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Crystal structure of *Thermus thermophilus* tRNA m$^1$A$_{58}$ methyltransferase and biophysical characterization of its interaction with tRNA.

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Short running title : Biophysical characterization of the interaction of *th*TrmI with tRNA

The abbreviations used are: AdoHcy, S-adenosyl-L-homocysteine; AdoMet, S-adenosyl-L-methionine; ESI-MS, Electrospray ionization mass spectrometry; MTases, methyltransferases; TrmI, m$^1$A$_{58}$ tRNA methyltransferase; thTrmI, *Thermus thermophilus* TrmI; mycoTrmI, *Mycobacterium tuberculosis* TrmI.
Abstract

Methyltransferases (MTases) from the TrmI family catalyse the S-adenosyl-L-methionine (AdoMet)-dependent N$_1$-methylation of tRNA adenosine 58. The crystal structure of *Thermus thermophilus* m$^1$A$_{58}$ tRNA MTase (*th*TrmI) in complex with S-adenosyl-L-homocysteine was determined at 1.7 Å resolution. This structure is closely related to that of *Mycobacterium tuberculosis* TrmI (*myco*TrmI), and their comparison enabled us to enlighten two grooves in the TrmI structure that are large enough and electrostatically compatible to accommodate one tRNA per face of TrmI tetramer. We have then conducted a biophysical study based on electrospray ionization mass spectrometry, site-directed mutagenesis and molecular docking. First, we confirmed the tetrameric oligomerisation state of TrmI and we showed that this protein remains tetrameric upon tRNA binding with formation of complexes involving one to two molecules of tRNA per TrmI tetramer. Secondly, three key residues for the methylation reaction were identified, the universally conserved D170, and two conserved aromatic residues Y78 and Y194. We then used molecular docking to position a N$_9$-methyladenine in the active site of TrmI. The N$_9$-methyladenine snugly fits in the catalytic cleft where the side chain of D170 acts as a bidentate ligand binding the amino moiety of AdoMet and the exocyclic amino group of the adenosine. Y194 interacts with the N$_9$-methyladenine ring whereas Y78 can stabilize the sugar ring. From our results, we propose that the conserved residues that form the catalytic cavity (D170, Y78 and Y194) are essential to fashion an optimized shape of the catalytic pocket.

**Keywords:** TrmI, m1A58 methyltransferase, X-ray structure, noncovalent mass spectrometry, Protein-RNA interactions
Introduction

Functional tRNAs carry a number of chemically modified nucleosides that are formed enzymatically after transcription, during the tRNA maturation process. To date, more than 90 different modifications have been identified in tRNAs from various organisms. Among these post-transcriptional nucleoside modifications, N$_1$-methyladenosine (m$^1$A) is found at a highly conserved A$_{58}$ position in the TΨC loop of many tRNAs in the three domains of life (Bacteria, Archaea and Eukarya). This modification occurs infrequently in bacteria and, for instance, is absent in E. coli tRNAs. On the contrary, it is common in the tRNAs of most eukaryotes and archaea, and this modification seems to be essential for some organisms. Actually, in the yeast Saccharomyces cerevisiae, mutants defective in N$_1$-methylation of A$_{58}$ are non-viable. Likewise, in Thermus thermophilus, gene disruption studies have shown that m$^1$A$_{58}$ modification is required for growth of this bacterium at high temperatures.

S-Adenosyl-L-methionine (AdoMet) dependent methyltransferases (MTases) are involved in a wide variety of biological processes involving methylation of nucleic acids, proteins, phospholipids and small molecules using the ubiquitous methyl donor AdoMet. Recently, two essential genes of S. cerevisiae, GCD10 and GCD14 (renamed TRM6 and TRM61), were identified to encode the two types of subunits of the yeast m$^1$A$_{58}$ MTase. The recombinant enzyme behaves as an $\alpha_2\beta_2$ heterotetramer, Trm61p being responsible for AdoMet binding and presumably catalysis of the methyl transfer reaction, whereas both types of subunits are essential for tRNA-binding. Indeed, the purified recombinant Trm61p subunit, which binds AdoMet, cannot bind tRNA in vitro in the absence of Trm6p. Interestingly, Trm61p was found to be closely related to a group of prokaryotic proteins which share not only the same AdoMet-binding site but also other highly conserved motifs. Therefore, the corresponding prokaryotic proteins were presumed to act as m$^1$A$_{58}$ MTases. This hypothesis was demonstrated experimentally for the bacterial orthologs in Thermus thermophilus and Mycobacterium tuberculosis and for the archaeal ortholog in Pyrococcus abyssi. These proteins are $\alpha_4$ homotetramers of Trm61p-like proteins, hereafter called TrmI. Interestingly, orthologs of the Trm6p protein could only be found in eukaryotes. It was shown that Trm6p, despite the absence of the characteristic MTases motifs, is structurally and evolutionary related to Trm61p suggesting that the eukaryotic m$^1$A$_{58}$ MTases evolved by gene duplication and speciation to form a heteromultimeric protein, whereas their prokaryotic orthologs remained homomultimers.

Despite the low level of sequence similarity between the various families of AdoMet-MTases, most of them contain a structurally highly conserved catalytic AdoMet-binding domain organized in a Rossman-like fold (For a review see). Only one crystal structure of an m$^1$A$_{58}$ tRNA MTase has been reported to date, that of the M. tuberculosis Rv2118c protein (mycoTrmI) in complex with S-adenosyl-L-methionine. It shows that the catalytic C-
terminal domain is very similar to that of other AdoMet-dependent MTases whereas the N-terminal domain which is mainly composed of β-sheets is not found in other MTases of known structure.

