Evolutionary Adaptation of the Essential tRNA Methyltransferase TrmD to the Signaling Molecule 3′,5′-cAMP in Bacteria*§

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The nucleotide signaling molecule 3′,5′-cyclic adenosine monophosphate (3′,5′-cAMP) plays important physiological roles, ranging from carbon catabolite repression in bacteria to mediating the action of hormones in higher eukaryotes, including human. However, it remains unclear whether 3′,5′-cAMP is universally present in the Firmicutes group of bacteria. We hypothesized that searching for proteins that bind 3′,5′-cAMP might provide new insight into this question. Accordingly, we performed a genome-wide screen and identified the essential \textit{Staphylococcus aureus} tRNA m1G37 methyltransferase enzyme TrmD, which is conserved in all three domains of life as a tight 3′,5′-cAMP-binding protein. TrmD enzymes are known to use S-adenosyl-l-methionine (AdoMet) as substrate; we have shown that 3′,5′-cAMP binds competitively with AdoMet to the \textit{S. aureus} TrmD protein, indicating an overlapping binding site. However, the physiological relevance of this discovery remained unclear, as we were unable to identify a functional adenylate cyclase in \textit{S. aureus} and only detected 2′,3′-cAMP but not 3′,5′-cAMP in cellular extracts. Interestingly, TrmD proteins from \textit{Escherichia coli} and \textit{Mycobacterium tuberculosis}, organisms known to synthesize 3′,5′-cAMP, did not bind this signaling nucleotide. Comparative bioinformatics, mutagenesis, and biochemical analyses revealed that the highly conserved Tyr-86 residue in \textit{E. coli} TrmD is essential to discriminate between 3′,5′-cAMP and the native substrate AdoMet. Combined with a phylogenetic analysis, these results suggest that amino acids in the substrate binding pocket of TrmD underwent an adaptive evolution to accommodate the emergence of adenylate cyclases and thus the signaling molecule 3′,5′-cAMP. Altogether this further indicates that \textit{S. aureus} does not produce 3′,5′-cAMP, which would otherwise competitively inhibit an essential enzyme.

3′,5′-Cyclic adenosine monophosphate (3′,5′-cAMP) is a second messenger molecule found in all three domains of life (1). It is involved in the regulation of a variety of physiological processes ranging from carbon catabolite repression (CCR)³ in bacteria to mediating the action of hormones in eukaryotes (1). CCR exists in most bacteria and describes the phenomenon that certain carbon sources (usually glucose) are preferentially catabolized over other secondary carbon sources. This is achieved through complex positive and negative regulatory transcription networks (2). CCR is well studied in \textit{Escherichia coli}, where 3′,5′-cAMP and the catabolite receptor protein (CRP) together form an active transcriptional factor that activates the expression of genes coding for proteins involved in catabolizing secondary carbon sources when glucose is exhausted (2). 3′,5′-cAMP is synthesized from ATP by adenylate cyclases (ACs), a large family of enzymes with divergent sequence, domain, and structural features (1). So far, six classes of ACs have been reported, with class III enzymes found in all domains of life and classes I, II, and IV only present in bacteria (1). The ACs identified in \textit{Prevotella ruminicola} and \textit{Rhizobium etli} are distinct from the existing families and were proposed to form Class V and VI enzymes, respectively (3, 4). Class I ACs are exemplified by the \textit{E. coli} CyaA enzyme; class II ACs are bacterial toxins most often secreted into eukaryotic host cells where they perturb host cell functions (5). Lastly, class IV ACs are a unique group of proteins only found in bacteria but forming part of a larger protein family called CYTH domain proteins (6, 7). CYTH proteins are an ancient protein family that exists in all three domains of life and are named after the type IV AC CyaB from \textit{Aeromonas hydrophila} and the human thiamine triphosphatase. It has been proposed that these enzymes were originally inorganic triplyphosphatases and subsequently evolved to contain other enzymatic activities such as adenylate cyclase, mRNA triphosphatase, and thiamine triphosphatase activity (6, 7). CYTH proteins contain a characteristic and highly conserved EXEK amino acid motif at their N terminus and have a

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conserved fold with eight β-sheets forming a tunnel-like structure (6). Various other conserved charged amino acids with their side chains projecting into the tunnel have been identified, and these are involved in coordinating the different polyphosphate substrates or are involved in enzyme catalysis (6, 8–10).

Although ACs enzymes are in general widely distributed among bacteria, there is conflicting evidence if 3',5'-cAMP is produced and plays a physiological role in the Firmicutes group of bacteria. Although a bioinformatics analysis performed by Galperin et al. (11) on 555 complete bacterial and archaean proteomes indicated that adenylate cyclase enzymes are absent in the majority of Firmicutes bacteria, including Staphylococcus aureus, a protein corresponding to SACOL1008 of S. aureus strain COL is nevertheless often annotated as adenylate cyclase. However, the predicted cyclase activity of this protein has never been tested. It has also been reported that 3',5'-cAMP is present in Bacillus subtilis when grown under oxygen limitation conditions, and its level was shown to decrease in the presence of nitrate (12, 13). However, in these studies the molecule suggested to be 3',5'-cAMP was identified only through chromatographic methods and its actual chemical structure was never confirmed by other methods, such as mass spectrometry, which is now routinely used. In S. aureus, ArcR (SACOL2653 in strain COL), a member of the CRP/FNR family of bacterial transcriptional regulators, plays a role in mediating catabolite repression by inducing the arginine deiminase operon genes arcABDC under anaerobic conditions (14). Furthermore, 3',5'-cAMP was shown in in vitro assays to enhance the ability of ArcR to bind to the promoter region of the ictE gene, coding for an L-lactate dehydrogenase (14). However, it was never tested if 3',5'-cAMP is actually present in S. aureus and has a similar effect on ArcR in vivo. In this study we wanted to shed further light on whether or not 3', 5'-cAMP is produced and plays a physiological function in S. aureus. Using an S. aureus ORFeome protein expression library, we screened for 3',5'-cAMP-binding proteins and identified TrmD as a tight 3',5'-cAMP-binding protein. However, we were unable to detect 3',5'-cAMP under various growth conditions nor a functional adenylate cyclase in S. aureus. TrmD is a highly conserved tRNA methyltransferase and present in all three domains of life (15). It converts Gly-37 into m^G37 by transferring the methyl group from S-adenosylmethionine (AdoMet) to a subset of tRNA species (15, 16). We further found that TrmD proteins from E. coli and Mycobacterium tuberculosis do not bind 3',5'-cAMP. Subsequent bioinformatics and extensive biochemical analyses suggested that 3',5'-cAMP competes with AdoMet for binding, presumably inhibiting the essential function of TrmD in S. aureus. Together with a phylogeny analysis, our data suggest that 3',5'-cAMP is absent in Staphylococcus. Finally, our work also highlights that the emergence of 3',5'-cAMP as a signaling molecule in bacteria required essential evolutionary adaptations of AdoMet-binding proteins such as TrmD.

