio harveyi luciferase gene was obtained as a 2.4-kb BamHI-HindIII insert from plasmid pLUX10 (30) (a kind gift of Richard Friedmann, University of Arizona) and cloned into BamHI-HindIII-digested vector pBKS-II(luxAB) (31) (a kind gift of Richard Friedmann, University of Arizona) and cloned into BamHI-HindIII-digested vector pBKS-II(luxAB) to generate plasmid pBKS-luxAB. The MSMEG_3780 promoter (obtained as an XbaI-XbaI insert from plasmid pBKS-PMSMEG_3780) and the luciferase gene (luxAB, obtained as a 2.4-kb BamHI-HindIII insert from plasmid pBKS-luxAB) were cloned into XhoI-HindIII-digested vector pMV-10-25 (8) to generate plasmid pMV-PMSMEG_3780-luxAB. Primers MS3780promHindIII-trunc (5’-GGGGCCGAAAACGTCTGACCCGA CGATTG-3’) and MS3780promXbaStu (5’-ACTATGCACTTCGGCTGACCCCAACCC-3’) were used to amplify genomic DNA to ±250 bp of the annotated MSMEG_3780 gene from M. smegmatis genomic DNA. The amplicon was digested with XbaI and XhoI and cloned into similarly digested plasmid pBKS-PMSMEG_3780, and the sequence was veriﬁed. The KpnI-SpeI fragment from pBKS-PMSMEG_3780 truncated and the SpeI-HindIII fragment containing luxAB from plasmid pBKS-luxAB were then cloned into KpnI-HindIII-cutter vector pMV-10-25 to generate pMV-PMSMEG_3780-luxAB.

To generate the shortest promoter fragment, pBKS-PMSMEG_3780 truncated was digested with Smal and HindII to release a 180-bp fragment. The vector was then religated to generate pBKS-PMSMEG_3780_short. A KpnI-XbaI fragment from this plasmid was cloned into KpnI-SpeI-digested pMV-PMSMEG_3780-luxAB to generate plasmid pMV-PMSMEG_3780_short-luxAB. The region encompassing the putative Rv1647 promoter (~400 bp) was ampliﬁed by PCR using primers Rv1647PknFwd (5’-CCGATTGTGTGGATCCCTTCTTCCGTTCCGTT-3’) and Rv1647PknSpeRev (5’-CTCAGGTATCCGATCCGTCCTC-3’) (12) to generate plasmid pBKS-PMSMEG_3780_rvC-luxAB. The region encompassing the putative Rv1647 promoter (~400 bp) was ampliﬁed by PCR using primers Rv1647PknFwd (5’-CCGATTGTGTGGATCCCTTCTTCCGTTCCGTT-3’) and Rv1647PknSpeRev (5’-CTCAGGTATCCGATCCGTCCTC-3’) to generate plasmid pBKS-PMSMEG_3780_rvC-luxAB. The product was digested with XbaI and SpeI and cloned into plasmid pBKS-PMSMEG_3780-luxAB, and the sequence was veriﬁed. The KpnI-SpeI fragment from pBKS-PMSMEG_3780_rvC-luxAB and the SpeI-HindIII fragment containing luxAB from plasmid pBKS-luxAB were then cloned into KpnI-HindIII-cutter vector pMV-10-25 to generate pMV-PMSMEG_3780_rvC-luxAB.