In the present work, we have first solved the crystal structure of the m^1A_{s8} tRNA MTase from *T. thermophilus* (*thTrmI*) at 1.7 Å resolution in complex with S-Adenosyl-L-homocysteine (AdoHcy), the product after methyl transfer. We then focused our attention on the tRNA-binding properties of *thTrmI* using several biophysical techniques, including electrospray ionization mass spectrometry (ESI-MS), site-directed mutagenesis, steady-state kinetic assays, fluorescence spectroscopy and molecular docking. ESI-MS has demonstrated its particular suitability for the investigation of noncovalent complexes. More particularly, so-called noncovalent ESI-MS has been extensively used to characterize supramolecular assemblies involving oligonucleotides in interaction with drugs and proteins (For reviews see 14-17). As protein:RNA systems are partly driven by electrostatic-based interactions which are strongly enhanced in the gas phase, ESI-MS is well suited for their characterization. Indeed, numerous studies of protein-RNA complexes have revealed a strong agreement between mass spectrometric gas-phase measurements and results obtained by solution phase techniques. In our work, we used noncovalent ESI-MS: i) to unambiguously assess the oligomerisation state of *thTrmI*; ii) to monitor the *thTrmI* oligomerisation state upon tRNA binding and iii) to determine the stoichiometry of the *thTrmI*-tRNA complexes. In addition, a mutagenesis study was performed to identify residues potentially crucial for the methylation reaction. We thus showed that the universally conserved D170 residue, together with two conserved aromatic residues, Y78 and Y194, which line the catalytic pocket are key residues for the enzymatic catalysis. Actually, D170A and Y78A variants are severely altered in their catalytic efficiency. Finally, we have used molecular docking to position a N^6-methyladenine in the active site of TrmI to further investigate the possible role of these residues.
Results and Discussion

Structure determination of *T. thermophilus* TrmI

Production and purification of *T. thermophilus* TrmI (thTrmI) have been described by Droogmans and coworkers. However, the recombinant His<sub>6</sub>-tagged protein exhibited a low solubility (<0.5 mg/mL) unless high salt concentrations were added (0.2 M imidazole-HCl and 0.5 M KCl). This was a serious issue for crystallization assays, in which precipitating solutions must be added to concentrated protein. In order to improve solubility, the N-terminal His<sub>6</sub>-tag of thTrmI was cleaved with thrombin. Indeed, after His<sub>6</sub>-tag removal, thTrmI became highly soluble (>10 mg/mL), even at moderate ionic strengths (150 mM). This type of behaviour has already been reported for some His-tagged proteins.

We then carried out crystallization assays on the complex between thTrmI, AdoHcy and *E. coli* tRNA<sub>iMet</sub>. The latter tRNA was selected because it is devoid of modification on A<sub>58</sub> and is a substrate of the *T. thermophilus* enzyme (see paragraph on steady-state kinetic assays). Although the tRNA was present in all crystallization conditions, crystals only contained the free thTrmI protein in complex with AdoHcy. Table 1 summarizes the data collection and refinement statistics. The structure of *T. thermophilus* TrmI protein was then solved by molecular replacement with the structure of m<sup>1</sup>A<sub>58</sub> tRNA MTase from *M. tuberculosis* TrmI (PDB entry code 1I9G) using PHASER (Z score = 23.7). Actually, thTrmI shares 39% identical and 13% similar residues with mycoTrmI. The model was then refined to 1.7 Å, to a final R<sub>free</sub> factor of 20.9% (Table 1). The last four residues at the N-terminus were too poorly ordered to be included, and therefore the final model consisted of 251 residues per monomer. The asymmetric unit contained a dimer of thTrmI. However, the symmetry-related subunits interact extensively leading to a tetrameric organization of thTrmI (Figure 1(b)).

Overall structure of thTrmI

The thTrmI monomer is composed of two distinct domains connected by an extended linker containing a short α-helix (helix 2, Figure 1(a)). The large C-terminal domain (residues 63-255) adopts a typical class I MTase fold which consists of a central seven-stranded β sheet (β<sub>1</sub> to β<sub>6</sub> and β<sub>8</sub> – Figure 1(a)) flanked by α helices on both sides (α<sub>A</sub> to α<sub>E</sub> – Figure 1(a)). The first five strands of the β sheet (β<sub>1</sub> to β<sub>5</sub> on Figure 1(a)) are parallel whereas the last two (β<sub>6</sub> to β<sub>8</sub>) are antiparallel. The C-terminal domain contains the AdoMet binding site and the catalytic pocket. The smaller N-terminal domain (residues 5-62) is largely a β structure composed of six β-strands (β<sub>1</sub> to β<sub>6</sub> Figure 1(a)) and one small α helix (α<sub>1</sub>).

The thTrmI tetramer is organised as a dimer of tight dimers. Two monomers interact via their extended β<sub>6</sub> strands which form a cross-subunit, antiparallel β-sheet (Figure 1(b), pale and dark blue). This dimer is further stabilized by ionic interactions between helices α<sub>2</sub> and α<sub>3</sub> from each subunit (Figure 1(b)). A pair of β<sub>6</sub> strands from each dimer packs on the related
β₆ strands of the other dimer to form a central anti-parallel β-sheet structure. The tetramer interaction is limited to this β-sheet region and stabilized by both ionic and hydrophobic interactions, mainly a salt bridge between E220 and R224, hydrophobic interactions between W226 and F245, and between H234 and F237 from one dimer to H242 from the other dimer. The salt bridge and the hydrophobic interactions between W226 and F245 have been also observed on the mycoTrmI structure⁷ and involve conserved residues.

In the tight dimer found in the asymmetric unit, the two cofactor-binding sites are occupied by AdoHcy molecules in identical conformations, similar to that of AdoMet in the M. tuberculosis TrmI structure. The binding cleft is formed by residues belonging to conserved motifs, named I, II, III, IV, (Figures 1(c), 1(d) and 7(a)).⁷ The adenine ring of the AdoHcy moiety is bound by motifs II, III, and IV via water-mediated hydrogen bonds between the exocyclic amino group and residues E155 and E173, and between the N₇ position of the adenine and the backbone amide of E173 (Figure 1(c)). Adenine binding is also reinforced by hydrophobic contacts between its aromatic ring and residues L171, V177, K153, L154, Y124, A126 and A100. The ribose moiety of AdoHcy is bound by motif II, essentially by hydrogen bonds involving the 2’ and 3’ hydroxyl groups and residues E125 and H130. The amino-acid part of the cofactor is mainly bound by motif I. Interestingly, an aspartate side chain that could be involved in the methylation reaction, the carboxylate chain of D170 from motif IV, participates to a direct polar interaction with the ammonium group of AdoHcy. The GXGXGG pattern of motif I binds to the amine and the carboxylate through a tight network of water-mediated hydrogen bonds involving the backbone (Figure 1(c)).

