

Structural Basis for Binding of RNA and Cofactor by a KsgA Methyltransferase

Chao Tu,¹ Joseph E. Tropea,¹ Brian P. Austin,¹ Donald L. Court,¹ David S. Waugh,¹ and Xinhua Ji^{1,*}

¹Center for Cancer Research, National Cancer Institute, National Institutes of Health, Frederick, MD 21702, USA

*Correspondence: jix@ncicrf.gov

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SUMMARY

Among methyltransferases, KsgA and the reaction it catalyzes are conserved throughout evolution. However, the specifics of substrate recognition by the enzyme remain unknown. Here we report structures of *Aquifex aeolicus* KsgA, in its ligand-free form, in complex with RNA, and in complex with both RNA and S-adenosylhomocysteine (SAH, reaction product of cofactor S-adenosylmethionine), revealing critical structural information on KsgA-RNA and KsgA-SAH interactions. Moreover, the structures show how conformational changes that occur upon RNA binding create the cofactor-binding site. There are nine conserved functional motifs (motifs I–VIII and X) in KsgA. Prior to RNA binding, motifs I and VIII are flexible, each exhibiting two distinct conformations. Upon RNA binding, the two motifs become stabilized in one of these conformations, which is compatible with the binding of SAH. Motif X, which is also stabilized upon RNA binding, is directly involved in the binding of SAH.

INTRODUCTION

Ribosome biogenesis has been a subject of great interest for decades. Many nucleotide modifications have been found in ribosomal RNA (rRNA), among which the dimethylation of residues A1518 and A1519 in helix 45, the terminal helix near the 3' end of *Escherichia coli* 16S rRNA, by KsgA is universally conserved throughout evolution (Xu et al., 2008). KsgA, first identified in *E. coli* (Helser et al., 1972), has been found in almost all species. Helix 45, essential for both ribosomal subunit association (Mitchell et al., 1992; Poldermans et al., 1980) and protein synthesis initiation (Poldermans et al., 1979b), is one of the most conserved regions in the small-subunit rRNA (Van Knippenberg et al., 1984). Recently, direct evidence for the interaction of KsgA with small-subunit rRNA near its functional center has been reported by Xu et al. (2008). KsgA methylates the two adjacent adenosines through the transfer of four methyl groups from four molecules of the S-adenosylmethionine (SAM) cofactor to atom N6 of each nucleotide while giving rise to four molecules of S-adenosylhomocysteine (SAH). Mutations in KsgA which result in the loss of dimethylation at A1518 and A1519 are the most common cause of resistance to the amino-

glycoside antibiotic kasugamycin in *E. coli* and other bacteria (Helser et al., 1971, 1972; Van Buul et al., 1983).

Ligand-free KsgA from *E. coli* (Ec-KsgA) is not able to bind cofactor until it binds to RNA (Poldermans et al., 1979a; Thammana and Held, 1974). It was hypothesized that the interaction of KsgA with a part of the 30S subunit separate from helix 45 caused a conformational change, allowing SAM to bind in the binding pocket (O'Farrell et al., 2004). Although KsgA is able to bind naked 16S rRNA (van Gemen et al., 1989), it will not methylate 16S rRNA (Thammana and Held, 1974). A recent study showed that KsgA can only methylate 30S subunits in a translationally inactive form (Desai and Rife, 2006). KsgA shares a high level of sequence homology with the Erm MTases (van Buul and van Knippenberg, 1985), which methylate the large-subunit 23S rRNA in bacterial ribosomes and thus mediate resistance to the macrolide-lincosamide-streptogramin B (MLS-B) group of antibiotics (Skinner et al., 1983). Unlike KsgA, ErmC' from *Bacillus subtilis* (Bs-ErmC') can methylate a fragment of 23S RNA as small as 32 nucleotides (Schluckebier et al., 1999; Weisblum, 1995).

KsgA orthologs from nonbacterial sources are represented by Dim1 from *Saccharomyces cerevisiae* (Sc-Dim1) (Lafontaine et al., 1995, 1998) and from human (Hs-Dim1) (Law et al., 1998; Oh et al., 2005), Pfc1 from *Arabidopsis thaliana* (Tokuhisa et al., 1998), and mitochondria mtTFB from human (Hs-mtTFB) (Cotney and Shadel, 2006; McCulloch et al., 2002; Seidel-Rogol et al., 2003). The mtTFB from yeast (Sc-mtTFB), however, has lost its MTase activity (Klootwijk et al., 1975). Orthologs from archaeobacteria, eubacteria, and eukaryotes have been shown to compensate for the loss of KsgA function in bacteria (Lafontaine et al., 1994; O'Farrell et al., 2006; Seidel-Rogol et al., 2003). Unlike Ec-KsgA, however, Hs-Dim1 appears to be able to bind the cofactor in the absence of RNA (Protein Data Bank [PDB] ID code 1ZQ9).

Structures of KsgA and Erm proteins have been reported for *E. coli* KsgA (Ec-KsgA; PDB ID code 1QYR) (O'Farrell et al., 2004), Hs-Dim1 (PDB ID code 1ZQ9) and Sc-mtTFB (PDB ID code 1I4W) (Schubot et al., 2001), Bs-ErmC' (PDB ID codes 2ERC, 1QAM, 1QAN, 1QAO, and 1QAQ) (Bussiere et al., 1998; Schluckebier et al., 1999), and *Streptococcus pneumoniae* ErmAM (PDB ID code 1YUB) (Yu et al., 1997). However, none of these structures is bound to RNA. Here we present the crystal structures of *Aquifex aeolicus* KsgA (Aa-KsgA) in complex with RNA with and without a bound SAH molecule, along with two high-resolution ligand-free structures, providing insights into the molecular details of ligand recognition by KsgA and the relationship between RNA binding and cofactor binding.

Table 1. Data Collection and Refinement Statistics

Crystal	KsgA1	KsgA2	KsgA-RNA	KsgA-RNA-SAH
X-Ray Diffraction Data				
Space group	P2 ₁ 2 ₁ 2 ₁	P2 ₁ 2 ₁ 2 ₁	C2	P2 ₁
Unit cell parameters				
a (Å)	43.1	42.9	85.2	43.1
b (Å)	57.4	53.6	58.3	58.1
c (Å)	87.6	101.0	91.1	69.0
α (°)	90.0	90.0	90.0	90.0
β (°)	90.0	90.0	107.4	109.8
γ (°)	90.0	90.0	90.0	90.0
Resolution (Å) ^a	30.0–1.44 (1.44–1.48)	30.0–1.68 (1.68–1.74)	30.0–3.0 (3.0–3.11)	30.0–2.78 (2.78–2.88)
Measured reflections	265,573	173,774	38,284	26,715
Unique reflections	38,509	25,134	7,560	6,848
Completeness (%)	96.9 (80.1)	92.3 (54.8)	88.5 (52.9)	84.2 (49.9)
Multiplicity	6.9 (2.7)	6.9 (2.9)	5.1 (2.9)	3.9 (2.1)
R _{merge} (%) ^b	6.8 (44.6)	6.3 (50.0)	10.3 (43.0)	14.4 (39.7)
I/σ(I)	24.6 (1.9)	25.7 (1.7)	13.4 (1.9)	9.0 (2.4)
Crystal Structure				
Effective resolution (Å) ^c	1.44	1.72	3.16	3.05
Number of nonhydrogen atoms				
Protein	2071	1947	1876	1986
Nucleic acid	—	—	950	950
Water	287	180	—	37
R _{factor} (%) ^d	15.9 (29.3)	20.6 (35.3)	23.3 (31.0)	23.9 (34.4)
R _{free} (%) ^d	20.0 (33.6)	24.6 (40.9)	28.0 (44.0)	28.8 (60.3)
Rmsd bonds (Å)	0.005	0.009	0.006	0.005
Rmsd angles (°)	0.9	1.2	1.1	1.0
Overall B factor (Å ²)				
B factor for protein (Å ²)	20.3	30.0	85.5	30.6
B factor for RNA (Å ²)	—	—	79.4	25.1
B factor for SAH (Å ²)	—	—	—	26.3
B factor for solvent (Å ²)	34.4	42.9	—	22.1
Ramachandran statistics (%)				
Most favored φ/ψ	94.4	94.3	86.1	84.0
Additionally allowed φ/ψ	5.6	5.7	13.0	14.2
Disallowed φ/ψ	0.0	0.0	0.0	0.0

^a Values in parentheses are for the highest resolution shell.

^b $R_{\text{merge}} = \sum (|I - \langle I \rangle|) / \sum I$, where I is the observed intensity.