The ireA promoter of approximately 330 bp (12) was ampliﬁed by PCR from genomic DNA using primers MSsigA_Kpnfwd (5’-AGCTGATACGACGTCTCCGGATCCGAGA-3’) and MSsigA_XhorRev (5’-TTGAGCAGCCGCGTCATGTG-3’). The PCR product was digested with KpnI and XhoI and cloned into the similarly cut vector pBKS-II(luxAB) to generate plasmid pBKS-PMSMEG_3780-ireA-luxAB. The luciferase gene was obtained as a 2.4-kb BamHI-HindIII insert from plasmid pBKS-luxAB and cloned into KpnI-XbaI-digested pMV-H94 to generate pMV-H94/luxA. The luciferase gene was obtained as a 2.4-kb BamHI-HindIII insert from plasmid pBKS-luxAB and cloned into KpnI-XbaI-digested pMV-H94 to generate pMV-H94/luxA. This plasmid allowed the expression of Rv1647 under the MSMEG_3780 promoter with a translational fusion to residue 144 (as annotated by TIGR) of the MSMEG_3780 protein. To conﬁrm the integration of the Rv1647 gene, PCR was carried out with primers Rv1647PSpeI allele (5’-TGGCTAACAATGTTCGATCC-3’) on genomic DNA prepared from M. smegmatis wild-type, knockout, and ΔMSMEG_3780RvC strains. The 16S rRNA gene was ampliﬁed from the genomic DNA of the three strains using primers Mc1061f-wfd (5’-CGAGCGGT GTTGCCGGATAAT-3’) and Mc1061s-rvs (5’-TCTACGCTCCTAAAGGAA-3’) to conﬁrm the integrity of the genomic DNA. Southern blot analysis was performed onSacI-digested genomic DNA prepared from the complement strain to conﬁrm the integration of the Rv1647 catalytic domain at the att site. The probe used was a 250-bp fragment released by HindIII-SpeI digestion of pBKS-PMSMEG_3780 truncated.

Membrane fractions were prepared from the complement strain and subjected to Western blot analysis using an afﬁliated anti-ribob body to Rv1647 (35) and a monoclonal antibody to gyrase A characterized previously (23).

RESULTS

Cyclases and cAMP levels in M. smegmatis. M. smegmatis has orthologs of eight cyclases from M. tuberculosis (38), six of which are predicted to be homodimeric adenylyl cyclases (MSMEG_0228, MSMEG_3243, MSMEG_3780, MSMEG_4279, MSMEG_4924, MSMEG_5018, and MSMEG_6154). In addition to a cyclase that is unique to its genome (MSMEG_3243), M. smegmatis has an ATPase domain-containing cyclase (MSMEG_0228) that has an ortholog in M. leprae and M. avium but not in M. tuberculosis. This variety of cyclases with diverse domain fusions suggests that M. smegmatis could serve as a model for studying CAMP-based signaling in mycobacteria with respect to several cyclases that it shares with other pathogenic members.

Given the large number of putative adenylyl cyclases, we decided to measure CAMP levels in M. smegmatis under various conditions. Intracellular CAMP levels in wild-type M. smegmatis were high during the early phases of growth (Fig. 1A). CAMP was also efﬁciently extruded by M. smegmatis into the growth medium, and concentrations of CAMP reaching ~4 μM accumulated in the culture medium of the cells (Fig. 1B) at 24 h and thereafter. The total amount of CAMP in the medium did not decrease during culture, indicating very little degradation of CAMP in the extracellular medium (Fig. 1B). The amount of CAMP found extracellularly appears to be dependent on the intracellular concentration of CAMP. Thus, extracellular levels of CAMP normalized to cell number (Fig. 1B) showed their peak at 24 h, after the intracellular levels had peaked at 12 h (Fig. 1A). Moreover, CAMP levels per million living cells declined extracellularly in the late log and stationary phases, reﬂecting the reduced intracellular concentrations of CAMP at these times. One can only speculate as to the function of these high extracellular levels of CAMP; perhaps mycobacteria use this second messenger to signal to each other, as is seen in Dictostelium (10). Early reports on CAMP in mycobacteria showed no change in CAMP levels during growth of the bacteria when expressed as CAMP concentration per gram (dry weight) (26). We chose
to express our data per viable cell in order to remove artifacts that might arise from altered cell wall composition during growth of the organism, which would contribute to measurements of dry weight and therefore mask changes in cAMP levels in the bacterial cell. Assuming a cellular volume of $1 \mu\text{m}^3$, the molar concentrations of intracellular cAMP are $\sim 3$ mM, which is comparable to levels reached in mammalian cells upon forskolin stimulation of the adenyl cyclases.