**Comparison with mycoTrmI structure and tRNA recognition**

Globally, the overall structure of thTrmI closely resembles that of mycoTrmI. Superimposition of T. thermophilus and M. tuberculosis TrmI structures gives a root mean square deviation of 4.47 Å for Ca, the largest variation occurring in the β-strands (β₆, β₇, β₈) that protrude out of the structure. When comparing the active sites more accurately, that of mycoTrmI is narrower by about 1 Å than that of thTrmI. Moreover, the protruding R249 in mycoTrmI makes like a supplementary step at the exit of the pocket that does not exist in thTrmI as R249 is replaced by a shorter side chain residue (V240). As a consequence, the accessible surface of Y78 is twice larger in thTrmI than that of Y84 in mycoTrmI. All these features seem to make the methyl donor less easily accessible in mycoTrmI. Overall, in both structures, the adenine should enter deeply in the binding cleft to have access to the methyl donor therefore implicating a major deformation of the phosphodiester backbone around A₅₈. It is also highly interesting to compare electrostatic potentials of their molecular surfaces (Figure 2). These calculations uncover that both proteins have two linked grooves of positive electrostatic potentials indicated by large blue surfaces on Figure 2 that are large enough to accommodate double-helical RNA. The first groove that covers the β-sheet region forming
the tetrameric architecture is less positively charged in *th*TrmI than in *myco*TrmI; whereas the second groove that encompasses the N-terminal domain of one TrmI subunit and that looks like an open hand is more positively charged in *th*TrmI than in *myco*TrmI. Several residues with basic side chains are conserved among the TrmI protein family, namely K13 (R in some sequences), R15 (K), R72, R127, R217 and R229 (K) together with the histidines involved in the formation of the tetramer H234 and H242. R15 and R217 are only conserved in bacterial TrmI proteins. These grooves have the dimensions to accommodate the acceptor arm (groove 1, dimensions: 25 Å large and 45 Å long) and the anticodon one (groove 2, dimensions: 25 Å large and roughly 30 Å long) without steric clashes, enabling us to position the adenosine 58 near the catalytic pocket (manual docking, data not shown). The grey line in Figure 2(a) indicates a possible position of the helix axis of tRNA. Numerous clashes occur within the T-arm that undoubtedly undergoes huge conformational changes that probably involves the entire T-arm and not only a simple flip of the adenosine 58. These changes are difficult to predict and, for this reason, we could not pursue this docking. In conclusion, the conservation of two grooves of positively charged surfaces supports the idea that TrmI binds tRNA as a tetramer and that two molecules of tRNA can interact simultaneously with the TrmI tetramer. The fact that, in eukaryotes, the enzyme has evolved from an $\alpha_4$ homotetramer to an $\alpha_2\beta_2$ heterotetramer is compatible with a two tRNA per TrmI tetramer stoichiometry, one tRNA being expected to interact with an $\alpha\beta$ subsystem.

*th*TrmI is a tetramer and binds up to two tRNA molecules as a tetramer.

Close inspection of the crystal structures of *th*TrmI and *myco*TrmI revealed that the tetramers are formed by two pairs of extensively interacting subunits stabilized by relatively small numbers of contacts between the two dimers. Given the architecture of TrmI, it was of interest to investigate the oligomerization state of *th*TrmI alone and upon tRNA binding, and to determine the binding stoichiometry of the *th*TrmI-tRNA complexes. Previously reported gel filtration experiments have indicated that both *myco*TrmI and *th*TrmI should form tetramer in solution. However, in the case of *th*TrmI, due to an inherent limited precision of this technique, it is difficult to unambiguously state whether the protein is tetrameric or pentameric. In this study, we used ESI-MS, which has been proven to be a valuable technique for the determination of oligomerization states of noncovalent assemblies to first confirm the oligomerization state of *th*TrmI. Figure 3(a) presents the ESI mass spectrum obtained for *th*TrmI under non-denaturing conditions and carefully controlled operating conditions (see Materials and Methods): a single ion distribution is observed in the m/z 4000-5000 range. With a measured molecular weight (MW) of 115450 ± 5 Da, this distribution can be assigned to the +23 to +30 charge states of a *th*TrmI tetramer (theoretical MW = 115453 Da). Thus, ESI-MS allowed us to unambiguously assess the tetrameric nature of the protein in agreement with our crystallographic results.
The tRNA binding stoichiometry was first investigated by gel retardation assay. As shown on Figure 4, the free tRNA band progressively disappears upon addition of increasing amounts of thTrmI evidencing thus the existence of thTrmI-tRNA interactions. Moreover, the free tRNA band completely disappears at 1:2 thTrmI:tRNA ratio, substantiating the formation of complexes involving two molar equivalents of thTrmI monomer per tRNA. To further confirm gel retardation experiments, we also used the potentialities of noncovalent ESI-MS to investigate the oligomerization state of thTrmI upon tRNA binding and to determine the tRNA binding stoichiometry. Titration experiments, monitored by noncovalent ESI-MS, involving a fixed concentration of thTrmI and increasing amounts of tRNA$^{\text{Met}}$, revealed the presence of three ion distributions (Figure 3(b-d)). The first one in the m/z 4000-5000 region with a molecular mass of 115450 ± 5 Da is related to the thTrmI tetramer. For the second ion distribution in the m/z 5000-5900 region, the molecular mass of 140347 ± 6 Da corresponds to the thTrmI$_4$:(tRNA)$_1$ complex. Finally, a third ion distribution within the m/z 5900-6800 range with a molecular mass of 165455 ± 10 Da refers to the thTrmI$_4$:(tRNA)$_2$ complex. Even a ten-fold molar excess of tRNA per thTrmI tetramer does not lead to the detection of complexes with higher tRNA binding stoichiometries. Interestingly, upon increase of the tRNA concentration from 1.5 to 10 molar equivalents per thTrmI tetramer, the relative abundances of thTrmI$_4$:(tRNA)$_1$ and thTrmI$_4$:(tRNA)$_2$ complexes statistically increase, which is in favour of non-cooperative tRNA binding system and further suggests the presence of two independent tRNA binding sites. The same experiments were also carried out in presence of AdoHcy (data not shown) and no effect either on the binding stoichiometry of the complex or on the binding affinity of the tRNA for thTrmI was observed.