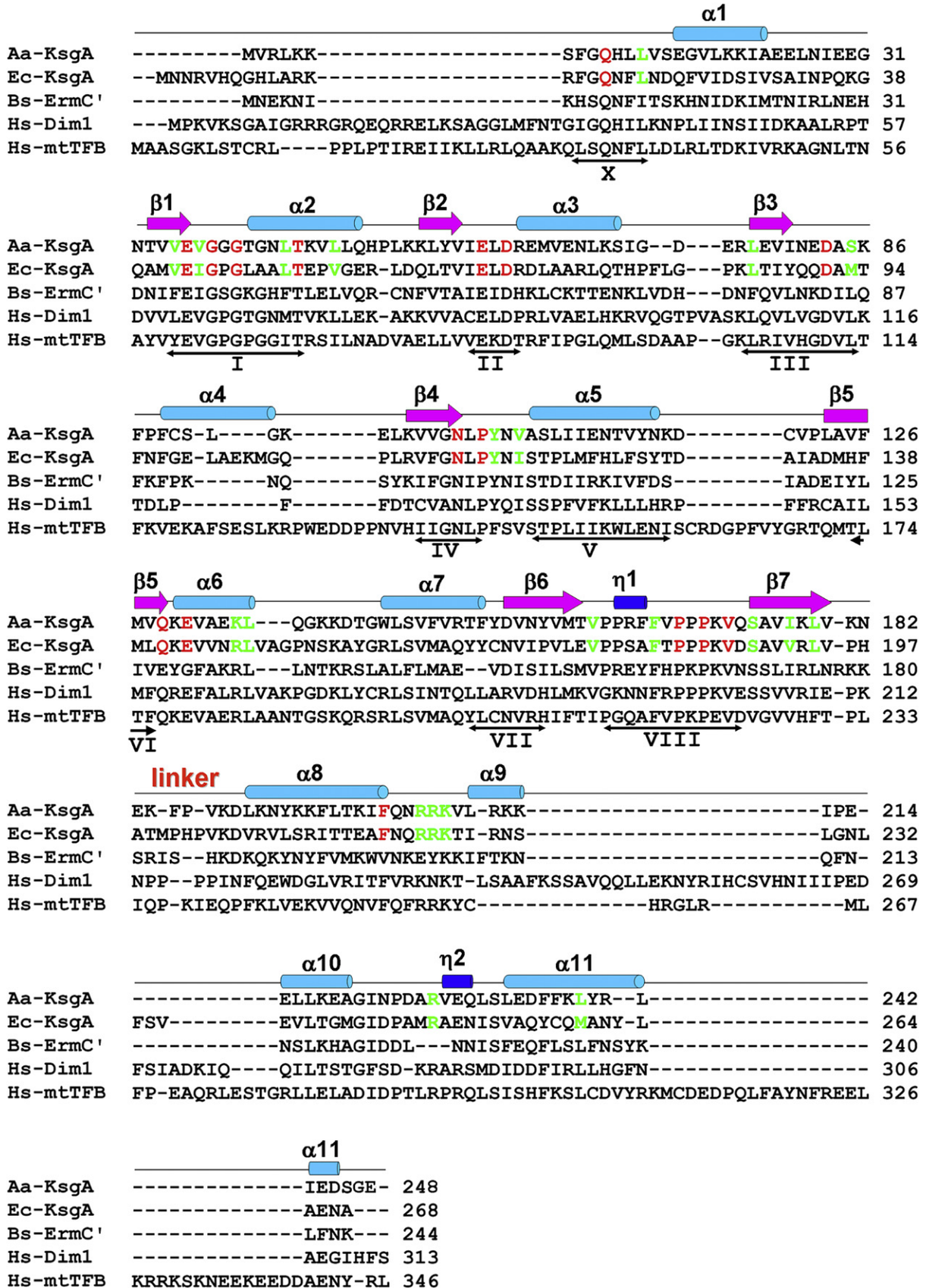
^c At the effective resolution of a structure, the completeness of X-ray data > 93% and the observable data > 70% for the highest resolution shell according to Notes for Authors 2008, *Acta Crystallographica Section D, Biological Crystallography*.

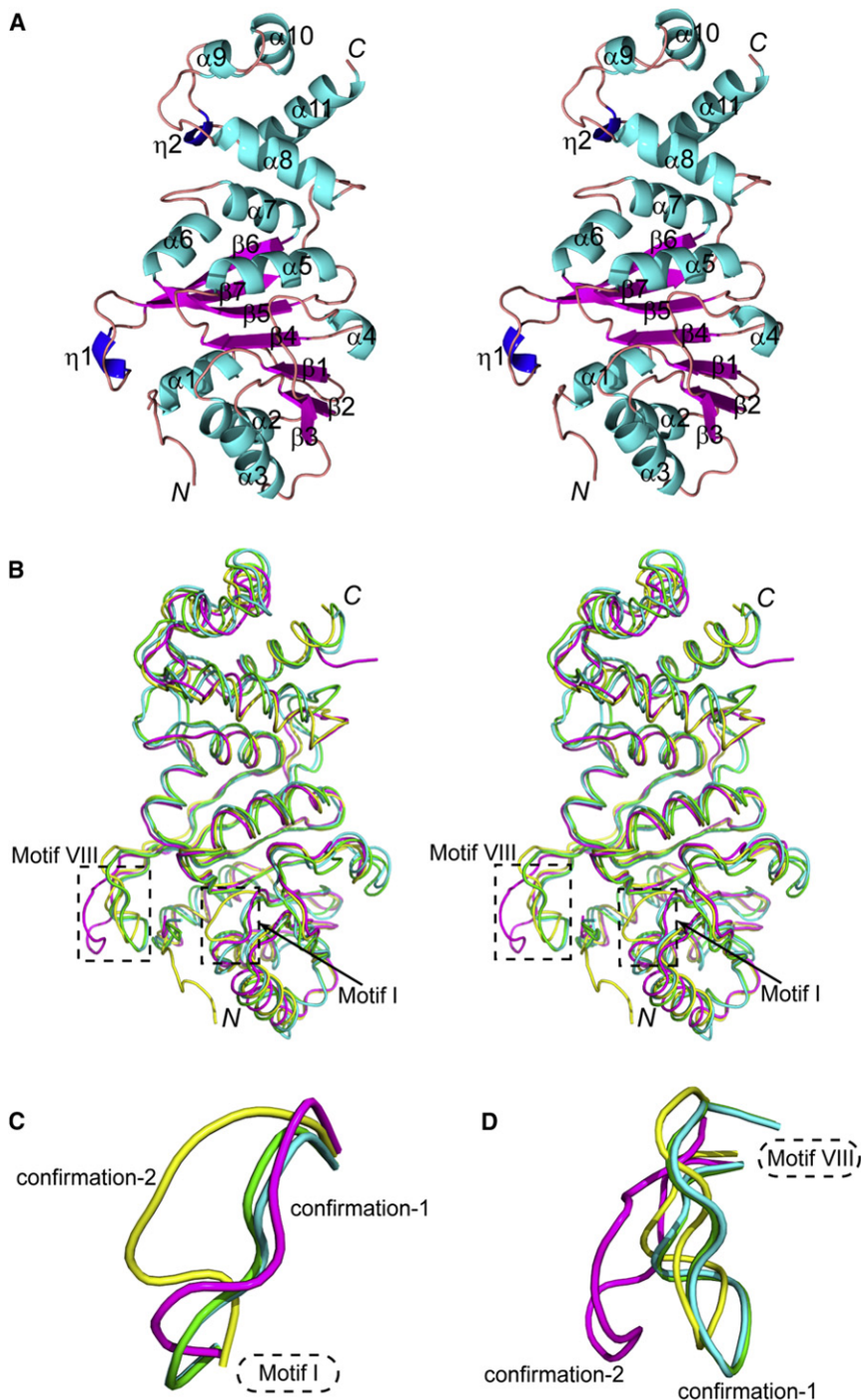
^d R_{factor} and $R_{\text{free}} = \sum ||F_{\text{obs}}| - |F_{\text{calc}}|| / \sum |F_{\text{obs}}|$, where R_{free} was calculated over 5% of the amplitude chosen at random and not used in the refinement. Both values were calculated for the resolution range of data collection.

RESULTS AND DISCUSSION

We report four crystal structures of Aa-KsgA: two in its ligand-free form (KsgA1 and KsgA2), one in complex with RNA (KsgA-RNA), and one in complex with both RNA and SAH (KsgA-RNA-SAH). We found SAH in the ternary complex structure, although SAM was present in the crystallization drops, for which the reason is not clear. Our crystallization trials with SAH, in the presence or absence of RNA, did not yield any crystals. The

Aa-KsgA sequence contains 248 amino acid residues. A nonnative Ser at the N terminus from cloning is not visible in any of the four structures. KsgA1 contains residues 7–246 and 287 water oxygen atoms; KsgA2 contains residues 12–246 and 180 water oxygens; KsgA-RNA contains residues 14–246 and 44 nucleotide residues; and KsgA-RNA-SAH contains residues 1–246, 44 nucleotide residues, an SAH molecule, and 37 water oxygens. X-ray data collection and refinement statistics are summarized in [Table 1](#).



**Figure 2. Ligand-free Aa-KsgA Structures**

(A) Stereo view showing the ribbon diagram of the KsgA1 structure. The α helices, β strands, and 3_{10} helices are numbered and shown in cyan, magenta, and blue, respectively.

(B) C α superposition of KsgA1 (yellow), KsgA2 (magenta), Ec-KsgA-chain A (cyan; PDB ID code 1QYR), and Ec-KsgA-chain B (green, PDB entry 1QYR).

(C) Two distinct conformations of motif I.

(D) Two distinct conformations of motif VIII.

MTases, there are nine common SAM-dependent MTase functional motifs, motifs I–VIII and X (Malone et al., 1995). Motifs I, II, III, IV, and VIII are highly conserved, whereas motifs V, VI, and VII are less conserved. Motifs I, II, and III contain amino acid residues important for cofactor binding, whereas motifs IV and VIII contains residues important for catalysis (Goeddecke et al., 2001; O'Farrell et al., 2004; Schluckebier et al., 1999). The location of motif X in the primary sequence is one of the major differences between C5-cytosine MTases and amino MTases. In the C5-cytosine MTases, this motif is located in the carboxy terminus, whereas in amino MTases, it is always to the amino side of motif I (Malone et al., 1995). Motif X and the upstream N-terminal residues are not visible in any of the previously reported N6-adenine RNA MTase structures, including Ec-KsgA (PDB ID code 1QYR), Hs-Dim1 (PDB ID code 1ZQ9), and ErmC' (PDB ID codes 2ERC, 1QAM, 1QAN, and 1QAO). In our KsgA-RNA-SAH structure, motif X and the entire N terminus of the protein are visible.