Results

Identification of the S. aureus TrmD Protein as a 3', 5'-cAMP-binding Protein—There is conflicting evidence whether or not 3',5'-cAMP exists in the Firmicutes group of bacteria. We reasoned that if 3',5'-cAMP is indeed present and functionally relevant in S. aureus, specific 3',5'-cAMP-binding protein(s) must exist. To investigate this, we made use of a S. aureus ORFeome protein expression library and the differential radial capillary action of ligand assay (DRAcALA), a simple and fast method for the detection of small molecule-protein interactions (17–19). The DRAcALA method is based on the principle that free radiolabeled small ligand will diffuse outward once spotted on a nitrocellulose membrane but will stay as a tight spot when bound to a protein (17). In previous studies this assay and the ORFeome protein expression library was successfully used to identify c-di-AMP and ppGpp-binding proteins in S. aureus (18–20). The ORFeome protein expression library is a collection of 2337 E. coli strains allowing for the overproduction of 86% of the annotated S. aureus strain COL proteins as His-MBP-fusion proteins (18–20). To apply this assay to the identification of potential 3',5'-cAMP-binding proteins, the expression of the S. aureus proteins was induced, and E. coli lysates were prepared. Next, radiolabeled [32P]cAMP was synthesized using a C-terminal truncated form of the E. coli adenylate cyclase enzyme CyaA. As assessed by thin layer chromatography (TLC), 97% of the input [α-32P]ATP was converted to [α-32P]cAMP (data not shown). The genome wide DRAcALA screen was subsequently performed as previously described (18–20). Two technical replicates were performed, and the lysate from one strain expressing the S. aureus COL protein SACOL1256 (plate 13 well F05) gave a positive result for 3',5'-cAMP binding (data not shown). SACOL1256 (or SAUSA300_1133 in the USA300 strain FPR3757) codes for the tRNA methyltransferase TrmD, termed from here on out as TrmD_{SA}. To further investigate if TrmD_{SA} can bind 3',5'-cAMP with a physiological relevant affinity, SAUSA300_1133 from the USA300 strain LAC* was cloned into vector pET28b, and the protein was expressed and purified as N-terminally His-tagged fusion protein. DRAcALAs were carried out with [32P]cAMP and serially diluted TrmD_{SA} protein ranging from 200 μM to 1.5 nm and a K_d of 1.97 ± 0.24 μM was determined (Fig. 1A). This binding affinity is in a similar range as the reported K_d of 2 μM for the interaction between the E. coli transcription factor CRP and 3',5'-cAMP (21). To test if the binding is specific to 3',5'-cAMP, an excess of the cold competitor nucleotides 3',5'-cAMP, 3',5'-cGMP, c-di-GMP, and c-di-AMP was added to the binding reaction. This analysis revealed that only cold 3',5'-cAMP but none of the other nucleotides tested could compete for binding with the radiolabeled [32P]cAMP (Fig. 1B). Two forms of cAMP have been detected in cells, 3',5'-cAMP, the classic signaling nucleotide, and 2',3'-cAMP, suggested to be a nucleotide intermediate formed during the RNA degradation process (22, 23). As revealed by competitive binding assays, only the classic signaling molecule 3',5'-cAMP but not 2',3'-cAMP could prevent the binding of radiolabeled 3',5'-cAMP to TrmD_{SA} (Fig. 1B). Taken together, these data show that the S. aureus TrmD protein is able to bind the 3',5'-cAMP signaling nucleotide with high affinity. However, it is also of note that no interaction between 3',5'-cAMP and the S. aureus ArcR protein, a transcription factor with homology to the E. coli CRP protein, could be detected (Fig. 1C).
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3′,5′-cAMP Competitively Binds at the AdoMet Substrate Binding Site of the S. aureus TrmD Protein—TrmD is a highly conserved and essential enzyme and present in nearly all bacteria. TrmD is responsible for methylating the Gly-37 residue at the N1 position in a subset of tRNAs using AdoMet as methyl donor (15). Because 3′,5′-cAMP is chemically similar to AdoMet and both contain an adenine moiety and a ribose ring (Fig. 1D), this raised the possibility that 3′,5′-cAMP binds at the AdoMet substrate binding site of TrmD<sub>SA</sub>. Indeed, as revealed by a competitive binding assay, AdoMet could inhibit in a dose-dependent manner the binding of radiolabeled 3′,5′-cAMP to TrmD<sub>SA</sub> (Fig. 1B). Next, isothermal titration calorimetry (ITC) experiments were performed, and a $K_d$ of 121.4 μM was determined for AdoMet binding to TrmD<sub>SA</sub>. Of note, the TrmD<sub>SA</sub> and AdoMet interaction was determined by ITC and not DRA-CALA, as the latter method can only be used for high affinity binding interactions (low μM $K_d$ or below) and using a radiolabeled ligand. Taken together, these data indicate that 3′,5′-cAMP binds at the same site as the natural substrate AdoMet, consistent with a competitive binding mechanism.

2′,3′-cAMP, but Not 3′,5′-cAMP, Can Be Detected in S. aureus Extracts—Previous work indicated that cAMP is produced in *B. subtilis* when grown without aeration under oxygen limitation conditions (12, 13). However, it should be noted that cAMP production was only assessed using chromatographic methods, and its chemical structure was never confirmed by NMR- or mass spectrometry-based methods (12, 13). Given the fact that we uncovered a 3′,5′-cAMP-binding protein in *S. aureus*, we next set out to determine if and when 3′,5′-cAMP is produced in *S. aureus* using a sensitive mass spectrometry-based method (24). Cytosolic extracts were prepared from the wild-type *S. aureus* strain JE2 and strain NE1299, containing a transposon insertion in SAUSA300_0905, coding for an uncharacterized protein often annotated as adenylate cyclase. The strains were grown in tryptic soy broth (TSB) medium under aerobic or micro-aerobic conditions as well as in B-medium supplemented with either glucose or sucrose as the only carbon source to reflect carbon catabolite repression in *S. aureus* (25). Extracts were prepared from both exponential and stationary phase cultures, and nucleotides were detected by LC-MS/MS as described in Bähre and Kaever (24). Using this sensitive method, 3′,5′-cAMP concentrations can be detected up to a lower limit of 0.412 pmol per sample and also discriminated from 2′,3′-cAMP. Large amounts of 2′,3′-cAMP were detected in all samples (Fig. 2). Normalization based on total protein concentrations revealed higher 2′,3′-cAMP levels in extracts prepared from strains grown in TSB medium under micro-aerobic than under aerobic conditions (Fig. 2). The 2′,3′-cAMP levels were even higher when bacteria were grown in B-medium (Fig. 2). Of note, more 2′,3′-cAMP was also present in extracts prepared from stationary than exponential phase cells, when the bacteria were grown in B-medium supplemented with glucose. Although high levels of 2′,3′-cAMP could be detected in all samples, 3′,5′-cAMP was not detected in any of the extracts, suggesting that *S. aureus* does not produce 3′,5′-cAMP at least under the conditions tested.
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SAUSA300_0905 and Its Homologs Are Distinct from Type IV AC Enzymes—An initial bioinformatics analysis indicated that the predicted S. aureus adenylate cyclase SAUSA300_0905 (USA300 FPR3757 nomenclature) is most closely related to type IV AC enzymes. It is possible that the SAUSA300_0905 protein was not expressed or active under the growth conditions tested, and hence, we were unable to detect 3',5'-cAMP in the S. aureus extracts. To test if SAUSA300_0905 is able to synthesize 3',5'-cAMP in vitro, the protein was expressed and purified from E. coli as the N-terminal His-tag fusion protein. The E. coli CyaA_{2–446} protein was purified and used as positive control. Radiolabeled \[^{32}P\text{ATP}\] was used as the substrate, and reactions were set up in three different buffers, as previously reported for in vitro enzyme assays with the E. coli CyaA (26) or the Yersinia pestis type IV AC CyaB (8). The enzyme reactions were incubated for 1 h or overnight at 37 °C, and the reaction products were subsequently analyzed by TLC. Within 1 h, the E. coli CyaA_{2–446} enzyme converted >91 and 61% of the ATP to cAMP in the Mg\(^{2+}\) - and Mn\(^{2+}\)-containing buffers, respectively; however, none of the \[^{32}P\text{ATP}\] was converted by SAUSA300_0905 (Fig. 3A). Overnight reactions essentially yielded similar results (data not shown). Genes coding for active AC enzymes have previously been identified through their ability to complement an E. coli cyaA mutant strain using simple plate assays, as 3',5'-cAMP-producing E. coli strains appear red or blue on MacConkey- or X-Gal-containing plates, respectively (3, 4, 27). To test if SAUSA300_0905 is able to produce 3',5'-cAMP when expressed in E. coli, SAUSA300_0905 and as a positive control the E. coli cyaA_{2–446} gene were cloned with an N-terminal His tag in vector pBAD33 and expressed under the control of the arabinose-inducible promoter. The resulting plasmids, pBAD33-SAUSA300_0905-His\(_{10}\) and pBAD33-cyaA_{2–446}-His\(_{10}\) and the empty vector pBAD33 as control were introduced into the cyaA mutant E. coli strain DHM1. The transformants were spotted onto LB plate supplemented with 50 μg/ml X-Gal and 0.02% arabinose. Whole cell lysates were prepared as described under “Experimental Procedures,” proteins were separated on a 12% SDS-PAGE gel, and His-tagged proteins were detected using a His-tag-specific antibody.