As ESI-MS detects species in the gas phase of the mass spectrometer, control experiments are always needed to ensure that mass spectra faithfully reflect the behaviour in solution. Thus, since electrostatic interactions are sensitive to the ionic strength of the medium, experiments were carried out at different ammonium bicarbonate concentrations (Figure 5). Decreasing the ammonium bicarbonate concentration from 200 mM to 100 mM displaces the equilibrium towards the formation of thTrmI$_4$:(tRNA)$_1$ and thTrmI$_4$:(tRNA)$_2$ complexes (Figure 5(a-b)). However, the signal intensities were dramatically lower when 100 mM ammonium bicarbonate was used rather than 200 mM buffer, mainly because of protein precipitation. Therefore, no ESI-MS analysis could be performed at buffer concentrations lower than 100 mM. While both thTrmI$_4$:(tRNA)$_1$ and thTrmI$_4$:(tRNA)$_2$ complexes are favoured at low ionic strengths, the use of higher salt concentrations destabilizes these assemblies (Figure 5(c-d)), leading even to a complete dissociation of thTrmI$t$-tRNA complexes into thTrmI tetramer at 1250 mM ammonium bicarbonate (Figure 5(d)). The fact that mass spectra do reflect expected changes induced by modification of the solution conditions is a further evidence that allows us to definitely rule out the possibility that the TrmI-tRNA complexes result from an artefact of the technique. Moreover, as described in
other publications, an additional ESI-MS control experiment was performed in strictly identical experimental and operating conditions with a non-substrate RNA as a negative control. The absence of any thTrmI-control RNA complex on ESI mass spectra even in presence of a five-fold molar excess of control RNA (Figure 6) indicates that the gas phase detection of the TrmI-tRNA complexes arises from a specific recognition in solution and not from any gas phase artefact.

In conclusion, noncovalent ESI-MS results clearly support that thTrmI remains tetrameric upon tRNA binding and that thTrmI binds up to two molecules of tRNA.

The D170A and Y78A variants of thTrmI are severely altered in their catalytic efficiency.

On the basis of multiple sequence alignments of the TrmI family members (Figure 7(a)) and on the comparison of the crystallographic structures of TrmI from M. tuberculosis and T. thermophilus, we chose to mutate three conserved residues, D170, Y194 and Y78 in order to study their involvement in adenosine binding and/or in catalysis. Figure 7(b) presents the thTrmI residues that are conserved or semi-conserved in the catalytic pocket, except the conserved residues that bind to AdoMet since their role in the catalysis mechanism is known. Yet, the universally conserved AdoMet-binding D170 was selected as it forms the back of the catalytic pocket and may also bind the adenine 58 ring of the tRNA substrate (Figure 7(b)). Y194 and Y78 were mutated because they respectively form the left side and the floor of the adenosine-binding pocket in the 3D structure (Figure 7(b)) and are conserved as aromatic residues among m1A58 tRNA MTases, belonging respectively to motifs V and X (Figure 7(a)). Therefore, these aromatic residues could be involved in the stabilization of the target adenine that needs to be flipped out of the tRNA structure to be methylated. P196 that is involved in making the shape of the catalytic cavity was not selected for mutation, because in M. tuberculosis, it is naturally replaced by an alanine. Therefore, we individually mutated Y78, D170 and Y194 to alanines. We expressed and purified the variant proteins as described in Materials and Methods section. We then determined the kinetic and RNA-binding parameters for wild-type and variant proteins (Table 2).

First, we investigated whether the mutations altered tRNA binding by the enzyme. Interaction with tRNA induces a quenching of the intrinsic tryptophan fluorescence of the protein of about 40 %. We therefore used fluorimetric titrations to determine the apparent dissociation constant between thTrmI variants and tRNAiMet (Table 2). All mutants showed quite unchanged dissociation constants (K_d about 15 ± 3 nM), thereby indicating that they retained a native folding and similar RNA binding ability compared to the wild-type enzyme. We then analysed their catalytic properties by determining their kinetic parameters kcat and KM for both AdoMet and tRNA substrates. The overall results are summarized in Table 2. Interestingly, all mutants retain some catalytic activity, but are altered to various extents,
essentially in their catalytic turnover $k_{cat}$. The D170A mutant retains some activity, but its $k_{cat}$ value is severely reduced, by a factor of about 300. D170A also shows an increased $K_M$ for AdoMet substrate in agreement with the crystal structure. It seems therefore to be the most important residue for the catalysis of the reaction. The Y78A mutant leads to a roughly 20-fold decrease of the catalytic turn-over constant. Its binding constant for the methyl donor is unchanged compared to the wild-type protein. The Y194A mutant shows a smaller decrease of $k_{cat}$ by a factor of 3 (Table 2). The $K_M$ value for AdoMet is increased suggesting that this residue is involved in cofactor binding. Yet, in the crystal structure, it is not involved in a direct interaction with AdoHcy in contrast to D170. The mutation of Y194 to an alanine probably slightly destabilizes the catalytic pocket or indirectly weakens AdoMet binding. In conclusion, D170A is the variant that is most largely altered in its catalytic efficiency followed by Y78A and then Y194A. And, therefore, D170, Y78 and Y194 are key residues in the catalytic mechanism.