We also measured cAMP levels under different stress conditions to investigate if the level of this second messenger is sensitive to environmental conditions that \textit{M. smegmatis} or \textit{M. tuberculosis} could experience. No change in viability was seen under the stress conditions utilized in Fig. 1C. In contrast to the response that has been well studied in \textit{E. coli}, where the cAMP level increases dramatically upon catabolite repression (15), no change in cAMP levels were evident in carbon-starved \textit{M. smegmatis} cells (Fig. 1C). Neither NO nor H$_2$O$_2$ caused a change in intracellular cAMP levels, but acid stress reduced cAMP levels by $\sim 50\%$. Additional stresses were tested, but under these conditions, \textit{M. smegmatis} could tolerate a shorter duration (1.5 h) without a significant loss in viability. A dramatic increase in intracellular cAMP levels was seen in cells treated with SDS, while a reduction in cAMP levels was seen in cells treated with PBS (Fig. 1D).

Given the high levels of cAMP in \textit{M. smegmatis}, we wished to assess the contribution of individual cyclases to the total cAMP levels. The MSMEG\textsubscript{3780} gene is an adenyl cyclase that is present in all mycobacterial genomes, and its orthologs in \textit{M. tuberculosis} and \textit{M. leprae} have been recently biochemically characterized (35). However, their role in the physiology of mycobacteria remains unknown. We describe below the importance of MSMEG\textsubscript{3780} in contributing to intra- and extracellular cAMP levels during growth and other environmental insults.

Characterization of the MSMEG\textsubscript{3780} cyclase. The MSMEG\textsubscript{3780} gene is predicted to encode a protein of 392 amino acids in length (TIGR annotation), with a predicted molecular mass of 42.8 kDa, while annotation by the Sanger Centre suggests that the N-terminal residue of the protein begins at residue 133 by TIGR annotation, which would result in a protein with a molecular mass of $\sim 30$ kDa (Fig. 2A).

In MSMEG\textsubscript{3780}, Asp-212 and Asp-256 are predicted to be metal-binding residues, Asn-313 and Arg-317 are the transition-state stabilizing residues, and Lys-252 and Asp-306 contribute to the substrate-specifying residues, i.e., ATP binding residues (Fig. 2A).

FIG. 1. cAMP levels in \textit{M. smegmatis} during growth and stress conditions. (A) Intracellular cAMP levels during growth of wild-type \textit{M. smegmatis}. (B) Total amount of extracellular cAMP (solid line) present in a culture (total volume, 10 ml). Extracellular cAMP (dotted line) of the same cultures is expressed per $10^8$ cells. C. \textit{M. smegmatis} cells were cultured for 30 h, resuspended in the medium indicated, and cultured for 5 h, following which the intracellular cAMP level was measured. (D) \textit{M. smegmatis} cells were cultured for 30 h, and cells were resuspended in fresh medium and cultured for another 1.5 h at 37°C or at 42°C. Alternatively, cells were resuspended in medium containing 0.05% SDS or in PBS and cultured for 1.5 h. The intracellular cAMP level was measured at the end of 1.5 h. All data shown represent the means $\pm$ standard errors of the means (SEM) of duplicate determinations, with each assay being performed three times.

Similar to observations with Rv1647, full-length MSMEG\textsubscript{3780} (TIGR overexpressed in \textit{E. coli}) was prone to proteolysis (35). Cloning, expression, and purification of the catalytic domain of MSMEG\textsubscript{3780} from residues 149 to 392 showed that the enzyme had significant adenyl cyclase activity in vitro (Fig. 2B). Many of the biochemical properties of MSMEG\textsubscript{3780} were similar to those of Rv1647 (35). For example, the pH optimum for adenyl cyclase activity was pH 7.9, and the activity was also stimulated 1.5- to 2-fold by CHAPS but not by 500 mM NaCl (Fig. 2B), indicating some distinct biochemical properties. The apparent $K_m$ for MnATP ($\sim 600$ $\mu$M) was also comparable to that observed for Rv1647 (Fig. 2C).