FIGURE 2. 2',3'-cAMP, but not 3',5'-cAMP, is present in S. aureus. Semi-quantitative measurements of 2',3'-cAMP in extracts prepared from wild-type S. aureus strain JE2 and the isogenic SAUSA300_0905 mutant strain (Δ905). Bacteria were grown in TSB medium with/without agitation (A) or in B-medium supplemented with 25 mM of glucose or sucrose (B). As indicated in the graph, extracts were either prepared from log phase cultures or stationary (stat) phase cultures.

FIGURE 3. SAUSA300_0905 is likely not a genuine adenylate cyclase. A, in vitro adenylate cyclase activity assay. Radiolabeled ATP was incubated as the negative control (NC) in the absence of protein and as the positive control with purified E. coli CyaA(2–446) protein (CyaAEC) or with purified histidine-tagged SAUSA300_0905 protein (0905). The reactions were set up in three different buffer systems as specified under “Experimental Procedures” and incubated at 37 °C for 1 h. Aliquots were separated by TLC, and radiolabeled compounds visualized using phosphorimaging. A representative result of three experiments is shown. B, in vivo adenylate cyclase activity. The empty vector pBAD33 (EV) and plasmids pBAD-cyaA_{2–446}-His\(_{10}\) (CyaAEC) or pBAD-SAUSA300_0905-His\(_{10}\) (0905) were introduced into the cyaA mutant E. coli strain DHM1. The transformants were spotted onto LB plate supplemented with 50 μg/ml X-Gal and 0.02% arabinose. C, detection of CyaAEC-His\(_{10}\) and SAUSA300_0905-His\(_{10}\) by Western blot. E. coli DHM1 containing plasmid pBAD33, pBAD-cyaAEC-His\(_{10}\) or pBAD-SAUSA300_0905-His\(_{10}\) was propagated in LB medium without or with 0.02% arabinose. Whole cell lysates were prepared as described under “Experimental Procedures,” proteins were separated on a 12% SDS-PAGE gel, and His-tagged proteins were detected using a His-tag-specific antibody.
protein similar to bona fide type IV adenylate cyclases. Next, homologs of SAUSA300_0905 were retrieved from the NCBI non-redundant (nr) protein sequence database, and this yielded 998 sequences with a minimum of 30% sequence identity and 60% sequence coverage. To compare SAUSA300_0905 with a genuine type IV adenylate cyclase, CyaB from Y. pestis was used as a query sequence to retrieve its closest homologs from the NCBI nr database. This yielded 562 sequences with a minimum of 30% sequence identity and 60% sequence coverage. But none of the CyaB homologs were from the Firmicutes group of bacteria. Next, multiple sequence alignments were performed within and across the two groups as outlined under “Experimental Procedures.” Key sections of the sequence logo motifs for CyaB homologs (and likely genuine adenylate cyclases (AC)) and SAUSA300_0905 homologs (hypothetical adenylate cyclases (hAC)) are shown above or below the alignment, respectively. The sequences of a few representative proteins are shown, which include: 3N10, the CyaB adenylate cyclase class IV from Y. pestis; ACah, WP_048207795.1, adenylate cyclase from A. hydrophila; ACvf, WP_044367144.1, adenylate cyclase from Vibrio fluvialis; ACpp, CAG22648.1, putative adenylate cyclase CyaB from Photobacterium profundum SS9; hACsa, SAUSA300_0905 from S. aureus; hACbs, KJ42798.1, hypothetical protein UM89_04625 from B. subtilis; hAClm, CCG36369.1, uncharacterized protein Yipk from Listeria monocytogenes LL195; hACsp, CIG15488.1, adenylate cyclase from Streptococcus pneumoniae; hACII, WP_010905301.1, adenylate cyclase from Lactococcus lactis. Green stars indicate residues conserved in both groups of proteins; green stars indicate residues specific for homologs of genuine adenylate cyclases; orange stars indicate amino acids conserved in SAUSA300_0905 and its homologs. B, structural model of the S. aureus protein SAUSA300_0905 predicted with Phyre2 and modeled on bh2851, a putative adenylate cyclase from B. halodurans (PDB code 2GFG). 94% of residues modeled at >90% confidence. Gray sticks indicate residues conserved in both groups of proteins; green and orange sticks indicate highly conserved residues specific for CyaB and SAUSA300_0905 homologs, respectively.

FIGURE 4. SAUSA300_0905 is distinct from genuine type IV adenylate cyclases. A, partial sequence alignment and Logo motifs based on amino acid identity for Y. pestis CyaB and S. aureus SAUSA300_0905 homologs. Homologs of CyaB and SAUSA300_0905 were identified by a BLASTP search, and multiple sequence alignments were performed within and across the two groups as outlined under “Experimental Procedures.” Key sections of the sequence logo motifs for CyaB homologs (and likely genuine adenylate cyclases (AC)) and SAUSA300_0905 homologs (hypothetical adenylate cyclases (hAC)) are shown above or below the alignment, respectively. The sequences of a few representative proteins are shown, which include: 3N10, the CyaB adenylate cyclase class IV from Y. pestis; ACah, WP_048207795.1, adenylate cyclase from A. hydrophila; ACvf, WP_044367144.1, adenylate cyclase from Vibrio fluvialis; ACpp, CAG22648.1, putative adenylate cyclase CyaB from Photobacterium profundum SS9; hACsa, SAUSA300_0905 from S. aureus; hACbs, KJ42798.1, hypothetical protein UM89_04625 from B. subtilis; hAClm, CCG36369.1, uncharacterized protein Yipk from Listeria monocytogenes LL195; hACsp, CIG15488.1, adenylate cyclase from Streptococcus pneumoniae; hACII, WP_010905301.1, adenylate cyclase from Lactococcus lactis. Gray stars indicate residues conserved in both groups of proteins; green stars indicate residues specific for homologs of genuine adenylate cyclases; orange stars indicate amino acids conserved in SAUSA300_0905 and its homologs. B, structural model of the S. aureus protein SAUSA300_0905 predicted with Phyre2 and modeled on bh2851, a putative adenylate cyclase from B. halodurans (PDB code 2GFG). 94% of residues modeled at >90% confidence. Gray sticks indicate residues conserved in both groups of proteins; green and orange sticks indicate highly conserved residues specific for CyaB and SAUSA300_0905 homologs, respectively.
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113, and Glu-136 (green stars in Fig. 4A), which have been shown to be required for adenylate cyclase activity (8), are absent in SAUSA300_0905 and its homologs. On the other hand, a unique and highly conserved DXEEXE motif (yellow stars in Fig. 4A) was identified within the C-terminal region of SAUSA300_0905 and its homologs that is absent from type IV ACs. Phyre2 was then used to generate a structure model of SAUSA300_0905 (using bh2851, a putative adenylate cyclase and purified as N-terminal His-tagged proteins (Fig. 5A). This model was then used in DRaCALAs, the average fraction-bound values and S.D. of four values were plotted, and the curve fitted, and 