**The substrate adenine snugly fits in the active site pocket of thTrmI.**

To investigate the positioning of the target A$_{58}$ into the catalytic pocket of *T. thermophilus* and *M. tuberculosis* TrmI, we decided to locate a N$_9$-methyladenine in the catalytic pocket of these proteins by molecular docking. We docked a N$_9$-methyladenine and not an adenosine or an AMP molecule since it is impossible to predict the conformation of the sugar of this flipped-out nucleotide. Moreover Y78 that makes the exit of the catalytic pocket and which may interact with the ribose of the adenosine 58 can also flip to prevent from steric clashes with the A$_{58}$ ribose. As the molecules of AdoHcy present in the active sites of thTrmI have identical conformations to the AdoMet co-factor present in mycoTrmI, we modelled an AdoMet molecule in the catalytic pocket of thTrmI by superimposing the AdoMet from mycoTrmI with AdoHcy in thTrmI. We used the program HADDOCK$^{37}$ that can take into account mutagenesis results by introducing data as Ambiguous Interaction Restraints (AIRs) to drive the docking process. An AIR is defined as an ambiguous distance between all residues shown to be involved in the interaction. Ambiguous intermolecular distance restraints were applied between D170, Y194, Y78 and the AdoMet cofactor on the one hand, and the N$_9$-methyladenine, on the other hand. A distance restraint between the N$_1$ atom of N$_9$-methyladenine and the methyl of AdoMet was also added. The model resulting from the docking with thTrmI is presented in Figure 8(a) and shows that the N$_9$-methyladenine fits snugly in the cavity where methylation takes place. The same result was obtained for mycoTrmI (data not shown). Moreover, the methyl group introduced to substitute the ribose is pointing above Y78 (Figure 8(b)) enabling us to add a ribose without steric hindrance. Therefore, Y78 is likely to strongly interact with the ribose of the adenosine 58 probably participating to stabilization of the flipped-out conformation. The distance between the N$_9$-methyladenine N$_1$ and the AdoMet methyl group is 2.3 Å. This corresponds to a distance
between the N9-methyladenine N1 and the AdoMet sulphur atom of 4.1 Å. This is in keeping with distances observed in co-crystal structures of other endocyclic purine N-MTases (distances between 3.6 Å and 4.5 Å). The docked model of the N9-methyladenine ring suggests a binding mode involving essentially Van der Waals interactions with Y194, and a polar contact between the exocyclic amino group of the purine and the carboxylate of D170 (Figure 8(b)). Therefore, the side chain of D170 could act as a bidentate ligand binding the amino terminal moiety of AdoMet and the N6 exocyclic amino group of the adenine ring substrate. Thus, the sulphur-methyl bond of AdoMet is almost coplanar with the N9-methyladenine ring and is pointing to the same direction as the N1 lone-pair orbital, which can be of significant importance for the chemical mechanism. We thus propose that D170 could minimize the transition state energy by positioning both the AdoMet and the target base in an optimal orientation for the methylation reaction. D170 could also deprotonate the N6 group to activate the N1 lone-pair orbital in order to attack the methyl group of AdoMet and to generate the exocyclic imino tautomer of N1-methyladenosine as the initial product. The nucleophilic attack leads to the m1A basic form which has to be subsequently protonated to give the m1A cationic form, stable at physiological pH (pKa = 9.3). As the deprotonation of N6 of m1A is easier than the deprotonation of N6 of the non-methylated adenine ring (pKa = 16.7), it seems more reasonable that the deprotonation and the nucleophilic attack occur in the same elementary step. The model perfectly situates D170 to serve that role, which is most definitely consistent with the kinetic results.

The chemical mechanism of adenine N1 methylation of RNA had not yet been investigated in contrast to the N1 methylation of guanine. In the case of m1G RNA methyltransferase (TrmD), an aspartate residue was shown to be involved in the deprotonation step of N1 of guanine. The two processes are likely to be different, as under physiological conditions, the N1 of guanine is protonated but that of adenine is not. DNA methyltransferases have been extensively studied, but the m1A modification is not encountered in DNA. The methylation of the N6 position of adenine has been investigated in DNA (for structural studies see) and more recently in rRNA. Interestingly, the main catalytic residue in m6A RNA or DNA MTases has been clearly identified by mutagenesis studies to be the D/N/S residue of the conserved pattern (D/N/S)PP(Y/F/W) which belongs to MTase motif IV (for reviews on DNA MTases see). In known structures of m6A DNA MTases (for example M•TaqI, PDB code 2ADM) or m6A RNA MTases (ErmC’, PDB code 1QAN) in complex with cofactor (AdoMet or AdoHcy), the position of the D/N/S residue of motif IV relative to the cofactor is closely similar to that of D170 residue in the T. thermophilus TrmI crystal structure. Additionally, two other residues have been identified to be important in catalysis in m6A DNA methyltransferases. These residues are aromatic and stabilize the flipped-out target base and/or act as cation-π catalysts. The first one is the aromatic Y/F/W residue of motif IV, which is proposed to stabilize the cationic transition state by cation-π interaction.
second one is the aromatic residue Y/F/W of motif VIII, which stabilizes the flipped base outside the DNA helix.\textsuperscript{49,50} These aromatic residues are conserved in motifs IV and VIII in m\textsuperscript{1}A tRNA MTases (Figure 7(a), symbol o), but they are not part of the catalytic pocket in the TrmI structure. Strikingly, two other aromatic residues, i.e. Y78 and Y194, are located in the active site cavity of thTrmI (Figure 7(b)) at relative positions with respect to the AdoMet cofactor similar to those of the aromatic residues listed above for DNA MTases. These residues are also conserved as aromatic residues among m\textsuperscript{1}A\textsubscript{58} tRNA MTases and belong respectively to motifs X and V (Figure 7(a)). The noticeable structural similarity between the catalytic pocket of m\textsuperscript{1}A tRNA MTases and m\textsuperscript{6}A DNA MTases suggests a similar role in catalysis for these residues. Y78 is thus the most probable candidate for the stabilization of the flipped-out adenosine given its position in the catalytic pocket and the dramatic decrease of $k_{cat}$ of the Y78A mutant. Then, Y194 could act as a cation-\pi catalyst to stabilize a cationic transition state. These roles are also in agreement with our docking results.

Coordinates and structure factors for \textit{T. thermophilus} TrmI protein in complex with AdoHcy have been deposited at the Protein Data Bank with accession code 2PWY.
Materials and Methods

Expression and purification of thTrmI for structural studies

Recombinant *T. thermophilus* TrmI protein (255 residues) was overexpressed and purified as previously described. The N-terminal His₆-tag was removed by thrombin cleavage (25 units of thrombin per mg of protein) performed overnight at ambient temperature. The thTrmI protein was then further purified by gel filtration on a HiLoad 26/60 Superdex 200 prepgrade chromatography column (Amersham Biosciences) equilibrated in 50 mM Tris-HCl buffer (pH 8.0), 100 mM KCl. Fractions containing pure recombinant protein were pooled, concentrated to 8-10 mg/mL using Amicon® Ultra (Millipore) and stored at 4 °C.