To provide further biochemical evidence for the structural and possible functional equivalence of MSMEG\textsubscript{3780} and Rv1647, we performed GST pull-down experiments. We had successfully utilized this heterodimerization assay to understand the oligomeric status of Rv1625c and its mutants (18). As shown in Fig. 2D, GST-MSMEG\textsubscript{3780} could associate not only with itself but also with Rv1647. Similarly, GST-Rv1647 could heterodimerize with MSMEG\textsubscript{3780} and homodimerize with histidine-tagged Rv1647.

Nucleotide cyclases form heterodimers only with structurally similar proteins (15). Therefore, given the shared biochemical properties of Rv1647 and MSMEG\textsubscript{3780}, it appears that a study of MSMEG\textsubscript{3780} not only could provide information on cAMP levels in \textit{M. smegmatis} but also could suggest the function of the Rv1647 ortholog in \textit{M. tuberculosis}. In order to understand the role of MSMEG\textsubscript{3780} further, we generated and characterized a deletion mutant as described below.

Generation and characterization of the \textbf{ΔMSMEG\textsubscript{3780}} strain. The ΔMSMEG\textsubscript{3780} knockout strain, lacking the entire catalytic domain of MSMEG\textsubscript{3780}, was generated with an in-frame deletion in MSMEG\textsubscript{3780} to avoid polar effects on other genes that could be present in an operon with MSMEG\textsubscript{3780}. The knockout strain was also complemented with a genomic insertion at the \textit{attl} site of a fragment encompassing 500 bp of upstream sequences of the annotated MSMEG\textsubscript{3780} gene (to include the putative promoter of MSMEG\textsubscript{3780}) and the full-length gene (Fig. 3A, B, and C). In order to ensure that the transcription of the downstream MSMEG\textsubscript{3779} gene was not compromised in the ΔMSMEG\textsubscript{3780} strain,
reverse transcription-PCR was performed with RNA prepared from the wild-type and knockout strains using primers specific for MSMEG_3779. As shown in Fig. 3D, the expression levels of MSMEG_3779 RNA in the wild-type and the knockout strains were similar, thus indicating the absence of polar effects of the deletion.

The expression of MSMEG_3780 during growth was assessed by Western blot analysis. Western blot analysis of lysates from the wild-type and knockout strains using an antibody against MSMEG_3780 showed a prominent band of ~30 kDa in the membrane fraction of wild-type cells, which was absent in membranes prepared from the ΔMSMEG_3780 strain (Fig. 3E). The band reappeared in the complement strain. Therefore, the protein in M. smegmatis associates with the particulate fraction in cells despite the absence of any predicted transmembrane domain in its coding sequence. The Rv1647 protein in M. tuberculosis was also found in the particulate fraction (35). The activation of the enzymatic activities of these two proteins by detergents suggested a preference for a hydrophobic environment, and perhaps, this is provided by the membrane environment in the cell. Rv1647 could be identified in a proteomic analysis of membrane proteins from M. tuberculosis (14), in agreement with the localization of Rv1647 that we reported previously (35). This also strengthens our