FIGURE 5. The TrmD proteins from E. coli and M. tuberculosis did not bind 3′,5′-cAMP. A, Coomassie-stained SDS-PAGE gel. About 7.5 μg of the indicated WT or mutant TrmD protein variants were separated on a 12% SDS-PAGE gel, and proteins were visualized by Coomassie staining. B, binding curve and Kd determination between 3′,5′-cAMP and the E. coli, M. tuberculosis, and S. aureus TrmD proteins. Radiolabeled 3′,5′-cAMP and purified TrmDΔA, TrmDΔC, and TrmDΔMT protein ranging from 1.5 nM to 200 μM were used in DRaCALAs, the average fraction-bound values and S.D. of four values were plotted, and the curve fitted, and 

Tyr-86 is Critical for Discriminating 3′,5′-cAMP from AdoMet in E. coli TrmD—The difference in the capacity of the S. aureus, E. coli, and M. tuberculosis TrmD proteins to bind 3′,5′-cAMP indicates a critical difference in their primary sequences and tertiary structures. To gain further insight into this, each of the TrmD proteins was used to retrieve their close homologs from their respective groups (that is, Firmicutes, \( \gamma \)-Proteobacteria, and Actinobacteria). Multiple sequence alignments were performed individually within each group, and sequence logos were generated using Jalview (Fig. 6A) (28). This analysis revealed three highly conserved motifs termed here A, B, and C, which based on previous structural and functional analysis are known to form the active site and the AdoMet substrate binding pocket in TrmD proteins (Fig. 6B) (29, 30). Among these motifs, the motifs B and C with consensus sequences CG(H/R)YEGAD(E/Q/R) and ExSXDQD(Y/F) VLGGE, respectively, are essentially the same in all three groups of bacteria (Fig. 6A). However, motif A differs significantly, with the consensus sequence YLSPQG in Proteobacteria, VPTPAG in Actinobacteria, and the degenerated consensus sequence (L/Y)/(L/M)XP(Q/A)G in Firmicutes (Fig. 6A). In particular, the first three amino acids of motif A, referred to from here on out as motif A1, show very low conservation in the Firmicutes group of bacteria. To visualize the location of these motifs in a structural context, a multiple sequence alignment was created for the TrmD proteins from all three groups. The alignment was subsequently mapped on the crystal structure of the TrmD protein from Hemophilus influenzae (PDB code 1UAK), a member of the \( \gamma \)-Proteobacteria group and displayed in the Consurf view (31). As expected, the three motifs, including the A1 motif, form the binding pocket for AdoMet on
Tyr-86 in TrmDEC might play a critical role in discriminating and hence would not be available for such a group (Fig. 7). The corresponding 3'-OH group in *S. aureus* TrmD protein (PDB code 3KY7) and the *H. influenzae* TrmD protein (PDB code 1UAK), the latter of which has been crystallized in the presence of AdoMet. As shown in the *H. influenzae* TrmD-AdoMet complex structure, the Tyr residue at the beginning of the A1 motif (corresponding to Tyr-86 in *E. coli* TrmD<sub>EC</sub>) and highly conserved in Proteobacteria, forms hydrogen bonds with the 3'-OH and 2'-OH of the ribose ring in AdoMet (Fig. 7A). The corresponding 3'-OH group in 3',5'-cAMP forms a phosphoester bond with the 5'-phosphate group (Fig. 7A) and hence would not be available for such a hydrogen bond interaction. This observation indicates that Tyr-86 in TrmD<sub>EC</sub> might play a critical role in discriminating AdoMet from 3',5'-cAMP by forming an additional hydrogen bond with AdoMet. Consistent with this idea, a Leu residue is present at the corresponding position in the A1 motif of the *S. aureus* TrmD<sub>SA</sub> protein, which could not form a hydrogen bond with the 3'-OH of AdoMet, and this may allow TrmD<sub>SA</sub> to bind AdoMet as well as 3',5'-cAMP. To test this hypothesis, the LMC amino acid residues of the TrmD<sub>SA</sub> A1 motif were replaced with YLS residues as found in TrmDEC. This variant was expressed and purified as N-terminal His-tagged protein (Fig. 5A). As assessed by DRAcalAs, this variant showed a decreased binding affinity for 3',5'-cAMP (Fig. 7B). Conversely, the Tyr-86 residue in the TrmD<sub>EC</sub> protein was replaced with a Phe (lacking the phenol hydroxyl group of Tyr) or Leu residue as present in TrmD<sub>SA</sub>, yielding the TrmD<sub>EC</sub>Y86F and TrmD<sub>EC</sub>Y86L variants, respectively (Fig. 5A). The TrmD<sub>EC</sub>Y86F variant was still unable to bind 3',5'-cAMP (similar to WT TrmD<sub>EC</sub>); however, the TrmD<sub>EC</sub>Y86L variant had a much increased binding affinity for 3',5'-cAMP (Fig. 7C). These findings were corroborated further by ITC experiments. Because of the weak binding, no *K<sub>d</sub>* values could be determined for the interaction between 3',5'-cAMP and the WT TrmD<sub>EC</sub> protein and the TrmD<sub>EC</sub>Y86F variant; however, the TrmD<sub>EC</sub>Y86L variant was able to bind 3',5'-cAMP with a *K<sub>d</sub>* of 41.2 μM. On the other hand, this variant now had a decreased binding affinity for AdoMet with a *K<sub>d</sub>* of 69.9 μM, whereas the WT TrmD<sub>EC</sub> and the TrmD<sub>EC</sub>Y86F variants had similar and high binding affinities for AdoMet with *K<sub>d</sub>* values of 21.5 μM and 27.9 μM, respectively. Taken together, these data indicate that the highly conserved Tyr-86 residue in *E. coli* TrmD and likely also in other γ-Proteobacteria is important for discriminating between 3',5'-cAMP and AdoMet, preventing the binding of the former, which otherwise could competitively inhibit the enzyme.

**Coevolution of TrmD with the Emergence of the 3',5'-cAMP Signaling Pathway**—The fact that TrmD proteins from bacteria that have been shown to produce 3',5'-cAMP (*E. coli* and *M. tuberculosis*) do not bind 3',5'-cAMP with high affinity, whereas TrmD from *S. aureus* tightly binds 3',5'-cAMP raises the possibility that TrmD has evolved immunity to 3',5'-cAMP in bacteria producing this signaling nucleotide. To test this hypothesis, the phylogeny of TrmD was analyzed and compared with that of adenylate cyclases. To do this we searched for homologs of *S. aureus* TrmD in the 555 complete bacterial and archaeal proteomes used by Galperin et al. (11) (supplemental Table S1). BLASTP queries identified a total of 503 homologs with a large taxonomic distribution. Specifically, homologues are found in one copy in all bacterial phyla (except for *Syntrophobacter fumaroxidans* MPOB that have two TrmD proteins: YP_847151 and YP_847109; supplemental Table S1). Close homologs are absent from Archaea, consistent with previous report that TrmD from Archaea is more similar to eukaryotic counterparts. A phylogenetic tree of TrmD homologs was built (supplemental Fig. S1) as well as all sequences were aligned, and careful inspection of the alignment showed that motifs B and C of the AdoMet binding site are well conserved among all TrmD homologs (supplemental Fig. S2). However, the first three residues of motif A are variable (supplemental Fig. S2). Based on the data of Galperin et al. (11) concerning the presence/absence of adenylate cyclase in bacterial proteomes, there appears to be a positive correlation between the presence or absence of ACs...
Protein Adaption with the Emergence of 3',5'-cAMP (and therefore the ability of bacteria to produce 3',5'-cAMP) and the divergence of the first three residues of A motif in TrmD proteins, especially in γ-Proteobacteria and Firmicutes (Fig. 7 and supplemental Fig. S2). In fact, in the proteomes where an AC is present (such as in E. coli), the first residue of motif A is predominantly a tyrosine/phenylalanine (49.62% of the cases). However, in the proteomes where an AC is absent (such as in S. aureus) the first residue in motif A is in a large number of cases a leucine (46.08% of the cases). In particular, for Firmicutes bacteria, there appears to be a good correlation between the absence and presence of a tyrosine/phenylalanine residue in the first position of TrmD motif A1 and the absence and presence of an AC in an organism (Fig. 7D). Taken together, this indicates an underlying evolutionary pressure and protein evolution to prevent the binding of 3',5'-cAMP to TrmD in organisms utilizing this cyclic-nucleotide as the signaling molecule.