Crystallization and crystal structure determination

Crystallization was performed at 19 °C using the sitting drop vapour diffusion method. Protein samples were prepared at 3 mg/mL in 50 mM Tris-HCl buffer (pH 8.0) containing 100 mM KCl, 2 mM AdoHcy and 1 equivalent of trNA².Met. Drops of 1 µL were prepared using a Cybi-Disk robot system that mixes equal volume of protein and reservoir solutions. Reservoir volumes of 100 µL were used. Crystals of thTrmI were obtained in 2 M ammonium sulphate, isopropanol 5 % (v/v) (condition number 5 of Hampton Research Crystal Screen II kit). The crystals were harvested, soaked in a cryoprotectant solution (2.2 M ammonium sulphate, isopropanol 5 % (v/v), glycerol 20 % (v/v)) and flash-frozen in liquid nitrogen before data collection. Diffraction data were collected at beam line ID14-1 of the European Synchrotron Radiation Facility (ESRF, Grenoble, France).

All crystallographic calculations were performed using the CCP4 suite version 6 as implemented in the graphical user interface. X-ray diffraction data were processed using MOSFLM and scaled with SCALA. The structure of thTrmI was solved by molecular replacement using the program PHASER and the structure of m¹A₅₈ tRNA MTase from *M. tuberculosis* (PDB entry code 1I9G) as a model. In the model, non-conserved residues were truncated to alanine. Model building of thTrmI was first performed with ARP/wARP using the warpNtrace automated procedure. Restrained refinements of the structure were performed with the program REFMAC5. Model and maps visualisations for manual reconstruction were performed with the program COOT. Solvent molecules were automatically added using ARP waters module implemented in REFMAC5. In the last stages of refinement, TLS parameters were refined using one group for each domain of the protein, *i.e.* the large C-terminal domain and the small N-terminal domain.

We computed the electrostatic surface potential using APBS Tools, an interface for performing APBS electrostatics calculation from within PyMOL (http://www.pymol.org).
Expression and purification of tRNA$_{i}^{\text{Met}}$

The *E. coli* tRNA$_{i}^{\text{Met}}$ was overexpressed from plasmid pBSTtRNA$_{i}^{\text{Met}}$ in *E. coli* JM101TR using a protocol derived from that of Meinnel and Blanquet.$^{60}$ Briefly, after phenol extraction of RNAs from bacteria, total tRNA was separated by gel filtration on a HiLoad 26/60 Superdex 75 prepgrade chromatography column (Amersham Biosciences) equilibrated in 25 mM Tris-HCl pH 7.0. The overexpressed tRNA$_{i}^{\text{Met}}$ was then separated from other tRNAs by an anion exchange step (Resource Q column) equilibrated in 25 mM Tris-HCl pH 7.0. tRNAs were eluted using a 350 mM to 550 mM NaCl gradient in the same buffer. The fractions containing the purified tRNA$_{i}^{\text{Met}}$ were pooled together, dialysed against 50 mM Tris-HCl, 100 mM KCl, pH 8.0, concentrated using Amicon® Ultra (Millipore) and stored at -20 °C.

Electrospray ionization mass spectrometry

Noncovalent mass spectrometry experiments were performed on an electrospray time-of-flight mass spectrometer (LCT, Waters). Samples were continuously infused into the mass spectrometer with a syringe pump (Harvard Apparatus) at a flow rate of 6 μl/min. Mass spectra were acquired in the positive ion mode on the mass range 2000-8000 m/z after calibration with the multiply charged ions produced by horse heart myoglobin diluted to 2 μM in a 1:1 water/acetonitrile mixture (v/v) acidified with 1% (v/v) formic acid. Deconvoluted mass spectra were obtained using the Transform algorithm of MassLynx 4.0 software. Instrumental parameters were optimized to get the best compromise between ion desolvation, ion transmission and preservation of noncovalent complexes during the ionization and desorption processes. This optimization concerned especially the pressure in the interface (Pi) and the accelerating voltage (Vc) which were set to 6.1 mbar and 120 V, respectively.

Purity and homogeneity of *th*TrmI were checked in denaturing conditions by diluting the protein to 5 μM in a 1:1 water/acetonitrile mixture acidified with 1% (v/v) formic acid. A good agreement was found between the measured molecular mass (28863.0 ± 0.3 Da) and the mass calculated from the amino acid sequence (28863.3 Da). For experiments performed in non-denaturing conditions, different ammonium bicarbonate concentrations ranging from 100 mM to 1250 mM were tested. A concentration of 200 mM ammonium bicarbonate was chosen as the best compromise to provide good quality ESI mass-spectra without extensive *th*TrmI:tRNA complexes dissociation (which takes place at high ionic strengths) or protein aggregation (occurring at low ionic strengths). Thus, this buffer concentration was used to study both *th*TrmI oligomerization state and its tRNA-binding properties. All experiments were carried out at 20 μM *th*TrmI (tetramer concentration) whereas *E. coli* tRNA$_{i}^{\text{Met}}$ concentration (molecular weight MW = 24888 Da) was increased up to 200 μM for titration experiments. To study the influence of AdoHcy on the *th*TrmI-tRNA complexation equilibrium, *th*TrmI and AdoHcy, both at 20 μM (tetramer concentration), were incubated in
the ammonium bicarbonate buffer in presence or in absence of tRNA (30 µM). Finally, a non-substrate control RNA of 59 nucleotides (MW = 19255 Da) was also used in a five-fold molar excess relative to the thTrmI tetramer concentration in order to investigate the specificity of the thTrmI-tRNA interactions. All solutions were incubated for 15 minutes at room temperature before analysis under non-denaturing conditions.

**Non-denaturing agarose gel electrophoresis**

Samples were prepared by mixing constant amount of *E. coli* tRNA$_{i}^{\text{Met}}$ (4 µg) with increasing amounts of thTrmI protein (from 0 to 1 molar equivalent of thTrmI tetramer) in loading buffer (Tris-HCl pH 8.0 50 mM, KCl 100 mM, glycerol 18 % (v/v)). Samples were kept at 4 °C for 12 hours and loaded on a 2 % (w/v) agarose gel. Migration was performed for one hour in Tris-Acetate buffer pH 8.0 at 4 °C under an electric field of 100 V. Bands were visualized by UV shadowing.