Structures of Ligand-free Aa-KsgA

The KsgA1 structure is shown in Figure 2A. The N-terminal domain (NTD) exhibits the SAM-dependent MTase fold. This fold, reminiscent of the Rossmann fold often encountered in mono- or dinucleotide-binding domains, consists of alternating

Figure 1 depicts a structure-based sequence alignment of Aa-KsgA with four related proteins: Ec-KsgA, Bs-ErmC', Hs-Dim1, and Hs-mtTFB. In amino (N4-cytosine or N6-adenine)

α helices and β strands, with a central seven-stranded β sheet sandwiched between a variable number of α helices. Strands 1–6 of the sheet are parallel, with strand 7 inserted antiparallel

Figure 1. Structure-Based Sequence Alignment

The sequences of Aa-KsgA, Ec-KsgA, Bs-ErmC', and Hs-Dim1 (GenBank accession numbers NP_214246, P06992, P13956, and NP_055288, respectively) were aligned based on their structures, whereas the Hs-mtTFB sequence (GenBank accession number NP_057104) was aligned based on sequence homology. Secondary structure elements and the position of the linker between the NTD and CTD are indicated above the sequences. The double-headed arrows under the sequences indicate the nine conserved functional motifs common for SAM-dependent MTases. Absolutely and highly conserved residues in KsgA sequences are indicated in red and green, respectively, most of which are also conserved in Bs-ErmC', although it is not a KsgA protein.

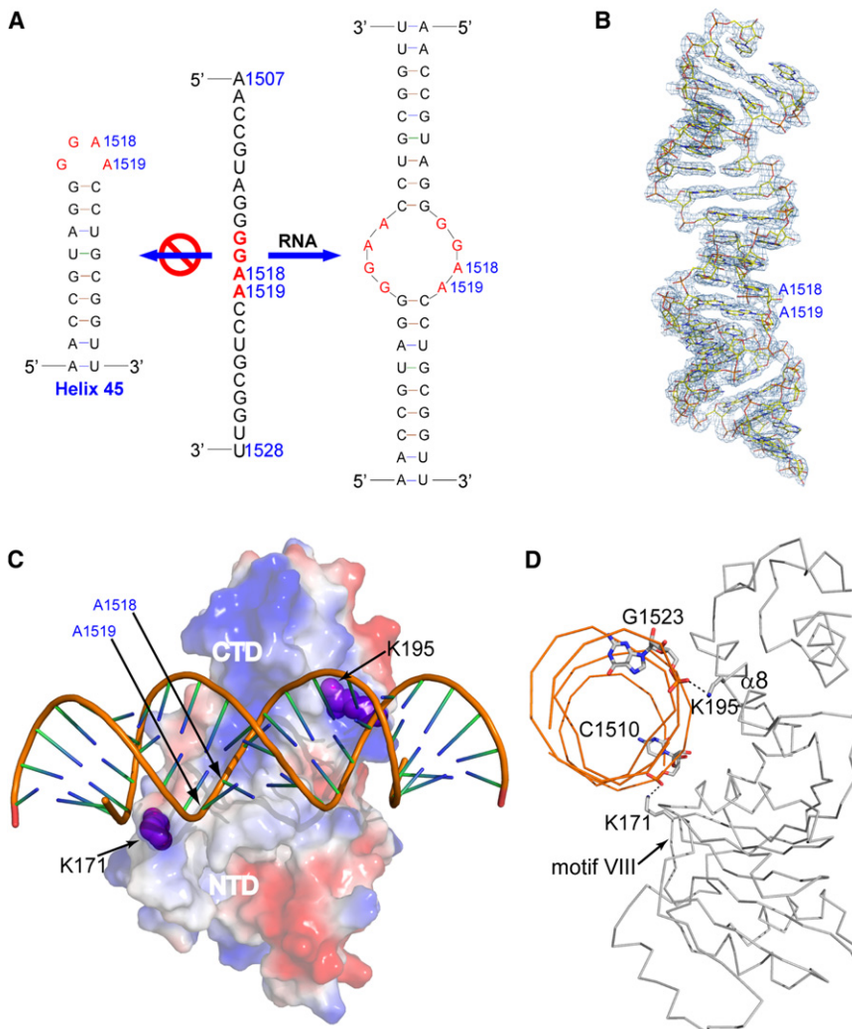


Figure 3. Structure of RNA and of the KsgA-RNA Complex

(A) The sequence of the substrate RNA (residues 1507–1528, *E. coli* numbering) is shown in the middle, while its presumed and observed secondary structures are shown on the left and right, respectively. The presumed tetraloop GGAA is highlighted in red.

(B) Structure of the RNA as observed in the ternary KsgA-RNA-SAH complex. The RNA is shown as a stick model in atomic colors (carbon in yellow, nitrogen in blue, oxygen in red, and phosphorus in orange) and outlined with the final $2F_o - F_c$ electron density map (contoured at 1.0σ in light blue). (C) Surface electrostatic potential for the protein in the binary KsgA-RNA complex. The positively charged area is indicated in blue and the negatively charged is in red. RNA is shown as a tube-and-stick model. Residues K171 (from motif VIII) and K195 (from $\alpha 8$) are highlighted, with sphere representation in magenta.

(D) Ribbon diagram of the binary KsgA-RNA complex viewed along the bound RNA. Shown as stick models are the amino acid or nucleotide residues involved in the two hydrogen bonds (dashed lines in black) between the protein and the RNA.

motif will be referred to as conformation 1, whereas the fourth will be termed conformation 2 (Figures 2C and 2D).

Structure of RNA

Although KsgA does not methylate free 16S rRNA, it binds to a small fragment of rRNA containing the target adenosine bases (van Gemen et al., 1989). The

RNA used in our study contains the 24 contiguous nucleotide residues of helix 45 (Figure 3A, middle) found in the 16S rRNA from *E. coli*. In the structure of the *E. coli* 30S subunit (PDB ID code 2AW7) (Schuwirth et al., 2005), helix 45 forms a hairpin structure capped by a central GGAA tetraloop (Figure 3A, left). In our protein-RNA structures, however, two RNA molecules form a duplex with four mismatched G-A pairs in the middle (Figure 3A, right). None of the mismatched nucleotides flips out of the duplex. The RNA duplex in both KsgA-RNA and KsgA-RNA-SAH are well defined (for an example, see Figure 3B).

Assuming that the substrate RNA has a stem-loop conformation, the hairpin RNA could be aligned with the duplex RNA in two possible ways based on the sequences shown in Figure 3A. Our structures suggest, however, that only one way of hairpin positioning, which brings the tetraloop of the hairpin into contact with K171/motif VIII, is likely to be catalytically compatible. A hypothetical model for the protein-substrate complex will be proposed in the section on implications for catalysis.

For Aa-KsgA, there is one molecule in the asymmetric unit, whereas for Ec-KsgA (PDB ID code 1QYR), there are two polypeptide chains (A and B) in the asymmetric unit. Thus, we have four ligand-free KsgA structures to analyze. Despite the relatively low sequence identity between Aa- and Ec-KsgA, they share very similar three-dimensional structures (Figure 2B). The overall root-mean-square deviation (rmsd) for the C α positions between KsgA1 and Ec-KsgA chain A is 1.8 Å, and the rmsd between KsgA1 and Ec-KsgA chain B is 1.6 Å. The most noticeable differences among the four structures are in the conformation of motifs I and VIII (Figure 2B). For each motif, three conformations are similar and the fourth is distinct. Thus, each motif exhibits two distinct conformations. The three similar conformations of each

RNA used in our study contains the 24 contiguous nucleotide residues of helix 45 (Figure 3A, middle) found in the 16S rRNA from *E. coli*. In the structure of the *E. coli* 30S subunit (PDB ID code 2AW7) (Schuwirth et al., 2005), helix 45 forms a hairpin structure capped by a central GGAA tetraloop (Figure 3A, left). In our protein-RNA structures, however, two RNA molecules form a duplex with four mismatched G-A pairs in the middle (Figure 3A, right). None of the mismatched nucleotides flips out of the duplex. The RNA duplex in both KsgA-RNA and KsgA-RNA-SAH are well defined (for an example, see Figure 3B).

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Structure of the Binary KsgA-RNA Complex

The cleft between the NTD and CTD is positively charged, providing a platform for the enzyme to bind RNA (Figure 3C).

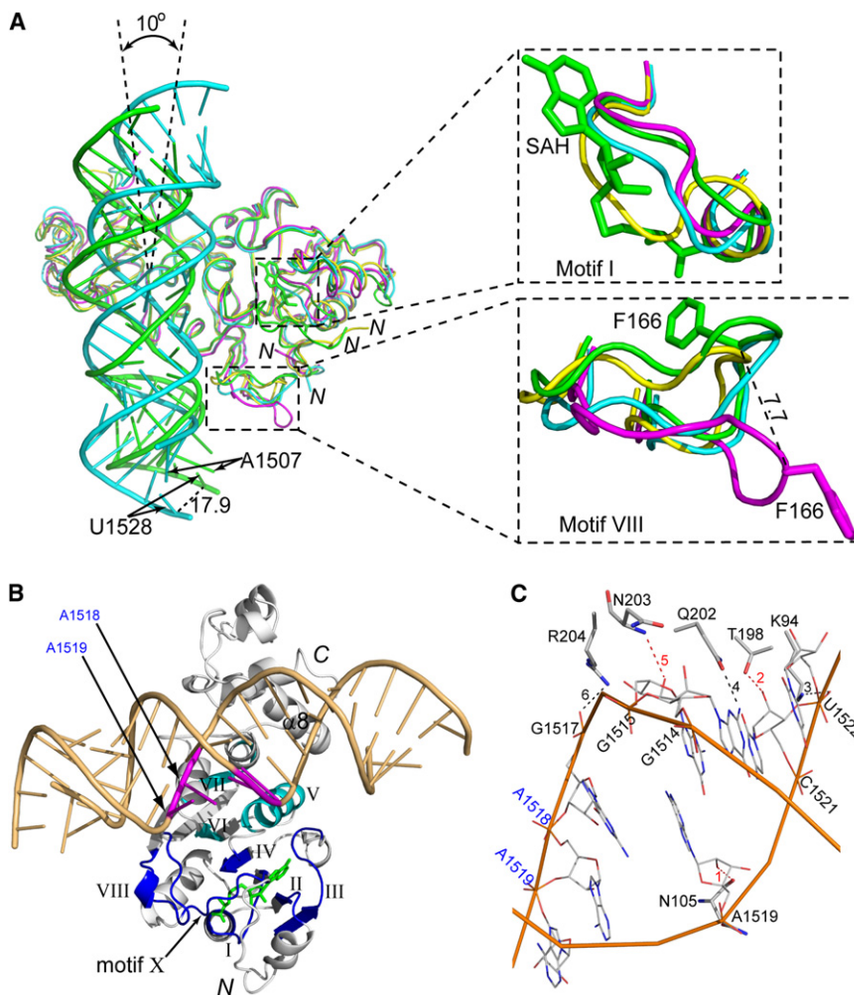


Figure 4. Structure of the KsgA-RNA-SAH Complex

(A) $C\alpha$ superposition of KsgA1 (yellow), KsgA2 (magenta), KsgA-RNA (cyan), and KsgA-RNA-SAH (green). Protein is shown as a ribbon diagram, RNA as tube-and-stick models, and cofactor SAH as a stick model. The rotation of RNA ($\sim 10^\circ$) and the distance of the U1528 shift (17.9 Å) relative to the RNAs in the two complexes are indicated. Close-up views of motifs I and VIII are shown in the upper and lower right panels. The distance (7.7 Å) of the $C\alpha$ shift of the highly conserved residue F166 between KsgA2 and KsgA-RNA-SAH is also indicated. The side chain of F166 is shown as a stick model.