Discussion

Evidence whether or not the signaling nucleotide 3',5'-cAMP is present in many of the well studied Firmicutes such as S. aureus has been elusive. In this study the S. aureus TrmD protein was identified as a 3',5'-cAMP-binding protein, and 3',5'-cAMP seems to bind competitively and with high affinity at the AdoMet substrate binding pocket (Fig. 1, A and B). However, using an LC-MS/MS analysis of bacterial extracts and based on other biochemical assays, we were unable to detect 3',5'-cAMP or an active adenylate cyclase in S. aureus (Figs. 2 and 3). 3',5'-cAMP is thus unlikely to be present in S. aureus. On the other hand, we found that TrmD proteins from the γ-proteobacterium E. coli and the actinobacterium M. tuberculosis do not bind 3',5'-cAMP with high affinity. Subsequent bioinformatics, mutagenesis, and biochemical analyses demonstrated that residue Tyr-86 in E. coli TrmD plays a pivotal role in discriminating between 3',5'-cAMP and the native substrate AdoMet. A further phylogenetic analysis suggests that amino acids in the substrate binding pocket of TrmD underwent an adaptive evolution to accommodate the presence of adenylate cyclases and the 3',5'-cAMP signaling molecule.

Several lines of evidence argue for the absence of 3',5'-cAMP in S. aureus and likely also in many other Firmicutes bacteria. The major function of 3',5'-cAMP in bacteria, the carbon catabolite repression, is executed through a modified pathway

![FIGURE 7. Tyr-86 in E. coli TrmD is critical to discriminate between 3',5'-cAMP and AdoMet. A, comparison of the AdoMet-binding pockets of the S. aureus and H. influenzae TrmD proteins. The apo structures of the S. aureus TrmD (PDB code 3KY7; shown in yellow) was overlaid in PyMOL with the AdoMet-bound structure of the H. influenzae TrmD (PDB 1UAK, shown in gray). A zoomed-in view of the ligand binding pocket is shown in schematic representation, with the AdoMet, Tyr-86 of TrmD E and Leu84 of TrmD A shown as a stick model. Tyr-86 of TrmD E formed two hydrogen bonds with 2'- and 3'-OH of the ribose ring of AdoMet, whereas Leu84 of TrmD A probably cannot. B and C, binding curve and K_d determination between 3',5'-cAMP and WT S. aureus TrmD and the indicated variant (B) and WT E. coli TrmD protein and the indicated variants (C), Radiolabeled cAMP and purified TrmD or the TrmD A-LMC-YLS variant TrmD E or the TrmD E-Y86F and TrmD E-Y86L variants ranging from 1.5 nM to 200 μM were used in DRaCALAs, the average fraction-bound values and S.D. of at least three values were plotted, the curve fitted, and K_d value determined as previously described (17). D, Logo motifs of the TrmD protein A motif amino acids of representative Firmicutes. The TrmD protein sequences from 78 representative Firmicutes, as described in Galperin et al. (11), were retrieved and grouped into sequences from bacteria likely lacking a bona fide adenylate cyclase (no AC (n = 69)) or containing an adenylate cyclase (with AC (n = 9)). The sequences were aligned separately, and logo motifs were generated and displayed in the ClustalX default color scheme. The TrmD A motif was split as described in Fig. 6A into motifs A1 and A2, and amino acid numbers indicated above each logo motif section is based on the S. aureus strain COL TrmD protein sequence.]
in several of the model Firmicutes bacteria (Bacillus, Listeria, and Staphylococcus) (2). Although CRP-like transcription factors are present in Firmicutes, they often lack key residues known to be required for the binding of 3',5'-cAMP (14). As shown in this study, ArcR, a CRP-family transcriptional factor in S. aureus, does not bind 3',5'-cAMP as determined by DRaCALA (Fig. 1C). Using a very sensitive LC-MS/MS-based method, 3',5'-cAMP could not be detected in S. aureus extracts prepared from cultures grown under different conditions including micro-aerobic and catabolite repression conditions, where 3',5'-cAMP was believed to be produced (Fig. 2) (12, 13). In vitro and in vivo experiments on the predicted adenylate cyclase SAUSA300_0905 (SACOL1008) enzyme indicated that this protein is not a genuine adenylate cyclase as this protein does not hydrolyze ATP nor produce 3',5'-cAMP (Fig. 3, A and B). Close homologues to SAUSA300_0905 are found in many other bacteria belonging to the Firmicutes group. Despite the fact that this protein likely has a similar overall fold as type IV ACs, we identified in our bioinformatics analysis distinct sequence features in SAUSA300_0905 and its homologs compared with bona fide Class IV ACs (Fig. 4A). The physiological function of SAUSA300_0905, its enzymatic activity, and substrate specificity remain to be studied.

A genome wide DRaCALA screen for 3',5'-cAMP-binding proteins using an S. aureus ORFeome library identified the essential tRNA methyltransferase TrmD.sa as 3',5'-cAMP-binding protein (Fig. 1A). 3',5'-cAMP binds with high affinity to TrmD.sa and competes for the binding with the native substrate AdoMet (Fig. 1A). The difference in binding affinity is indicated by the limited capacity of AdoMet to compete with 3',5'-cAMP for binding to TrmD.sa (Fig. 1B). Given that TrmD binds its substrate tRNA species in an AdoMet-dependent manner (16), this difference in binding affinity suggests a potential inhibitory effect on the essential function of TrmD would 3',5'-cAMP be present in S. aureus. As shown in this work, a tyrosine residue found at position 86 in the E. coli TrmD and highly conserved among TrmD proteins from γ-Proteobacteria, aids in the discrimination and preferential binding of AdoMet over 3',5'-cAMP in E. coli (Figs. 6A and 7). This residue is absent in S. aureus and most other Firmicutes (Figs. 6A and 7A). However, some non-type Firmicutes strains, which are predicted to encode a genuine adenylate cyclase (11), such as Clostridium acetobutylicum ATCC 824, Clostridium perfringens str. 13, Caldicellulosiruptor saccharolyticus DSM 8903, Desulfotomaculum reducens MI-1, Natranaerobius thermophilus JW/NM-WN-LF, and their sub-strains, contain a Tyr or Phe residue at this position, suggesting a coupling of the presence of a functional adenylate cyclase and 3',5'-cAMP production with the presence of a Tyr/Phe amino acid reside at this position (Fig. 7D). Altogether the findings presented in this study strongly suggest that 3',5'-cAMP and a functional adenylate cyclase enzyme are absent in S. aureus and, although not all, likely also a large number of other bacteria belonging to the Firmicutes group.