FIG. 2. Expression, purification, and characterization of MSEMG_3786 adenylyl cyclase and interaction with Rv1647. (A) Sequence similarity of the Rv1647 and the MSMEG_3780 proteins. The boxed residue represents the annotated start of MSMEG_3780 according to the Sanger Centre annotation. The predicted amino acid sequences were aligned using ClustalW. Dashed arrows represent substrate-specifying residues; gray arrows represent transition-state stabilizing residues. (B) Purified MSMEG_3780 (~500 ng) was assayed in the presence of either 10 mM Mg or 10 mM Mn as a free metal along with 1 mM ATP. CHAPS (0.5%) or NaCl (500 mM) was added in some assays, along with 10 mM Mn and 1 mM MnATP as a substrate, and assays were carried out at pH 9. The inset shows purified MSMEG_3780 used for adenylyl cyclase assays. Values represent the means ± SEM of duplicate determinations, with each assay being performed three times. (C) MSMEG_3780 activity was assayed in the presence of various concentrations of MnATP in the presence of 10 mM free Mn. The level of cAMP produced was measured by radioimmunoassay. All data shown represent the means ± SEM of triplicate determinations, with each assay being performed twice. (D) A GST pull-down assay was performed as described in Materials and Methods. Hexahistidine-tagged proteins that interacted with the GST fusion protein were detected by Western blot analysis using an anti-His antibody. An equal aliquot of proteins that were taken for interaction was subjected to SDS-polyacrylamide gel electrophoresis, and the gel was stained to ensure that approximately equal amounts of protein were used for the experiment. The result shown is representative of experiments performed twice.
observation that MSMEG_3780 is also located in the membrane (Fig. 3E).

The molecular mass of the protein as expressed in *M. smegmatis* was ~30 kDa, much smaller than the ~42.8 kDa according to TIGR annotation. We previously reported that the mass of Rv1647 in *M. tuberculosis* lysates using an affinity-purified antibody was ~30 kDa (35). Significant similarity between the Rv1647 and MSMEG_3780 proteins starts from residues L43 (Rv1647) (TIGR annotation) and L121 (MSMEG_3780) (TIGR annotation) (Fig. 1A). Proteins expressed from these residues would have a predicted molecular mass of ~29 kDa, close to the molecular mass that was observed in Western blot analyses. As described below, we were indeed able to show that a promoter for MSMEG_3780 lies within the region annotated by TIGR as being within the coding sequence of MSMEG_3780.

The expression of MSMEG_3780 was monitored during growth, and the highest levels of the MSMEG_3780 protein...
were seen at 12 h and 24 h (Fig. 3F), at times when cAMP levels were also found to be high.

The wild-type and ΔMSMEG_3780 strains showed no difference in their growth rates and cell or colony morphologies in vitro. However, we did observe differences in cAMP levels in the two strains. During the early and mid-log phases of growth, levels of cAMP in the ΔMSMEG_3780 strain were significantly lower than those in the wild-type or complement strains (Fig. 4A), indicating that the activity of MSMEG_3780 could contribute to the generation of cAMP at this time. The cAMP levels in the complemented strain were similar to those of wild-type M. smegmatis. Moreover, levels of extracellular cAMP were lower at 12 h and 24 h in the ΔMSMEG_3780 strain, which is reflective of the reduced intracellular levels of cAMP in this strain (Fig. 4A).

We also evaluated the contribution of the MSMEG_3780 gene to cAMP levels during various stresses. A reduction in cAMP levels was seen in both wild-type and knockout strains when cultured in PBS, indicating that cAMP levels under these conditions were regulated by the activities of perhaps a number of adenyl cyclases and not just MSMEG_3780 (Fig. 4B). Interestingly, however, the reduction in intracellular cAMP levels was not seen in the ΔMSMEG_3780 strain upon acid stress (Fig. 4C). This reduction in intracellular cAMP levels was again restored upon complementation of the knockout strain, indicating that the fall in intracellular cAMP levels in acid-stressed bacteria was indeed regulated by MSMEG_3780 (Fig. 4C). The knockout strain grew at rates comparable to those of the wild-type strain over 48 h in acidified medium but with both strains growing slower in acidified medium than in normal medium (data not shown).

The intracellular pH of M. smegmatis remains stable at 6.5 even upon lowering the external pH to 4 (28), suggesting that the decrease in intracellular cAMP production may not be because of a reduction in the catalytic activity of MSMEG_3780, which is poorly active at low pHs. Therefore, we believe that a major factor in reducing intracellular cAMP levels upon acid stress is the decreased transcription of the MSMEG_3780 gene. We therefore decided to identify and study the promoter of MSMEG_3780 and assess the role of transcriptional regulation in controlling total cellular cAMP levels.