(B) Front view of the KsgA-RNA-SAH structure: protein as a ribbon diagram in white; substrate RNA as a tube-and-stick model in light orange with the adjacent A1518 and A1519 highlighted in magenta; and cofactor SAH as a stick model in green. Roman numerals indicate the conserved structural motifs of the protein: well-conserved motifs I–IV, VIII, and X in blue; less-conserved motifs V–VII in cyan.

(C) Six strong hydrogen bonds (dashed lines) are observed between the protein and the RNA in the KsgA-RNA-SAH structure. Amino acid residues interacting with the bound RNA are shown as thicker sticks while the nucleotide residues are shown as thinner sticks. Hydrogen bonds 1, 2, and 5 (colored in red) involve the O2' hydroxyl groups of nucleotide residues.

Only residues on the NTD side of the cleft have been previously shown to be important in the binding of Bs-ErmC' to its substrate RNA (Maravic et al., 2003a). We show here that positively charged side chains on both sides of the cleft are involved in double-stranded (ds) RNA binding by Aa-KsgA. In the KsgA-RNA structure, K171 (in motif VIII) and K195 (in helix $\alpha 8$) each forms a strong hydrogen bond with a phosphate O2P of RNA (Figure 3D). As a result of RNA binding, the enzyme undergoes conformational changes in the relative positioning of the NTD and CTD as well as within the two domains, among which the most significant is the stabilization of motifs I and VIII. In ligand-free KsgA, each motif exhibits two distinct conformations (Figures 2C and 2D). In the protein-RNA complexes, however, both motifs are stabilized in conformation 1 (Figure 4A).

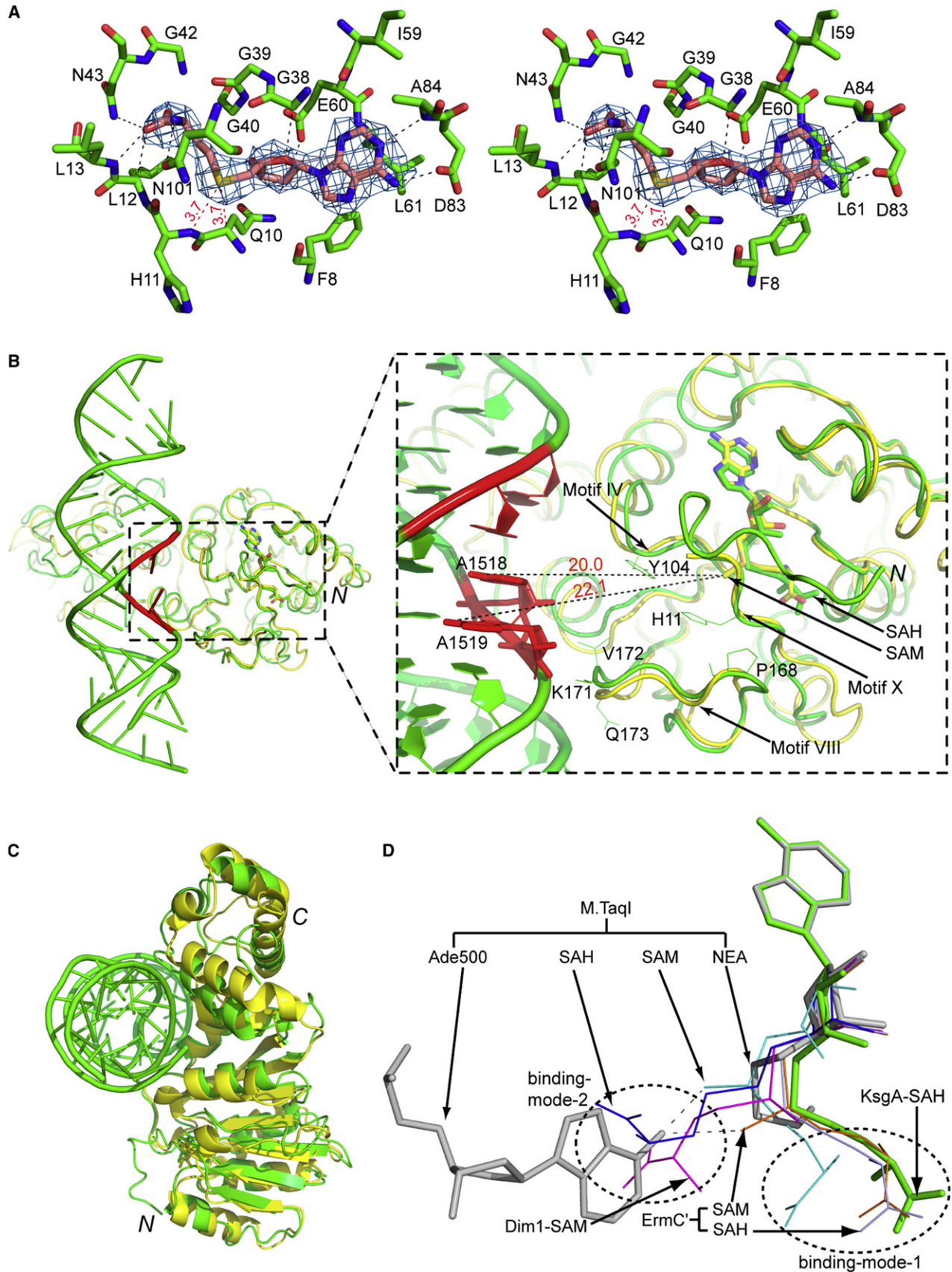
Structure of the Ternary KsgA-RNA-SAH Complex

Ec-KsgA does not bind cofactor until it binds to RNA (O'Farrell et al., 2004; Poldermans et al., 1979a; Thammana and Held, 1974). We have observed that Aa-KsgA does not crystallize in the presence of SAM alone, but crystallizes in the presence of either RNA alone or both RNA and SAH, suggesting that once KsgA binds the RNA in the cleft between the NTD and CTD, it can then bind the cofactor. RNA binding in the cleft stabilizes

motifs I and VIII in conformation 1 (Figure 4A). In conformation 1, motif VIII is pointing toward the cofactor-binding site, whereas in conformation 2, it is pointing away from the cofactor-binding site. In conformation 1, motif I is not in the way of the bound SAH, whereas in conformation 2, it blocks the cofactor-binding site. Thus, for both motifs, conformation 1 is compatible with cofactor binding and conformation 2 is not.

Previously, motif X was thought to be nonconserved and to comprise residues in the N-terminal α helix (O'Farrell et al., 2004). We show here that motif X contains conserved residues Q10 and L13, but does not include $\alpha 1$ residues (Figure 1). It appears that the N-terminal loop and motif X of KsgA become ordered only after binding both the RNA and the cofactor, although the functional role of the N-terminal loop is not clear. Among the better-conserved motifs, motifs I–IV and X are located in proximity to the cofactor, whereas motif VIII interacts with both the cofactor and the RNA. Less-conserved motifs V–VII do not contribute directly to the binding of RNA or cofactor (Figure 4B).

The structures of the bound RNA in the binary and ternary complexes are similar. However, the binding of the SAH occurs in concert with a change of the relative positioning between the protein and the RNA, which is equivalent to an $\sim 10^\circ$ counterclockwise rotation of the RNA duplex (Figure 4A). When the cofactor-binding site is occupied, the interaction between the protein and the RNA is significantly enhanced. Compared to



the binary complex, a total of 12 residues make contacts with the RNA, including N105, K140, K171, V172, Q173, K194, K195, T198, K199, Q202, N203, and R204. Among the contacts between the protein and the RNA, six strong hydrogen bonds are observed, three interacting with the O2' hydroxyls of G1515, A1519, and C1521, two with phosphate oxygen O2P of G1517 and U1522, and one with atom N2 of G1514 (Figure 4C). Apparently, the three strong hydrogen bonds from N105, T198, and N203 to the O2' hydroxyls of RNA distinguish the dsRNA from dsDNA.

Cofactor Binding

Five out of the nine common motifs are involved in the binding of SAH. Motif X (₈FGQHLL₁₃) makes extensive contacts with the bound SAH (Figures 4B and 5A). The distance between the sulfur of SAH and residue Q10 or H11 is 3.7 Å, suggesting that these two residues may assist in orienting the methyl group of SAM for catalysis. Motif I (₃₆VEVGGGTGNLT₄₅), that is, the G loop, forms part of the cofactor-binding pocket (Figure 5A). Motif II (₅₉LLED₆₂) contains two absolutely conserved residues, E60 and D62, among which E60 forms a hydrogen bond with the O2' hydroxyl on the ribose moiety of SAH. Motif III (₇₈LEVINEDAS₈₅) contains one absolutely conserved residue D83, which forms a hydrogen bond with the exocyclic N6 amine group of the SAH adenine moiety. Motif IV (₉₈VVGNNLP₁₀₃) contains residues that are important for both cofactor binding and catalysis. One of the absolutely conserved residues, N101, is hydrogen bonded with atom N of homocysteine (Figure 5A).