TrmD is a highly conserved tRNA methyltransferase found in all three domains of life that converts Gly-37 into m1G37 by transferring the methyl group from AdoMet to a subset of tRNAs species (15, 32). This modification on tRNAs is essential for maintaining the correct reading frame during protein translation (15, 33). Abolishing the function of TrmD increases +1 frameshift events during protein translation, and growth defects have been observed in its absence in bacteria and yeast (15, 33–35). To bind the AdoMet substrate, TrmD proteins assumes a particular protein fold composed of a deep trefoil knot that is a characteristic of SPOUT family RNA methyltransferases (MTases). SPOUT family MTases methylate ribosomal RNAs or tRNAs on a base or ribose ring using AdoMet as the methyl donor (36). The conserved AdoMet binding site and unique protein fold of SPOUT MTases raises the question as to why other MTases from S. aureus were not identified as 3',5'-cAMP-binding proteins. Several MTases have been crystallized in complex with AdoMet, its analogue sinefungin, or S-adonetyl-l-homocysteine (AdoHcy) (29, 37–39). Careful inspection of the substrate binding sites of these other MTases, namely the tRNA MTases TrmL (PDB code 1MXI), TrmH (PDB code 1V2X), MjNep1 (PDB code 3BBH), and the RNA MTases ScNep1 (PDB code 2V3K) and RsmE (PDB code 2Z0Y) (Fig. 8), revealed that in these cases a main chain peptidyl amine of a highly conserved glycine residue forms hydrogen bonds with the 2'- and 3'-OH of AdoMet/AdoHcy/sinefungin. This difference might explain why only TrmD(sa) was found to interact with 3',5'-cAMP, as other MTases in S. aureus are likely able to discriminate between 3',5'-cAMP and AdoMet.

In γ-Proteobacteria and Firmicutes, we found bioinformatically and experimentally a good correlation between the ability of TrmD proteins to discriminate between 3',5'-cAMP and AdoMet and the presence of a tyrosine or phenylalanine residue at a position equivalent to position 86 in the E. coli TrmD protein (Figs. 6 and 7). These data suggest that TrmD proteins in bacteria producing 3',5'-cAMP adapted to confer “immunity” to 3',5'-cAMP, binding of which would otherwise inhibit the essential function of TrmD. However, this simple correlation was not obvious in all groups of bacteria. In particular, we found that TrmD proteins from the Actinobacteria group had a unique set of highly conserved amino acid residues VPT in the motif A1 (Fig. 6A). Considering the numerous adenylate cyclases reported in this group of bacteria (up to 16 ACs in M. tuberculosis) (1), it is possible that the TrmD proteins underwent a global whole-protein adaption and optimization to properly function in the unique physiological conditions imposed by these ACs. It remains to be investigated how these “unusual” TrmD proteins cope with the presence of physiological concentrations of 3',5'-cAMP. Such studies could also have translational impact, as TrmD proteins from bacteria are distinct from those found in Eukaryotes and Archaea and thus remain attractive drug targets.

TrmD has been extensively studied, especially in pathogenic bacteria, where a number of biochemical and structural characterization were performed (16, 29) in order to pinpoint unique features that are drug-able. Indeed, the TrmD<sub>MT</sub> protein was highlighted as an excellent drug target candidate in M. tuberculosis (40), and screens for small molecule TrmD enzyme inhibitor were performed (41, 42). Lahoud et al. (42) found that adenosine and methionine fragments of AdoMet preferentially inhibit bacterial TrmD proteins over those of eukaryotic-Archaea origins. Hill et al. (41) screened a large col-
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**Experimental Procedures**

**Bacterial Growth Conditions**— *E. coli* strains were grown in Luria Bertani (LB) LB-M9 medium (49.3 mM Na₂HPO₄, 14.7 mM KH₂PO₄, 8.55 mM NaCl, 18.7 mM NH₄Cl, 3.7 mM sodium succinate, 11.1 mM glucose, 2 mM MgSO₄, 1% Tryptone, and 0.5% yeast extract) was used for the growth of the *E. coli* S. aureus ORFeome library strains. S. aureus strains were grown in TSB medium or B-medium (1% peptone, 0.5% yeast extract) supplemented with 25 mM glucose or sucrose (25). For micro-aerobic growth of *S. aureus*, cells were inoculated into 50-ml Falcon tubes filled up completely with TSB medium and incubated at 37°C without agitation (43). Micro-aerobic growth was validated by adding 0.001% resazurin (Sigma) to the cultures as a redox indicator. When appropriate, cultures were supplemented with antibiotics indicated in Table 1.

**Strain and Plasmid Constructions**—Bacterial strains and primers used in this study are listed in Tables 1 and 2, respectively. For construction of plasmids pET28b-His-trmD_EC and pET28b-His-trmD_SA primer pairs, ANG1918/1919 and ANG1920/1921 were used to amplify the trmD genes using *E. coli* MG1655 or *S. aureus* LAC* chromosomal DNA as template. The PCR products were digested with NheI/EcoRI and ligated with plasmid pET28b that has been cut with the same enzymes. For construction of plasmids pET28b-His-trmD₁,c(Y86F) and pET28b-His-trmD₁,c(Y86L), primer pairs ANG1918/2152 and ANG1919/2153 primer pairs ANG1918/2154 and ANG1919/2155 were used in the first round of PCR. The resulting products were gel-purified and fused using primer pair ANG1918/1919. The final product was digested with NheI and EcoRI and ligated with plasmid pET28b that had been cut with the same enzymes. Plasmid pET28b-His-trmD₁,c(LMC-YLS) was constructed in a similar manner using primers ANG1920/2156 and ANG1921/2157 in the first PCR and primers ANG1920/1921 in the second PCR. The final product was digested with enzymes NheI and EcoRI and ligated with plasmid pET28b. For construction of plasmid pET23b-His-trmD_TRP Primer pair ANG2158/2159 and *M. tuberculosis*...
TABLE 1

Bacterial strains used in this study

| Strain | Relevant features | Reference |
|--------|------------------|-----------|
| Escherichia coli strains | | |
| XL1 Blue Cloning strain, TetR, ANG127 | | Stratagene |
| BL21(DE3) E. coli strain used for protein expression, ANG191 | | Novagen |
| DHDM1 cyaA mutant E. coli strain, ANG3110 | | (53) |
| ANG1824 XL1 Blue pET28b, KanR | | Novagen |
| ANG1867 BL21(DE3) pET28b, KanR | | Novagen |
| ANG3168 BL21(DE3) pET28b-His, cyaEc, (2–446); KanR | | This study |
| ANG3412 DH5α pBAD33, CamR | | (54) |
| ANG3813 XL1 Blue pBAD33-cyaAc, (2–446); His6, CamR | | This study |
| ANG3414 XL1 Blue pBAD33-SAU300_0905, His6, CamR | | This study |
| ANG3434 DHM1 pBAD33-cyaAc, (2–446); His6, CamR | | This study |
| ANG3435 DHM1 pBAD33-SAU300_0905, His6, CamR | | This study |
| ANG3436 DHM1 pBAD33, CamR | | This study |
| ANG3290 T7IQ pVL791-His6-SACOL1008; CamR, Carbr | | This study |
| ANG3650 BL21(DE3) pET28b-His, trmDTB; KanR | | This study |
| ANG3851 BL21(DE3) pET28b-His, trmDTB; KanR | | This study |
| ANG3682 XL1 Blue pET28b-His, trmDTB; Y66F, KanR | | This study |
| ANG3683 XL1 Blue pET28b-His, trmDTB; Y66F, KanR | | This study |
| ANG3864 XL1 Blue pET28b-His, trmDTB; LMC-YLS; KanR | | This study |
| ANG3685 XL1 Blue pET28b-His, trmDTB; KanR | | This study |
| ANG3686 BL21(DE3) pET28b-His, trmDTB; Y86F, KanR | | This study |
| ANG3687 BL21(DE3) pET28b-His, trmDTB; Y86F, KanR | | This study |
| ANG3918 BL21(DE3) pET28b-His, trmDTB; LMC-YLS; KanR | | This study |
| ANG3688 BL21(DE3) pET28b-His, trmDTB; KanR | | This study |
| ANG3689 BL21(DE3) pET28b-His, trmDTB; KanR | | This study |
| ANG3895 T7IQ pVL791-His6, SACOL2653; CamR, Carbr | | This study |
| ANG3896 XL1 Blue pET28b-His, cyaAc, (2–446); KanR | | This study |
| ANG4001 XL1 Blue pET28b-His, trmDTB; KanR | | This study |
| ANG4002 XL1 Blue pET28b-His, trmDTB; KanR | | This study |
| S. aureus strains | | |
| JE1 USA300 MRSA strain (ANG2624) | | (55) |
| NE1299 JE2 SAUSA300_0905-Tn; Ernr (ANG3894) | | (55) |