**Promoter of the MSMEG_3780 gene.** In order to test if nucleotide sequences within the gene based on the annotation at TIGR (i.e., encoding amino acid residues 1 to 133) harbored a promoter, we cloned a fragment from nucleotides (nt) 5 to 448, according to TIGR annotation, upstream of the Vibrio harveyi luciferase (luxAB) gene (30) (Fig. 5A). Significant luciferase activity could be detected with the cloned region of the MSMEG_3780 gene (Fig. 5B), indicating the presence of a promoter in this region. A shorter construct, from nt 251 to 448, showed promoter activity comparable to that of the longer promoter, while a fragment from nt 361 to 448 showed low luciferase activity equivalent to that of a promoterless construct, thereby defining the 3' end of the promoter (Fig. 5B). We therefore predict that the start codon of MSMEG_3780 could be downstream of nt 251, and indeed, proteins made from any of the residues shown in Fig. 5A would correspond in size to the MSMEG_3780 protein detected by Western blot-
The activity of the MSMEG_3780 promoter was markedly lower than that of sigA or hsp60 (Fig. 5B), suggesting that MSMEG_3780 is a minor protein component in the cell. We monitored the activity of the MSMEG_3780 promoter during growth, and in agreement with the Western blot analysis, promoter activity was highest at 12 h and reduced dramatically thereafter (Fig. 5C). The activity of the sigA promoter remained constant at all stages of growth. Therefore, it appears that MSMEG_3780 could be responsible for the maintenance of cAMP levels in the bacterium in the logarithmic phase of growth. Since we observed that the transcriptional regulation of MSMEG_3780 could contribute to cAMP levels, we also assessed MSMEG_3780 promoter activity during various stresses.

A dramatic reduction in MSMEG_3780 promoter activity was seen upon acid stress, suggesting that the reduced transcription of MSMEG_3780 could account for the reduction in cAMP levels in acidified cultures. MSMEG_3780 promoter activity remained constant under other stress conditions after 5 h, and no changes in sigA promoter activity were seen even under conditions of acid stress (Fig. 5D). MSMEG_3780 promoter activity was reduced (~30%) under both SDS and PBS stresses in 1.5 h, and this reduction was also seen with the sigA promoter (Fig. 5E). Therefore, the increase in intracellular cAMP levels upon SDS treatment could not be correlated with the altered transcription of MSMEG_3780 but could be due to a general modulation of the catalytic activity of adenyl cyclases in the bacteria since many of these proteins are membrane associated (38) and perhaps activated by mild detergents.

The Rv1647 promoter is acid sensitive, and the gene can complement the ΔMSMEG_3780 strain. In the studies described thus far, we have observed changes in the expression levels of MSMEG_3780 that could be correlated with alterations in cAMP levels. In order to assess whether Rv1647 in M. tuberculosis shares these attributes, we studied the functional similarity between the Rv1647 and MSMEG_3780 proteins by generating a ΔMSMEG_3780 strain complemented by the expression of Rv1647 (Fig. 6A and B). Southern blotting confirmed the presence of the Rv1647 gene in the ΔMSMEG_3780 strain (Fig. 5C). The expression of Rv1647 in the complemented strain was seen by Western blot analysis using an antibody to Rv1647 (Fig. 6D). Complementation by the Rv1647 protein in the ΔMSMEG_3780 strain could indeed restore cAMP levels during early logarithmic growth in the strain (Fig. 6E), and acid stress also reduced cAMP levels in the complemented strain (Fig. 6F).
In order to show that Rv1647 gene transcription could be regulated by acid stress, we cloned the putative Rv1647 promoter (encompassing the 107-bp intergenic region between the Rv1646 and Rv1647 genes to 298 bp into the Rv1647 gene) upstream of luciferase and monitored the regulation of its activity under conditions of acid stress. Indeed, the promoter of Rv1647 was also downregulated upon acid stress (Fig. 6G).