Motifs V (residues 107–117), VI (residues 126–128), and VII (residues 151–156) are less conserved (Figure 1). These motifs do not directly interact with SAH (Figure 4B). The highly conserved motif VIII (₁₆₂PPRFFVPPPQV₁₇₃) may play multiple roles in KsgA catalysis. This motif contains the highly conserved residue F166 (Figure 4A). This residue in DNA N6-adenine MTase M·TaqI is F196, the side chain of which is in an edge-to-face π -stacking arrangement with the target adenine (PDB ID code 1G38). There is an extensive network of electrostatic and hydrophobic interactions between residues F166, V167, and P168 in motif VIII and residues H11 and L12 in motif X (not shown). Residue H11 also interacts with motif IV through highly conserved Y104 (Figure 5B). Thus, motifs IV, VIII, and X must be functionally dependent on one another.

Cofactor SAM and product SAH assume a common binding mode in Bs-ErmC' (Schluckebier et al., 1999). This binding mode is also observed in KsgA-RNA-SAH. Based on the C α positions in the NTD, three structures are aligned and depicted in Figure 5B for the superposition of KsgA-RNA-SAH and ErmC'-SAM (PDB ID code 1QAO), and in Figure 5C for the superposition of KsgA-RNA-SAH and ErmC'-SAH (PDB ID code 1QAN). The positioning of the adenine and ribose moieties of SAH in KsgA is virtually the same as that of SAM in EcmC', and the two molecules show only small conformational differences in their methionine moieties (Figure 5B). This cofactor-binding mode (designated binding mode 1) has also been observed for the SAM molecule in the M·TaqI-SAM structure (PDB ID code 2ADM).

For N6-adenine MTases, a second cofactor-binding mode has been observed in which the methionine moiety is pointing in the opposite direction (designated binding mode 2) as shown in Figure 5D. Among members of the N6-adenine MTase family, molecular details for the methyl group transfer have been revealed for M·TaqI on the basis of the structure of the enzyme in complex with both DNA and a cofactor analog NEA (PDB ID code 1G38). The arrangement of the substrate adenine base and the NEA in the M·TaqI-DNA-NEA structure suggests that in binding mode 1, the activated methyl group of SAM points toward the N6 of the substrate adenine, whereas in binding mode 2, the methyl group points away from the substrate and consequently the methionine moiety appears to block the proper positioning of the adenine base (Figure 5D). Therefore, cofactor binding mode 1 is compatible with the direct transfer of the methyl group from SAM to atom N6 of adenosine, whereas binding mode 2 may initiate product release.

Implications for Catalysis

In KsgA-RNA-SAH, the two RNA molecules form a duplex with four G-A mismatches in the middle (Figure 3A). Although this arrangement is not likely the catalytic assembly, it shows how KsgA binds a dsRNA. In addition, it provides guidance for a comparative analysis of available structural information, suggesting how KsgA may bind a substrate RNA. It has previously been suggested that methylation of A1518 and A1519 alters the RNA conformation significantly (Rife and Moore, 1998). If the substrate RNA assumes the stem-loop conformation,

Figure 5. The Cofactor-Binding Site in Aa-KsgA and Bs-ErmC'

(A) Stereo view of the SAH-binding site in the ternary KsgA-RNA-SAH complex. The SAH cofactor and the surrounding amino acid residues within hydrogen-bonding distance (3.5 Å) are shown as stick models in atomic colors (carbon in magenta in SAH and in green in the surrounding amino acids, nitrogen in blue, oxygen in red, and sulfur in yellow). The cofactor is outlined with the final $2F_o - F_c$ electron density map contoured at 1.0σ and in blue. Hydrogen bonds between SAH and nearby amino acid residues are indicated with dashed lines in black, and the distance between the SAH sulfur and the C α of Q10 and the nitrogen of H11 are indicated in red. The locations of motifs I–IV and X are indicated.

(B) C α -trace superposition between KsgA in the ternary KsgA-RNA-SAH complex (green) and Bs-ErmC' in the binary ErmC'-SAM complex (PDB ID code 1QAO). Proteins are illustrated as ribbon diagrams and RNA as a tube-and-stick model. The adjacent adenosines are highlighted in red. Residues A1518 and A1519, which are closer to the putative catalytic center, are shown as stick models. The cofactors are shown as stick models in atomic colors (carbon in green in SAH and in yellow in SAM, nitrogen in blue, oxygen in red, and sulfur in yellow). Conserved residues in KsgA which appear to play important roles in either the binding of cofactor/RNA or the stabilization of target adenosines are shown as thinner sticks in green for KsgA. The distances between atom S of SAM and the N6 of highlighted adenosines are indicated with dashed lines in black.

(C) C α -trace superposition between the ErmC'-SAH complex (yellow; PDB ID code 1QAN) and the KsgA-RNA-SAH complex (green).

(D) Superposition of SAH in KsgA-RNA-SAH (green), SAH in ErmC'-SAH (light blue; PDB ID code 1QAN), SAH in M·TaqI-SAH (blue; PDB ID code 1AQI) (Schluckebier et al., 1997), SAM in ErmC'-SAM (light orange; PDB ID code 1QAO), SAM in Dim1-SAM (magenta; PDB ID code 1ZQ9), and SAM in M·TaqI-SAM (cyan; PDB ID code 2ADM) (Labahn et al., 1994; Schluckebier et al., 1997). The substrate adenosine and the cofactor analog NEA in M·TaqI-DNA-NEA (gray; PDB ID code 1G38) are also aligned. The alignment is based on the superposition of the adenine base of the cofactors. Stick models are shown for KsgA-RNA-SAH and M·TaqI-DNA-NEA, and line models are used for other structures.

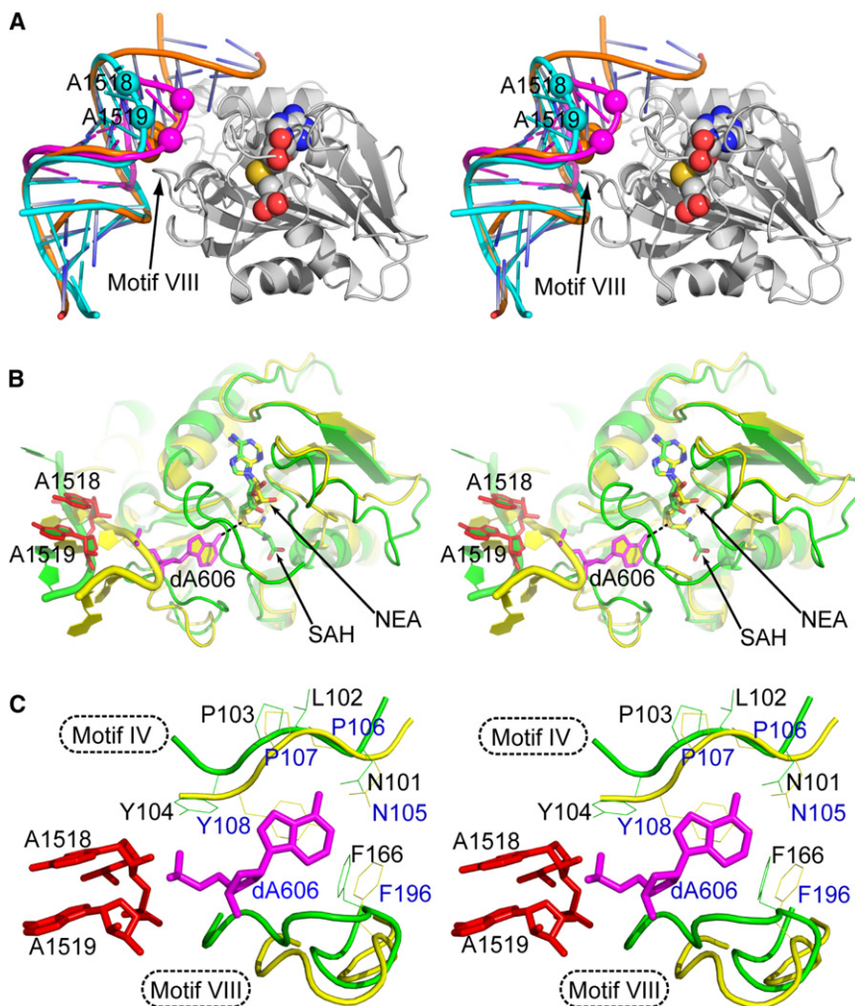


Figure 6. The Putative Catalytic Center of KsgA

(A) Stereo view showing the superposition of tetra-loop structures with the KsgA-RNA-SAH complex. SAH is shown as spheres (carbon in gray, nitrogen in blue, oxygen in red, and sulfur in yellow). KsgA is shown as a ribbon diagram in gray. RNAs are illustrated as tube-and-stick models with the positions of A1518 and A1519 indicated by spheres. RNA in KsgA-RNA-SAH is shown in orange, a structure of product RNA with dimethylated A1518 and A1519 (PDB ID code 2AW7) in cyan, and a model for substrate RNA with unmethylated adenosines (PDB ID code 1AFX) in magenta.

(B) Stereo view showing the superposition of the putative catalytic center of KsgA as observed in the ternary KsgA-RNA-SAH complex (green) with the catalytic center assembly of M·TaqI as observed in the M·TaqI-DNA-NEA structure (PDB ID code 1G38; yellow). The alignment was based on the cofactor SAH (stick model with carbon in green, nitrogen in blue, oxygen in red, and sulfur in yellow) and the cofactor analog NEA (stick model with carbon in yellow). The target adenine dA606 for M·TaqI is shown as a stick model in magenta, whereas the A1518 and A1519 for KsgA are shown as stick models in red. The contact between atom N6 of dA606 in the M·TaqI complex and atom S of SAH in the KsgA complex is indicated with a dashed line in black.