H37Rv chromosomal DNA was used to amplify the trmDB gene. The PCR product was digested with Ndel and HindIII and ligated with plasmid pET23b that had been cut with the same enzymes. For construction of plasmid pET28b-HisS-cyaAcEC (2–446) primer pair, ANG1788/1789 and E. coli MG1655 chromosomal DNA were used. The PCR product was digested with Ndel and EcoRI and ligated with plasmid pET28b. For construction of plasmids pBAD33-cyaAcEC (2–446)–His6 and pBAD33-SAU300_0905–His6 primer pairs ANG2003/2004 and ANG2005/2006 were used to amplify the cyaA or SAUSA300_0905 gene using E. coli MG1655 and S. aureus LAC* chromosomal DNA, respectively. The PCR products were digested with Kpnl and Xbal and ligated with plasmid pBAD33 that has been cut with the same enzymes. All plasmids were initially recovered in E. coli strain XL-1 Blue, and sequences of inserts were confirmed by fluorescent automated sequencing (GaTc Biotech). For protein expression and purification, the plasmids were introduced in E. coli strain BL21(DE3), yielding the strains as specified in Table 1.

Protein Expression and Purification—E. coli BL21(DE3) strains (Table 1) were used for the expression and purification of His-CyaAcEC (2–446), the His-tagged TrmD proteins, and its variants. One liter of cultures of the different strains were grown at 37 °C to an A600 of 0.5–0.7, protein expression was induced with 0.5 mM isopropyl 1-thio-β-D-galactopiranoside, and cultures were incubated overnight at 18 °C. For purification of His6-SACOL1008 and His6-SACOL2653, strain T7IQ containing plasmids pVL791-His6-SACOL1008 and pVL791-His6-SACOL2653 were grown at 30 °C overnight. The next day the cultures were diluted 1:50 into 1 liter of fresh LB medium and incubated for 4 h at 30 °C, and subsequently protein expression was induced with 1 mM isopropyl 1-thio-β-D-galactopyranoside for 4 h. Protein purifications by nickel-nitrilotriacetic acid affinity chromatography and size exclusion chromatography were performed as previously described (44). Protein containing fractions were pooled and concentrated to ~10 mg/ml using 10-kDa cut-off centrifugal filters. Protein concentrations were determined using the BCA protein assay kit from Pierce. The purity of the purified proteins was assessed on Coomassie-stained gels after separation of the indicated amounts of protein on 12% SDS-PAGE gels.

Synthesis of 32P 3',5'-cAMP—32P-Labeled 3',5'-cAMP was synthesized from [α-32P]ATP (PerkinElmer Life Sciences) by incubating 55.5 nm [α-32P]ATP with 20 μM purified His-CyaAcEC (2–446) protein in 40 mM Tris, pH 7.5, 100 mM NaCl, 10 mM MgCl2 buffer overnight at 37 °C. The sample was subsequently incubated for 10 min at 95 °C, the denatured His-CyaAcEC (2–446) protein was removed by centrifugation at 17,000 × g for 5 min, and the supernatant was transferred to a new tube and stored at −20 °C. The conversion of [α-32P]ATP to [α-32P]cAMP was estimated to be at least 97%, as assessed by TLC using H2O/ethanol/ammonium bicarbonate (30:70:0.2 m) as the mobile phase and densitometry analysis, which was performed as described previously (18).

Differential Radial Capillary Action of Ligand Assay and Screen for 3',5'-cAMP-binding Proteins—An S. aureus ORFeome library allowing the expression of 2337 N-terminally His-MBP-tagged S. aureus strain COL proteins in E. coli was utilized to identify potential 3',5'-cAMP target proteins. The construction and use of this ORFeome library has been described previously (19, 20). Protein expression and the preparation of whole cell lysates and the subsequent DRaCALA were performed as previously described, with the modification that 32P-labeled 3',5'-cAMP was used as the nucleotide ligand (17–19). For the determination of Kd values by DRaCALA, 2-fold serial dilutions of purified His-TrmD5A, His-TrmD5C, His-TrmD5V, or the different variants were prepared in binding buffer (40 mM Tris, pH 7.5, 100 mM NaCl, 10 mM MgCl2) starting at a concentration of 200 μM and subsequently mixed with ~2 nm 32P-labeled 3',5'-cAMP. The mixtures were incubated for 5 min at room temperature before spotting 2.5 μl of the reactions on nitrocellulose membranes (Amersham Biosciences Hybond-EC; GE Healthcare). The fraction of ligand bound and Kd values were calculated as previously described (17). For nucleotide competition assays, the specified purified protein at a final concentration of 100 μM was incubated with ~2 nm 32P-labeled 3',5'-cAMP in the presence of 400 μM concentrations of the competitor nucleotides AdoMet, 3',5'-cAMP, 3',5'-cGMP, c-di-AMP, c-di-GMP, ATP, ADP, AMP, or 2',3'-cAMP. The reactions were incubated for 5 min at room temperature, 2.5 μl was spotted onto nitrocellulose membranes, and fraction-bound values were determined as described above.

In Vitro Adenylate Cyclase Assay—The purified His6-SACOL1008 protein was tested for potential adenylate cyclase activity using previously described in vitro assay systems (8, 26). The purified His6-CyaAC (2–446) protein was used as the positive control, and enzyme assays were set up in three different buffer systems. Buffer 1 consisted of 40 mM Tris, pH 7.5, 100 mM NaCl, and 10 mM MgCl2 and was previously used to measure the activity of the E. coli CyaA enzyme (26). Buffer 2 con-
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TABLE 2
Primes used in this study

| Number | Name                  | Sequence                      |
|--------|-----------------------|-------------------------------|
| ANG1918 | 5-Nhel-MG1655 TrmD   | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG1919 | 3-EcoRI-MG1655 TrmD  | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG1920 | 5-Nhel-SACOL1256 TrmD| CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG1921 | 3-EcoRI-SACOL1256 TrmD | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG2152 | EcTrmDY86F-R          | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG2153 | EcTrmDY86F-F          | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG2154 | EcTrmDY86L-F          | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG2155 | EcTrmDY86L-F          | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG2156 | SaTrmDLMC-YLS-R       | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG2157 | SaTrmDLMC-YLS-F       | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG2158 | Rv2906c Forward       | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG2159 | Rv2906c Reverse       | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG2178 | 5-Nhel-CyaA (2–446)   | CTAGGCTAGCTGATAGGGTCTTGTGTTG |
| ANG1789 | 3-EcoRI-CyaA (2–446)  | CTAGGCTAGCTGATAGGGTCTTGTGTTG |

was added to the cultures at a final concentration of 0.001% w/v as previously described (43). The reduction potential of resazurin at pH 7.0 and 25 °C is +380 mV, sitting between oxygen gas (+820 mV) and cytochromes (+290 to +80). This makes this compound suitable for detecting aerobic respiration activity or lack thereof, as the former would lead to the reduction of resazurin to resorufin and a blue to pink color change (45). Next, bacteria were collected by centrifugation, and the pellets were suspended immediately in 1 ml of nucleotide extraction buffer containing acetonitrile-methanol-water (2:2:1, v/v) and heated for 15 min at 95 °C to minimize the effect of oxygen on the cell physiology and metabolites. Six samples were prepared for each culture condition for strains JE2 and JE2 Tn::SAUSA300_0905. For three of these samples the extraction buffer was spiced with 92.8 ng/ml isotope-labeled 3',5'-cAMP. To generate the isotope-labeled 3',5'-cAMP, 5 mM 13C,15N-ATP was converted into 13C,15N-cAMP with 5 μM E. coli CyaA(2–446) in 40 mM Tris, pH 7.5, 100 mM NaCl, and 10 mM MgCl2 buffer, and the sample was incubated at 37 °C overnight. The conversion rate was determined as 93.4% by LC-MS/MS analysis. S. aureus extracts were prepared, and 3',5'-cAMP and 2',3'-cAMP was detected by LC-MS/MS as described previously (24).