In a recent study where a global analysis of genes in M. tuberculosis that were sensitive to pH 5.5 was reported (31), the Rv1647 gene was not found to show a change in mRNA levels. That microarray analysis was performed 2 h after subjecting cells to acid stress, indicating that Rv1647 mRNA could be stable for many hours, as the transcription of the Rv1647 gene is reduced dramatically in as little as 1 h of acid stress (Fig. 6G).

In conclusion, the results presented here have shown that an adenyl cyclase from M. tuberculosis could functionally complement its ortholog in M. smegmatis, thereby validating the use of M. smegmatis as a model system to study the roles of cAMP and adenyl cyclases in mycobacteria.

**DISCUSSION**

A number of questions as to the role of cAMP in mycobacteria remain, and the importance of this second messenger in the physiology and perhaps pathophysiology of these bacteria remains largely speculative (39). Here, we have studied the functional importance of a highly conserved adenyl cyclase gene in mycobacteria and used M. smegmatis as a model system to address questions related to the synthesis of cAMP by MSMEG_3780 and levels of cAMP in M. smegmatis during bacterial growth and under conditions of stress. Under normal growth conditions, levels of cAMP appear to be high in the log phase, and at least some of this cAMP is generated by MSMEG_3780, since cAMP levels at 12 h and 24 h of growth were lower in the ΔMSMEG_3780 strain. The marked reduction in intracellular cAMP levels in the stationary phase (~10% of the levels seen at 24 h of growth) could not be accounted for by a reduction in MSMEG_3780 expression, since levels of cAMP fell even in the knockout strain. This indicates that either a very efficient degradation mechanism is operative in the cell or there is low adenyl cyclase activity at later stages of growth.

We have identified the product of the Rv0805 gene in M. tuberculosis as being a cAMP phosphodiesterase (32, 34), but no ortholog of Rv0805 has been seen in M. smegmatis. Perhaps other cAMP phosphodiesterases in M. smegmatis need to be identified.

The Rv1625c and Rv1264 adenyl cyclase knockouts generated in M. tuberculosis did not show dramatic phenotypic differences from wild-type H37Rv (9, 15), nor did they show...
Reduced virulence. In the latter studies, no measurements of cAMP were made, and therefore, the contribution of these genes to regulating cAMP levels in M. tuberculosis is not known.

Certain stress conditions (PBS, acid stress, and SDS) regulated the levels of intracellular cAMP. Therefore, genes (possibly involved in adaptation to these stresses) that are regulated under these conditions could be cyclic AMP receptor protein (CRP) responsive in mycobacteria. Indeed, the importance of cAMP and CRP in regulating the expression of acid shock proteins in E. coli was reported previously (6), suggesting the possibility that acid-inducible or -repressed genes in mycobacteria could also be regulated by cyclic AMP levels in the cell. It is interesting that the increase in cAMP levels normally associated with carbon starvation in E. coli was not seen in M. smegmatis. The major intrinsic protein family of glycerol facilitators was identified in the M. smegmatis genome, while no members of this family could be detected in the genome of M. tuberculosis (43), and it is likely that these transporters facilitate the uptake of glycerol in M. smegmatis, protecting it from severe carbon stress. There are far fewer genes involved in carbohydrate uptake in M. tuberculosis than in M. smegmatis (4 versus 28, respectively) (43), suggesting that cAMP responses to carbon deprivation could vary in these two bacteria.

It is intriguing that a large amount of cAMP is extruded by M. smegmatis into the culture medium. It was suggested previously that cAMP secreted by pathogenic Mycobacterium microti is responsible for the inhibition of phagosome-lysosome fusion (21), thereby implicating cAMP as being a virulence factor or a “toxin” in mycobacteria. This mechanism of elevating cAMP levels in host cells is distinct from the strategy placed under these conditions could be cyclic AMP receptor protein-like DNA binding protein. J. Bacteriol. 187:7795–7804.