(C) A close-up stereo view showing the details of the adenine-binding site. The amino acid side chains in motif IV (N105, P106, P107, and Y108) and motif VIII (F196) in M·TaqI are shown as line models in yellow. The corresponding side chains in KsgA (N101, L102, P103, Y104, and F166) are shown in green. The adenine substrate is shown in magenta.

a structure of dimethylated helix 45 (PDB ID code 2AW7) and a model of unmethylated stem loop (PDB ID code 1AFX; Butcher et al., 1997) can be superpositioned with the RNA duplex in the KsgA-RNA-SAH structure (Figure 6A). Because the RNA duplex in the structure is symmetric, the two stem-loop structures can also be aligned with the other half of the duplex. However, only the alignment shown in Figure 6A positions the target adenosines at the entrance of the catalytic center and in proximity to motif VIII. Although the stem-loop RNA may not form strong hydrogen bonds with the CTD as suggested by the dsRNA in the KsgA-RNA-SAH structure (Figure 4C), the crucial role of motif VIII in catalysis is clearly implicated (Figure 6A). Ec-KsgA can only methylate 30S subunits in a translationally inactive form (Desai and Rife, 2006). Likely, this is also true for Aa-KsgA. Thus, the enzyme may be embedded in the 30S and interact with RNA extensively, which should be sufficient to position the A1518-A1519 half of the tetraloop in contact with motif VIII.

In the KsgA-RNA-SAH structure, the distance between the acceptor (atom N6 of adenine) and the donor (atom S of SAH) of the methyl group transfer reaction is ~ 20 Å (Figure 5B). In the KsgA-substrate model, the acceptor-donor distance is also ~ 20 Å (Figure 6A). For a direct methyl group transfer mechanism, this distance needs to be ~ 5 Å (Goedecke et al., 2001; Klima-

sauskas et al., 1994). The DNA double helix in the M·TaqI-DNA-NEA structure (PDB ID code 1G38) is located at a distance similar to that for the RNA duplex in KsgA-RNA-SAH, suggesting that a base-flipping mechanism may bring the acceptor in proximity to the donor of the reaction (Figure 6B). Thus, the catalytic center assembly of KsgA may be modeled on the basis of KsgA-RNA-SAH and M·TaqI-DNA-NEA (PDB ID code 1G38).

Although the overall structures of the M·TaqI and KsgA complexes are different, the two N6-adenine MTases share a very similar cofactor-binding site (Figure 6B). Five residues of M·TaqI in motif IV (N105, P106, P107, and Y108) and motif VIII (F196) are involved in the binding and stabilization of the target adenosine (dA606). The counterparts of these five residues in KsgA are N101, L102, P103, Y104, and F166 (Figure 6C). The importance of these residues in catalysis has been demonstrated by mutational studies. Mutations of Asn and Phe residues are poorly tolerated in DNA MTases and ErmC' (Guyot et al., 1993; Kong and Smith, 1997; Maravic et al., 2003b; Pues et al., 1999; Roth et al., 1998; Willcock et al., 1994), whereas mutation of the Tyr residue totally abolishes the activity of both M·TaqI and ErmC', both in vitro and in vivo (Pues et al., 1999). Four out of these five residues are either absolutely (N101 and P103) or highly (Y104 and F166) conserved in KsgA (Figure 1).

Residue P106 in M·TaqI does not interact with the target adenine; its counterpart L102 in KsgA is not highly conserved either.

In addition to the conserved residues in M·TaqI which are involved in the stabilization of the target adenine, residues V21 from motif X and K199 from motif VIII are in contact with the substrate adenine. The counterparts of V21 and K199 in KsgA are residues H11 from motif X and P168 from motif VIII. Residue H11 may play multiple roles because (1) the amide of H11 is very close to the sulfur of SAH (Figure 5A) and (2) the side chain of H11 is hydrogen bonded with motif VIII residue P168 and in close contact with motif IV residue Y104 (Figure 5B). Thus, H11 seems to function as a hub in the active center of KsgA by interacting with both the cofactor and the target adenine and linking two catalytically very important motifs (IV and VIII). Residue P168 of KsgA may mimic the functional role of K199 in M·TaqI, which is part of the DNA-binding loop and undergoes a conformational change upon DNA binding. Similarly, P168 of KsgA is in the middle of flexible motif VIII that also undergoes significant conformational changes upon RNA binding (Figure 4A).

Conclusions

Our ligand-free structures reveal two distinct conformations for both motif I and motif VIII (Figure 2). Our complex structures provide insights into the molecular details of ligand recognition by Aa-KsgA and the relationship between RNA binding and cofactor binding. We show how the RNA is bound, how the RNA-bound KsgA is changed for SAH binding, and how SAH is bound. The protein binds to the RNA duplex through motif VIII in the NTD and helix $\alpha 8$ in the CTD (Figures 3C and 4B). It binds SAH mainly through motifs I, II, III, IV, and X (Figures 4B and 5A). Before RNA binding, motifs I and VIII are flexible (Figure 2B). Upon RNA binding, motif VIII is stabilized which in turn stabilizes motif I, allowing the cofactor-binding site to form. Motif X, along with residues 1–6 at the N terminus, exhibits ordered structure in KsgA-RNA-SAH. A comparison of protein-RNA interactions between KsgA-RNA and KsgA-RNA-SAH indicates that 10 more residues of the protein in the ternary complex are involved in RNA binding (Figures 3D and 4C). For catalysis, the substrate adenine must become an integral part of the catalytic center assembly, for which we have proposed a base-flipping mechanism that has yet to be elucidated. Whether the dimethylation of the two adenosines is carried out by a single- or multi-binding event remains to be seen.

EXPERIMENTAL PROCEDURES

Cloning, Protein Expression, and Purification

The ORF of Aa-KsgA was amplified from genomic DNA by PCR using the following oligonucleotide primers: 5'-GAG AAC CTG TAC TTC CAG TCT ATG GTA AGA CTG AAA AAA TCC TTC-3' and 5'-GGG GAC CAC TTT GTA CAA GAA AGC TGG GTT ATT ACT CTC CAG AAT CCT CAA TTA ATC-3' (primer R). The PCR amplicon was subsequently used as a template for a second PCR with the following primers: 5'-GGG GAC AAG TTT GTA CAA AAA AGC AGG CTC GGA GAA CCT GTA CTT CCA G-3' and primer R. The amplicon from the second PCR was inserted by recombinational cloning into the entry vector pDONR221 (Invitrogen, Carlsbad, CA, USA) and the nucleotide sequence was confirmed experimentally. The open reading frame (ORF) encoding the KsgA gene now with a recognition site for tobacco etch virus (TEV) protease (ENLYFQ/S) fused in-frame to its N terminus was moved by recombinational cloning into the destination vector pDEST-HisMBP (Nal-

lamsetty and Waugh, 2006) to produce pBA1939. pBA1939 directs the expression of Aa-KsgA as a fusion to the C terminus of *E. coli* maltose-binding protein (MBP) with an intervening TEV protease recognition site. The MBP contains an N-terminal hexahistidine tag for affinity purification by immobilized metal affinity chromatography. The fusion protein was expressed in *E. coli* strain BL21-CodonPlus(DE3)-RIL (Stratagene, La Jolla, CA, USA). Cells were grown to mid-log phase ($OD_{600} \sim 0.5$) at 37°C in Luria broth containing 100 $\mu\text{g/ml}$ ampicillin, 30 $\mu\text{g/ml}$ chloramphenicol, and 0.2% glucose. Overproduction of fusion protein was induced with isopropyl- β -D-thiogalactopyranoside at a final concentration of 1 mM for 4 hr at 30°C. The cells were pelleted by centrifugation and stored at -80°C .

All procedures were performed at 4°C–8°C. Ten grams of *E. coli* cell paste was suspended in 150 ml of ice-cold 50 mM sodium phosphate (pH 7.5), 200 mM NaCl, 25 mM imidazole buffer (buffer A) containing 1 mM benzamidine HCl (Sigma, St. Louis, MO, USA) and complete EDTA-free protease inhibitor cocktail tablets (Roche Molecular Biochemicals, Indianapolis, IN, USA). The cells were lysed with an APV-1000 homogenizer (Invensys, Røhølsvej, Denmark) at 10,000 psi, and centrifuged at 30,000 $\times g$ for 30 min. The supernatant was filtered through a 0.22 μm polyethersulfone membrane and applied to a HisPrep FF 16/10 column (GE Healthcare Bio-Sciences, Piscataway, NJ, USA) equilibrated in buffer A. The column was washed to baseline with buffer A and eluted with a linear gradient of imidazole to 500 mM. Fractions containing recombinant His₆-MBP-KsgA were pooled, concentrated using an Amicon YM30 membrane (Millipore, Bedford, MA, USA), diluted with 50 mM sodium phosphate (pH 7.5), 200 mM NaCl buffer (buffer B) to reduce the imidazole concentration to about 25 mM, and digested overnight at 4°C with His₆-tagged TEV protease. The digest was applied to the HisPrep FF 16/10 column equilibrated in buffer A and the KsgA emerged in the column effluent. The column effluent was incubated with dithiothreitol (10 mM), concentrated using an Amicon YM10 membrane, and applied to a HiPrep 26/60 Sephacryl S-100 HR column (GE Healthcare Bio-Sciences) equilibrated in buffer B. The peak fractions containing KsgA were pooled, heat treated at 90°C for 90 min, and clarified by centrifugation and filtration. The sample was concentrated as above and applied to a HiPrep 26/60 Sephacryl S-100 HR column equilibrated in 25 mM Tris (pH 7.2), 150 mM NaCl, 2 mM Tris(2-carboxyethyl) phosphine (TCEP) buffer. The peak fractions containing recombinant KsgA were pooled and concentrated to 30–40 mg/ml (estimated at 280 nm using a molar extinction coefficient of 15,930 $\text{M}^{-1}\cdot\text{cm}^{-1}$). Aliquots were flash-frozen in liquid nitrogen and stored at -80°C . The final product was judged to be >90% pure by sodium dodecyl sulfate-polyacrylamide gel electrophoresis. The molecular weight was confirmed by electrospray ionization mass spectroscopy.