Preparation of Cell Extract and Detection of Nucleotides by LC-MS/MS—S. aureus strains JE2 and JE2 Tn::SAUSA300_0905 (NE1299, ANG3894) were grown overnight in TSB medium as well as in B-medium supplemented with 25 mM concentrations of either glucose or sucrose (for stationary phase samples). The next day cultures were also back-diluted 1:50 to fresh medium and incubated for 3 h at 37 °C (for exponential phase samples). Bacterial cells corresponding to a 1-ml sample were collected by centrifugation, and the pellets were suspended immediately in 1 ml of nucleotide extraction buffer containing acetonitrile-methanol-water (2:2:1, v/v) and heated for 15 min at 95 °C to minimize the effect of oxygen on the cell physiology and metabolites. Six samples were prepared for each culture condition for strains JE2 and JE2 Tn::SAUSA300_0905. For three of these samples the extraction buffer was spiced with 92.8 ng/ml isotope-labeled 3',5'-cAMP. To generate the isotope-labeled 3',5'-cAMP, 5 mM 13C,15N-ATP was converted into 13C,15N-cAMP with 5 μM E. coli CyaA(2–446) in 40 mM Tris, pH 7.5, 100 mM NaCl, and 10 mM MgCl2 buffer, and the sample was incubated at 37 °C overnight. The conversion rate was determined as 93.4% by LC-MS/MS analysis. S. aureus extracts were prepared, and 3',5'-cAMP and 2',3'-cAMP was detected by LC-MS/MS as described previously (24).

Kd Determination by ITC—A MicroCal iTC200 instrument (GE Healthcare) was used to determine the dissociation constants of 3',5'-cAMP or AdoMet and the E. coli or S. aureus wild-type TrmD proteins or TrmD variants. To minimize the dilution effect, the purified TrmD proteins were dialyzed for 24 h at 4 °C against 4 liters of binding buffer (40 mM Tris, pH 7.5, 100 mM NaCl, 10 mM MgCl2, 5% v/v glycerol). Subsequently, the samples were spun down at 17,000 × g at 4 °C for 10 min to remove any insoluble material, the supernatant was transferred to new tubes, and the protein concentrations were measured using a BCA assay kit (Pierce). An aliquot of the dialysis buffer was used to make 1 mM 3',5'-cAMP and AdoMet solutions and also used to set the TrmD proteins to a concentration of 100 μM. After initial trials, the MicroCal iTC200 was set to a reference power of 6 μcal/s, a stirring speed of 500 rpm, and a temperature of 25 °C, and 20 injections were made at 180-s intervals. At least two technical replicates were performed with each
TrmD protein. As the negative control, the ligands were titrated against the dialysis buffer, and the obtained values were subtracted from the experimental data. Curve-fitting, data analysis, and $K_d$ calculations were performed using the Origin program.

**Western Blot**—The expression of SAUSA300_0905-His$_6$ and CyaA$_{E/a}(2-446)$-His$_6$ from the pBAD33 vectors in DH1 was confirmed by Western blot. Briefly, cultures of strain DH1 containing the different pBAD33-derived vectors were grown overnight in LB medium at 30 °C. The next day the overnight cultures were induced with 0.02% arabinose for 3 h. Bacteria from 1-ml culture aliquots before and after the induction were collected by centrifugation, and cells were suspended in 1× SDS sample buffer to get a final $A_{600 \text{ nm}}$ of 40. Samples were heated for 10 min at 95 °C, and proteins separated on a 12% SDS-PAGE gel. Proteins were then transferred to a PVDF membrane, and His-tagged proteins were detected using a monoclonal anti-poly-His-peroxidase antibody (Sigma A-7058).

**Sequence and Structure Analysis**—Homologs of the *S. aureus* protein SAUSA300_0905, the *Y. pestis* CyaB protein, and the TrmD proteins from *E. coli*, *S. aureus*, and *M. tuberculosis* proteins were identified as follows. For SAUSA300_0905 and CyaB from *Y. pestis*, the respective protein sequences were used as query sequences in BLASTP searches in the NCBI non-redundant (nr) protein sequence database using default settings. For the TrmD proteins, the respective protein sequences were used as query sequence in a BLASTP search confined to their respective groups of bacteria, namely *E. coli* TrmD for γ-Proteobacteria, *M. tuberculosis* TrmD for Actinobacteria, and *S. aureus* TrmD for Firmicutes. Subsequently, the identified protein homologs with a maximum expect (e) values below 3e-04 and a minimum sequence identity of 60 and 30%, respectively, were retrieved and used for further analysis in Jalview (28). A multiple sequence alignment was generated for each group of proteins by running 20 iterations, and a conserved logo-sequence was generated with Cluster Omega (46). To compare the sequences of the SAUSA300_0905 and CyaB homologs or the *E. coli*, *S. aureus*, and *M. tuberculosis* TrmD homologs, the respective groups of sequences were combined, and a multi-sequence alignment and/or a logo-sequence was generated as described above. A multiple sequence alignment of all TrmD proteins was also generated and subsequently used as the input sequence on the ConSurf server to visualize the AdoMet binding site in a structural context (31). To this end, chain A of the AdoMet-bound *H. influenzae* TrmD protein was used as the structural template (PDB code 1UAK). PyMOL (v1.7.4.4 Edu Enhanced for Mac OS X, Schrödinger, LLC.) was used to display the ConSurf data and also for the structural comparison of the AdoMet binding site of the *S. aureus* TrmD (PDB code 3KY7) and *H. influenzae* TrmD (PDB code 1UAK) proteins. A structural model of SAUSA300_0905 was generated in Phyre2 (47) and viewed in PyMOL.

**Phylogeny Analysis of TrmD and Adenylate Cyclases**—A local protein database containing the 555 complete bacterial and archaeal proteomes used by Galperin et al. (11) in his study on the distribution of bacterial signal transduction systems was built. This database was queried with the BLASTP program (default parameters; Ref. 48) using the full-length sequence of TrmD protein of *S. aureus* strain N315 as a seed (Ref_seq: NP_374356, Locus_tag: SA1083). The distinction between homologous and non-homologous sequences was assessed by visual inspection of the BLASTP output (no arbitrary cut-offs on E-value or score). To ensure the exhaustive sampling of homologs, iterative BLASTP queries were performed using homologs identified at each step as new seeds. The absence of a homolog in any complete proteome in the local database was systematically verified by TBLASTN queries against the nucleotide sequence of the corresponding genome. For each candidate protein, the retrieved homologs were added to the dataset. The retrieved sequences were aligned using MAFFT v7.045b (default parameters; Ref. 49). Regions where the homology between amino acid positions was doubtful were removed using the BMGE software (BLOSUM30 option; Ref. 50). Bayesian analyses were performed using MrBayes version 3.2.2 (51) with a mixed model of amino acid substitution including a gamma distribution (4 discrete categories) and an estimated proportion of invariant sites. MrBayes was run with 4 chains for 1 million generations, and trees were sampled every 100 generations. To construct the consensus tree, the first 2000 trees were discarded as “burn in” (51). The Sequence-logos of TrmD the alignments were generated using Phylo-mLogo visualization tool to highlight the three motifs involved in the AdoMet binding (52).

**Author Contributions**—Y. Z. and A. G. designed the study, acquired funding, and wrote the manuscript. Y. Z. and L. E. B. acquired the experimental data, Y. Z. and R. A. performed the bioinformatics analyses, Y. Z., R. A., L. E. B., J. C., V. K., and A. G. analyzed the data.

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