**Site-directed Mutagenesis**

Alanine site-directed mutagenesis was carried out using the Quickchange® kit and protocol (Stratagene). All oligonucleotides for mutagenesis were designed to have appropriate pairing stability (fusion temperature near 80 °C) and were between 27 to 32 nt long. All mutant genes were fully sequenced and contained only the desired substitutions.

**Expression and purification of wild-type and mutant thTrmI proteins for kinetic studies**

The wild type and mutant proteins were overexpressed and purified as for structural studies except that the thrombin cleavage was omitted. Protein concentrations were evaluated with the Bradford method. The proteins were then diluted in 50 mM Tris-HCl pH 8.0 to a final concentration of 0.25 mg/mL and stored at 4 °C.

**Fluorescence titrations of thTrmI by tRNA$_{i}^{\text{Met}}$**

Fluorescence measurements were performed at 20 °C on a JASCO spectrofluorimeter. Excitation and emission wavelengths were 280 nm and 343 nm respectively. The excitation and emission bandwidths were 5 nm and 10 nm respectively. Fluorescence titrations experiments were performed by adding increasing concentrations of nucleic acid to a fixed amount of thTrmI protein (15 nM) in 50 mM Tris-HCl, 200 mM KCl, 0.5 % (w/v) PEG 8000, pH 8.0. Fluorescence intensities were corrected for dilution and were fitted using Eq. (1) assuming two equivalent tRNA binding sites per thTrmI tetramer (see Mass Spectrometry Results).

$$I = I_0 - I_0 - I_0 \left( \frac{K_d + L_r + nN_r}{2nN_r} - \sqrt{(K_d + L_r + nN_r)^2 - 4L_r nN_r} \right)$$
where $I_0$, Fluorescence intensity without tRNA; $I$, fluorescence intensity at a given concentration of tRNA; $I_\infty$, fluorescence intensity at the plateau; $n$, number of tRNA binding sites on the protein; $N_t$, total concentration of protein; $L_t$, total concentration of tRNA. Confidence limits on the parameters were estimated by Monte-Carlo sampling.61

**Steady-state kinetic assay**

[Methyl-$^3$H]-AdoMet (555 GBq/mmol, 20.5 MBq/mL, MP Biomedicals) was mixed with non-radioactive AdoMet (Sigma) to achieve a specific radioactivity of about 500 cpm/pmol (16.7 Bq/pmol). The methylation kinetic assays were performed in 50 mM Tris-HCl pH 8.0 at 60 °C. The tRNA and [methyl-$^3$H]-AdoMet were thermally equilibrated in 1.25 x concentrated reaction buffer (96 µL) at 60 °C for 5 min, and the reaction was started by adding the enzyme (24 µL). Aliquots (25 µL) were removed after different incubation times and transferred into 800 µL of 5 % (w/v) trichloroacetic acid (TCA) at 0 °C for 30 min in order to quench the reaction and to precipitate the tRNA. The precipitates were collected by filtration through GF/C filters (Whatman). The filters were washed with cold 5 % TCA, dried, and the radioactivity was measured by liquid scintillation counting for 2 min, resulting in a counting error below 4 %. Data were corrected by subtracting the background radioactivity determined from a control without enzyme.

For the determination of AdoMet $K_M$ and $k_{cat}$, the reaction mixtures contained MTase (wild type, 25 nM ; Y194A, 50 nM ; Y78A, 100 nM ; D170A, 500 nM), tRNA$_{iMet}$ (20 µM) and $^3$H-AdoMet (wild type, 0.5-8 µM ; Y194A, 0.5-8 µM ; Y78A, 0.5-8 µM ; D170A, 5-50 µM). For tRNA$_{iMet}$ $K_M$ and $k_{cat}$, the reaction mixtures contained MTase (wild type, 25 nM ; Y194A, 50 nM ; Y78A, 100 nM ; D170A, 500 nM), tRNA$_{iMet}$ (wild type, 0.5-8 µM ; Y194A, 0.5-8 µM ; Y78A, 0.25-4 µM ; D170A, 0.5-8 µM) and $^3$H-AdoMet (wild type, 20 µM ; Y194A, 50 µM ; Y78A, 20 µM ; D170A, 50 µM). Initial rates ($v_i$) for each substrate concentration (tRNA$_{iMet}$) were determined from the slope of linear fits of time course data points (wild type, 9 min ; Y194A, 20 min ; Y78A, 40 min ; D170A, 40 min). Enzyme parameters were obtained by non-linear least square fitting using Eq. (2) of Michaelis-Menten kinetics (wild type and mutant Y194A) or Eq. (3) in cases where enzyme concentration was not negligible compared to substrate concentration (mutant Y78A and D170A). $k_{cat}$ values were calculated for two catalytic sites per TrmI tetramer. Confidence limits on the parameters were estimated by Monte-Carlo sampling using the MC-Fit program.61

\[
(2) \quad v_i = \frac{V_{max}S_0}{K_M + S_0} \\
(3) \quad v_i = \frac{V_{max}}{2E_0} \left( K_M + S_0 + E_0 - \sqrt{(K_M + S_0 + E_0)^2 - 4S_0E_0} \right)
\]
Molecular Docking

Docking of the \( N_9 \)-methyladenine ring into the active site of TrmI was performed using HADDOCK 1.3\textsuperscript{37} (http://www.nmr.chem.uu.nl/haddock/) that makes use of CNS\textsuperscript{62} as structure calculation software. HADDOCK allows to deal with ambiguous constraints, i.e. restraints can be applied between residues and not only between atoms. The protein structure used for the docking was the tetrameric \textit{T. thermophilus} TrmI crystal structure in which the AdoHcy ligand was replaced with the AdoMet methyl donor. A 2 Å distance was used to define ambiguous restraints (AIRs) that were applied between the adenine and a set of neighbours composed of AdoMet cofactor and \textit{thTrmI} residues identified by mutagenesis (i.e. Y78, D170 and Y194). One unambiguous restraint set to 2 Å was applied between the atoms that react in the methyl transfer reaction, i.e. the \( N_1 \) position of the \( N_9 \)-methyladenine ring and the \( C_\varepsilon \) atom of the AdoMet ligand. During the rigid body energy minimisation, 2000 structures were calculated and the 1000 best solutions based on the intermolecular energy were used for the semi-flexible refinement. The docking converged to a single solution. The same protocol was also applied to the \textit{M. tuberculosis} TrmI.
REFERENCES