Crystallization and Data Collection

Protein concentrations were adjusted to ~ 27 mg/ml in 25 mM Tris (pH 7.2), 150 mM NaCl, 2 mM TCEP buffer for crystallization trials. RNA oligonucleotides were purchased from Integrated DNA Technologies (Coralville, IA, USA) and used without further purification. The molar ratio of RNA to protein was kept at 1.2:1 for crystallization. Initially, the ratio was for single-stranded RNA versus protein, and later changed to dsRNA versus protein, but the concentration of RNA did not make any difference in the composition of the protein-RNA complexes. RNAs were heated at 75°C for 5 min and cooled on ice for 10 min before being mixed with the protein solution. The protein-RNA mixtures were heated at 70°C for 5 min and slowly cooled down to and kept at room temperature for 45 min before crystallization. For the ternary complex, ~ 10 mM SAM was added and the mixture was kept at room temperature for another 30 min, but SAH was observed in the ternary complex. Crystallization trials of the protein with either SAH alone or both SAH and RNA did not yield any crystals. A different RNA containing 28 nucleotide residues was present in the crystallization drops of KsgA1 and KsgA2, but only the protein crystallized.

Crystals were grown at $19^\circ\text{C} \pm 1^\circ\text{C}$. A Hydra II Plus One (Matrix Technologies, Hudson, NH, USA) crystallization robot system was employed. The sitting drops of KsgA1 and KsgA2 contained 0.2 μl of protein solution and 0.2 μl of reservoir solution. For KsgA1, the reservoir solution contained 20% PEG MME 5000 and 0.1 M bis-Tris (pH 6.5). For KsgA2, the reservoir contained 25% PEG 3000 and 0.1 M MES (pH 6.5). Crystals for both apo forms did not appear until 12 months later and reached a final size of 0.20 mm \times 0.15 mm \times 0.05 mm. The drops of KsgA-RNA crystals contained 0.3 μl protein-RNA

solution and 0.3 μ l reservoir solution containing 0.2 M NH_4NO_3 and 40% *m*-phenylenediamine (MPD), and the crystal reached a size of 0.15 mm \times 0.05 mm \times 0.01 mm in 10 days. The drops of KsgA-RNA-SAH contained 0.4 μ l protein-RNA-cofactor solution and 0.2 μ l reservoir solution containing 0.2 M KF and 40% MPD, and the crystals reached a final size of 0.4 mm \times 0.05 mm \times 0.01 mm in 7–10 days.

The crystals were flash-frozen in liquid nitrogen before data collection. The X-ray data of KsgA1, KsgA2, and KsgA-RNA-SAH were collected at beamline 22-BM, whereas the data of KsgA-RNA were collected at 22-ID, of the Southeast Regional Collaborative Access Team (SER-CAT) at the Advanced Photon Source, Argonne National Laboratory. Data processing was carried out with the HKL2000 program suite (Otwinowski and Minor, 1997). Crystal data and processing statistics are summarized in Table 1.

Structure Solution and Refinement

The structures were solved by molecular replacement (MR) using the program Phaser (McCoy et al., 2007). The search model for KsgA2 was chain A of the Ec-KsgA structure (PDB ID code 1QYR), whereas the search model for KsgA1 was the refined KsgA2 structure. The two ligand-free structures were refined using PHENIX (Adams et al., 2002) and REFMAC (Murshudov et al., 1997). To solve the two complex structures, the refined KsgA1 structure and the helix 45 structure (residues 1507–1528) of the 16S rRNA structure (PDB ID code 1J5E) (Wimberly et al., 2000) were used as search models. The MR solution contained one protein and two RNA molecules. The electron density revealed that the two RNA molecules formed a continued double helix. The two complex structures were refined using CNS (Brünger et al., 1998) and REFMAC (Murshudov et al., 1997).

Bulk solvent correction was employed in all of the refinements. The $2F_o - F_c$ and $F_o - F_c$ electron density maps were used to inspect and improve the models during the refinement. The SAH molecule was built using the REFMAC library (Vagin et al., 2004). Solvent molecules, as peaks $\geq 3\sigma$ on the $F_o - F_c$ electron density map with reasonable hydrogen-bond networks, were included as water molecules at the later stage of the refinement and verified with omit maps. All graphics work was carrying out using O (Jones et al., 1991) and Coot (Emsley and Cowtan, 2004). The refined structures were assessed using PROCHECK (Laskowski et al., 1993). Illustrations were prepared with PyMOL (DeLano Scientific LLC).

ACCESSION NUMBERS

The coordinates and structure factors have been deposited in the Protein Data Bank under ID code 3FTD for KsgA1, 3FTC for KsgA2, 3FTE for KsgA-RNA, and 3FTF for KsgA-RNA-SAH.

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REFERENCES

Adams, P.D., Grosse-Kunstleve, R.W., Hung, L.W., Ioerger, T.R., McCoy, A.J., Moriarty, N.W., Read, R.J., Sacchettini, J.C., Sauter, N.K., and Terwilliger, T.C. (2002). PHENIX: building new software for automated crystallographic structure determination. *Acta Crystallogr. D Biol. Crystallogr.* 58, 1948–1954.

Brünger, A.T., Adams, P.D., Clore, G.M., DeLano, W.L., Gros, P., Grosse-Kunstleve, R.W., Jiang, J.S., Kuszewski, J., Nilges, M., Pannu, N.S., et al. (1998). Crystallography & NMR system: a new software suite for macromolecular structure determination. *Acta Crystallogr. D Biol. Crystallogr.* 54, 905–921.

Bussiere, D.E., Muchmore, S.W., Dealwis, C.G., Schluckebier, G., Nienaber, V.L., Edalji, R.P., Walter, K.A., Lador, U.S., Holzman, T.F., and Abad-Zapatero, C. (1998). Crystal structure of ErmC', an rRNA methyltransferase which mediates antibiotic resistance in bacteria. *Biochemistry* 37, 7103–7112.

Butcher, S.E., Dieckmann, T., and Feigon, J. (1997). Solution structure of the conserved 16 S-like ribosomal RNA UGAA tetraloop. *J. Mol. Biol.* 268, 348–358.

Cotney, J., and Shadel, G.S. (2006). Evidence for an early gene duplication event in the evolution of the mitochondrial transcription factor B family and maintenance of rRNA methyltransferase activity in human mtTFB1 and mtTFB2. *J. Mol. Evol.* 63, 707–717.

Desai, P.M., and Rife, J.P. (2006). The adenosine dimethyltransferase KsgA recognizes a specific conformational state of the 30S ribosomal subunit. *Arch. Biochem. Biophys.* 449, 57–63.

Emsley, P., and Cowtan, K. (2004). Coot: model-building tools for molecular graphics. *Acta Crystallogr. D Biol. Crystallogr.* 60, 2126–2132.

Goedecke, K., Pignot, M., Goody, R.S., Scheidig, A.J., and Weinhold, E. (2001). Structure of the N6-adenine DNA methyltransferase M.TaqI in complex with DNA and a cofactor analog. *Nat. Struct. Biol.* 8, 121–125.

Guyot, J.B., Grassi, J., Hahn, U., and Guschlbauer, W. (1993). The role of the preserved sequences of Dam methylase. *Nucleic Acids Res.* 21, 3183–3190.

Helser, T.L., Davies, J.E., and Dahlberg, J.E. (1971). Change in methylation of 16S ribosomal RNA associated with mutation to kasugamycin resistance in *Escherichia coli*. *Nat. New Biol.* 233, 12–14.

Helser, T.L., Davies, J.E., and Dahlberg, J.E. (1972). Mechanism of kasugamycin resistance in *Escherichia coli*. *Nat. New Biol.* 235, 6–9.

Jones, T.A., Zou, J.Y., Cowan, S.W., and Kjeldgaard, M. (1991). Improved methods for building protein models in electron-density maps and the location of errors in these models. *Acta Crystallogr. A* 47, 110–119.

Klimasauskas, S., Kumar, S., Roberts, R.J., and Cheng, X. (1994). Hhal methyltransferase flips its target base out of the DNA helix. *Cell* 76, 357–369.

Klootwijk, J., Klein, I., and Grivell, L.A. (1975). Minimal post-transcriptional modification of yeast mitochondrial ribosomal RNA. *J. Mol. Biol.* 97, 337–350.

Kong, H., and Smith, C.L. (1997). Substrate DNA and cofactor regulate the activities of a multi-functional restriction-modification enzyme, Bcgl. *Nucleic Acids Res.* 25, 3687–3692.

Labahn, J., Granzin, J., Schluckebier, G., Robinson, D.P., Jack, W.E., Schildkraut, I., and Saenger, W. (1994). Three-dimensional structure of the adenine-specific DNA methyltransferase M.TaqI in complex with the cofactor S-adenosylmethionine. *Proc. Natl. Acad. Sci. USA* 91, 10957–10961.

Lafontaine, D., Delcour, J., Glasser, A.L., Desgres, J., and Vandenhaute, J. (1994). The Dim1 gene responsible for the conserved M(2)(6)Am(2)(6)a dimethylation in the 3'-terminal loop of 18-S ribosomal-RNA is essential in yeast. *J. Mol. Biol.* 241, 492–497.

Lafontaine, D., Vandenhaute, J., and Tollervey, D. (1995). The 18S ribosomal-RNA dimethylase Dim1p is required for pre-ribosomal-RNA processing in yeast. *Genes Dev.* 9, 2470–2481.

Lafontaine, D.L.J., Preiss, T., and Tollervey, D. (1998). Yeast 18S rRNA dimethylase Dim1p: a quality control mechanism in ribosome synthesis? *Mol. Cell. Biol.* 18, 2360–2370.

Laskowski, R.A., MacArthur, M.W., Moss, D.S., and Thornton, J.M. (1993). PROCHECK—a program to check the stereochemical quality of protein structures. *J. Appl. Crystallogr.* 26, 283–291.

Law, S.F., Zhang, Y.Z., Klein-Szanto, A.J., and Golemis, E.A. (1998). Cell cycle-regulated processing of HEF1 to multiple protein forms differentially targeted to multiple subcellular compartments. *Mol. Cell. Biol.* 18, 3540–3551.

Malone, T., Blumenthal, R.M., and Cheng, X. (1995). Structure-guided analysis reveals nine sequence motifs conserved among DNA amino-methyltransferases, and suggests a catalytic mechanism for these enzymes. *J. Mol. Biol.* 253, 618–632.

Maravic, G., Bujnicki, J.M., Feder, M., Pongor, S., and Flögel, M. (2003a). Alanine-scanning mutagenesis of the predicted rRNA-binding domain of ErmC' redefines the substrate-binding site and suggests a model for protein-RNA interactions. *Nucleic Acids Res.* 31, 4941–4949.

- Maravic, G., Feder, M., Pongor, S., Flogel, M., and Bujnicki, J.M. (2003b). Mutational analysis defines the roles of conserved amino acid residues in the predicted catalytic pocket of the rRNA:m6A methyltransferase ErmC'. *J. Mol. Biol.* **332**, 99–109.
- McCoy, A.J., Grosse-Kunstleve, R.W., Adams, P.D., Winn, M.D., Storoni, L.C., and Read, R.J. (2007). Phaser crystallographic software. *J. Appl. Crystallogr.* **40**, 658–674.
- McCulloch, V., Seidel-Rogol, B.L., and Shadel, G.S. (2002). A human mitochondrial transcription factor is related to RNA adenine methyltransferases and binds S-adenosylmethionine. *Mol. Cell. Biol.* **22**, 1116–1125.
- Mitchell, P., Osswald, M., and Brimacombe, R. (1992). Identification of intermolecular RNA cross-links at the subunit interface of the *Escherichia coli* ribosome. *Biochemistry* **31**, 3004–3011.
- Murshudov, G.N., Vagin, A.A., and Dodson, E.J. (1997). Refinement of macromolecular structures by the maximum-likelihood method. *Acta Crystallogr. D Biol. Crystallogr.* **53**, 240–255.
- Nallamsetty, S., and Waugh, D.S. (2006). Solubility-enhancing proteins MBP and NusA play a passive role in the folding of their fusion partners. *Protein Expr. Purif.* **45**, 175–182.
- O'Farrell, H.C., Scarsdale, J.N., and Rife, J.P. (2004). Crystal structure of KsgA, a universally conserved rRNA adenine dimethyltransferase in *Escherichia coli*. *J. Mol. Biol.* **339**, 337–353.
- O'Farrell, H.C., Pulicherla, N., Desai, P.M., and Rife, J.P. (2006). Recognition of a complex substrate by the KsgA/Dim1 family of enzymes has been conserved throughout evolution. *RNA* **12**, 725–733.
- Oh, J.H., Yang, J.O., Hahn, Y., Kim, M.R., Byun, S.S., Jeon, Y.J., Kim, J.M., Song, K.S., Noh, S.M., Kim, S., et al. (2005). Transcriptome analysis of human gastric cancer. *Mamm. Genome* **16**, 942–954.
- Otwinowski, Z., and Minor, W. (1997). Processing of X-ray diffraction data collected in oscillation mode. *Methods Enzymol.* **276**, 307–326.
- Poldermans, B., Roza, L., and Van Knippenberg, P.H. (1979a). Studies on the function of two adjacent N6,N6-dimethyladenosines near the 3' end of 16 S ribosomal RNA of *Escherichia coli*. III. Purification and properties of the methylating enzyme and methylase-30 S interactions. *J. Biol. Chem.* **254**, 9094–9100.
- Poldermans, B., Van Buul, C.P., and Van Knippenberg, P.H. (1979b). Studies on the function of two adjacent N6,N6-dimethyladenosines near the 3' end of 16 S ribosomal RNA of *Escherichia coli*. II. The effect of the absence of the methyl groups on initiation of protein biosynthesis. *J. Biol. Chem.* **254**, 9090–9093.
- Poldermans, B., Bakker, H., and Van Knippenberg, P.H. (1980). Studies on the function of two adjacent N6,N6-dimethyladenosines near the 3' end of 16S ribosomal RNA of *Escherichia coli*. IV. The effect of the methyl groups on ribosomal subunit interaction. *Nucleic Acids Res.* **8**, 143–151.
- Pues, H., Bleimling, N., Holz, B., Wolcke, J., and Weinhold, E. (1999). Functional roles of the conserved aromatic amino acid residues at position 108 (motif IV) and position 196 (motif VIII) in base flipping and catalysis by the N6-adenine DNA methyltransferase from *Thermus aquaticus*. *Biochemistry* **38**, 1426–1434.
- Rife, J.P., and Moore, P.B. (1998). The structure of a methylated tetraloop in 16S ribosomal RNA. *Structure* **6**, 747–756.
- Roth, M., Helm-Kruse, S., Friedrich, T., and Jeltsch, A. (1998). Functional roles of conserved amino acid residues in DNA methyltransferases investigated by site-directed mutagenesis of the EcoRV adenine-N6-methyltransferase. *J. Biol. Chem.* **273**, 17333–17342.
- Schluckebier, G., Kozak, M., Bleimling, N., Weinhold, E., and Saenger, W. (1997). Differential binding of S-adenosylmethionine S-adenosylhomocysteine and Sinefungin to the adenine-specific DNA methyltransferase M.TaqI. *J. Mol. Biol.* **265**, 56–67.
- Schluckebier, G., Zhong, P., Stewart, K.D., Kavanaugh, T.J., and Abad-Zapatero, C. (1999). The 2.2 Å structure of the rRNA methyltransferase ErmC' and its complexes with cofactor and cofactor analogs: implications for the reaction mechanism. *J. Mol. Biol.* **289**, 277–291.
- Schubot, F.D., Chen, C.J., Rose, J.P., Dailey, T.A., Dailey, H.A., and Wang, B.C. (2001). Crystal structure of the transcription factor sc-mtTFB offers insights into mitochondrial transcription. *Protein Sci.* **10**, 1980–1988.
- Schuwirth, B.S., Borovinskaya, M.A., Hau, C.W., Zhang, W., Vila-Sanjurjo, A., Holton, J.M., and Cate, J.H. (2005). Structures of the bacterial ribosome at 3.5 Å resolution. *Science* **310**, 827–834.
- Seidel-Rogol, B.L., McCulloch, V., and Shadel, G.S. (2003). Human mitochondrial transcription factor B1 methylates ribosomal RNA at a conserved stem-loop. *Nat. Genet.* **33**, 23–24.
- Skinner, R., Cundliffe, E., and Schmidt, F.J. (1983). Site of action of a ribosomal RNA methylase responsible for resistance to erythromycin and other antibiotics. *J. Biol. Chem.* **258**, 12702–12706.
- Thammana, P., and Held, W.A. (1974). Methylation of 16S RNA during ribosome assembly in vitro. *Nature* **251**, 682–686.
- Tokuhiya, J.G., Vijayan, P., Feldmann, K.A., and Browse, J.A. (1998). Chloroplast development at low temperatures requires a homolog of DIM1, a yeast gene encoding the 18S rRNA dimethylase. *Plant Cell* **10**, 699–711.
- Vagin, A.A., Steiner, R.A., Lebedev, A.A., Potterton, L., McNicholas, S., Long, F., and Murshudov, G.N. (2004). REFMAC5 dictionary: organization of prior chemical knowledge and guidelines for its use. *Acta Crystallogr. D Biol. Crystallogr.* **60**, 2184–2195.
- van Buul, C.P., and van Knippenberg, P.H. (1985). Nucleotide sequence of the ksgA gene of *Escherichia coli*: comparison of methyltransferases effecting dimethylation of adenosine in ribosomal RNA. *Gene* **38**, 65–72.
- Van Buul, C.P., Damm, J.B., and Van Knippenberg, P.H. (1983). Kasugamycin resistant mutants of *Bacillus stearothermophilus* lacking the enzyme for the methylation of two adjacent adenosines in 16S ribosomal RNA. *Mol. Gen. Genet.* **189**, 475–478.
- van Gemen, B., Twisk, J., and van Knippenberg, P.H. (1989). Autogenous regulation of the *Escherichia coli* ksgA gene at the level of translation. *J. Bacteriol.* **171**, 4002–4008.
- Van Knippenberg, P.H., Van Kimmenade, J.M., and Heus, H.A. (1984). Phylogeny of the conserved 3' terminal structure of the RNA of small ribosomal subunits. *Nucleic Acids Res.* **12**, 2595–2604.
- Weisblum, B. (1995). Erythromycin resistance by ribosome modification. *Antimicrob. Agents Chemother.* **39**, 577–585.
- Willcock, D.F., Dryden, D.T., and Murray, N.E. (1994). A mutational analysis of the two motifs common to adenine methyltransferases. *EMBO J.* **13**, 3902–3908.
- Wimberly, B.T., Brodersen, D.E., Clemons, W.M., Jr., Morgan-Warren, R.J., Carter, A.P., Vornrhein, C., Hartsch, T., and Ramakrishnan, V. (2000). Structure of the 30S ribosomal subunit. *Nature* **407**, 327–339.
- Xu, Z., O'Farrell, H.C., Rife, J.P., and Culver, G.M. (2008). A conserved rRNA methyltransferase regulates ribosome biogenesis. *Nat. Struct. Mol. Biol.* **15**, 534–536.
- Yu, L., Petros, A.M., Schnuchel, A., Zhong, P., Severin, J.M., Walter, K., Holzman, T.F., and Fesik, S.W. (1997). Solution structure of an rRNA methyltransferase (ErmAM) that confers macrolide-lincosamide-streptogramin antibiotic resistance. *Nat. Struct. Biol.* **4**, 483–489.