ACKNOWLEDGMENTS

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FIGURE LEGENDS

Figure 1: Crystal Structure of *T. thermophilus* TrmI.
(a) Overall structure of thTrmI monomer. The secondary structure elements are labelled following the nomenclature defined in Schluckebier *et al.* The cofactor AdoHcy is shown in blue sticks. (b) Organization of thTrmI as a tetramer. Each monomer is drawn with a different colour. The two tight dimers are respectively in red and yellow, and in pale and dark blue. The tetramerization interface involved strands β6 and β7.
(c) The AdoHcy binding site. The ligand (blue) and the enzyme residues (gray) are shown as sticks. Water molecules are drawn as blue spheres. Polar contacts are represented with dashed lines. Strictly conserved residue among m1A58 tRNA MTases are labelled with bold letters. Characteristic motifs from MTases are highlighted. (d) Density map (2Fo-Fc at 1σ) around the cofactor. The AdoHcy ligand (blue) and the enzyme residues (green) are shown as sticks. Water molecules are represented as red spheres.

Figure 2: Electrostatic surface potential of (a) *T. thermophilus* TrmI and (b) *M. tuberculosis* TrmI. Blue indicates positive charge and red negative one with the maximum color saturation corresponding to -3 kT (red) and +3 kT (blue). The figure was prepared using the APBS PyMOL plugin.

Figure 3: Noncovalent ESI-MS analyses of thTrmI and thTrmI-tRNA assemblies. ESI mass spectra obtained under non-denaturing conditions (200 mM ammonium bicarbonate buffer pH 8.0; Vc = 120 V; Pi = 6.1 mbar) for (a) thTrmI tetramer (20 µM) alone and in presence of (b) 30 µM, (c) 100 µM and (d) 200 µM of *E. coli* tRNAiMet. Inserts correspond to deconvoluted spectra showing molecular weights on the x-scale. T (MW = 115450 ± 5 Da) corresponds to (thTrmI)4, T+1tRNA (MW = 140347 ± 6 Da) corresponds to (thTrmI)4:(tRNA)1 complex and T+2tRNA (MW = 165455 ± 10 Da) corresponds to (thTrmI)4:(tRNA)2 complex. (*) correspond to tRNA signals.

Figure 4: Non-denaturing agarose gel electrophoresis
Constant amount of *E. coli* tRNAmet (4 µg) with increasing amounts of thTrmI protein (from 0 to 1 molar equivalent of thTrmI tetramer). Bands were visualized by UV shadowing.

Figure 5: Influence of the buffer ionic strength on thTrmI-tRNA complexes stabilities. ESI mass spectra obtained under non-denaturing conditions (ammonium bicarbonate pH 8.0; Vc = 120 V; Pi = 6.1 mbar) for thTrmI tetramer (20 µM) in presence of 100 µM of *E. coli* tRNAiMet. Ammonium bicarbonate concentration was set to (a) 100 mM, (b) 200 mM, (c) 400 mM and (d) 1250 mM. Inserts correspond to deconvoluted spectra showing molecular
weights on the x-scale. T corresponds to \((th\text{-}TrmI)_{4}\), T+1tRNA corresponds to \((th\text{-}TrmI)_{4}:(tRNA)_{1}\) complex and T+2tRNA corresponds to \((th\text{-}TrmI)_{4}:(tRNA)_{2}\) complex. (*) correspond to tRNA signals.

**Figure 6:** Specificity of \(th\text{-}TrmI\)-tRNA complexes formation. ESI mass spectra obtained under non-denaturing conditions (200 mM ammonium bicarbonate buffer pH 8.0; \(Vc = 120\) V; \(Pi = 6.1\) mbar) for \(th\text{-}TrmI\) tetramer (20 \(\mu\)M) in presence of (a) 100 \(\mu\)M *E. coli* tRNA\textsuperscript{Met} (b) 100 \(\mu\)M non substrate control RNA. Inserts correspond to deconvoluted spectra showing molecular weights on the x-scale. T corresponds to \((th\text{-}TrmI)_{4}\), T+1tRNA corresponds to \((th\text{-}TrmI)_{4}:(tRNA)_{1}\) complex and T+2tRNA corresponds to \((th\text{-}TrmI)_{4}:(tRNA)_{2}\) complex. (*) corresponds to RNA signals.

**Figure 7:** Sequence alignments of TrmI proteins and conserved residues in \(th\text{-}TrmI\) catalytic pocket.

(a) Part of sequence alignment of *T. thermophilus*, *M. tuberculosis*, *P. abyssi*, *M. musculus*, *H. sapiens* and *S. cerevisiae* m\(^1\)A\textsubscript{58} tRNA MTases. Residue numbers are those of *T. thermophilus* sequence. Highly conserved residues are shown on a dark blue background and residues with a similar physico-chemical character are on a pale blue background. The secondary structure of the *T. thermophilus* TrmI is shown below the sequences as arrows for \(\beta\)-strands and cylinder for \(\alpha\)-helices. The names of the secondary structure elements refer to Figure 1(a). The sequence motifs typical of the AdoMet dependent MTase family are also indicated. Residues marked with a star (*) (i.e. Y78, D170 and Y194) are involved in the catalytic pocket formation and are those studied by mutagenesis in the present work. Residues marked with a circle (o) (i.e. W175 and F245) are conserved aromatic residues in motifs IV and VIII, belonging to the catalytic pocket in m\(^6\)A DNA MTases. (b) Conserved and semi-conserved residues around the catalytic pocket of *T. thermophilus* TrmI. The strictly conserved residues among the m\(^1\)A\textsubscript{58} tRNA MTase family are shown as sticks on the *T. thermophilus* structure except the conserved residues that bind AdoMet. AdoMet was modeled from AdoHcy by adding a methyl group on the sulphur atom, the methyl is positioned by the sulfonium chirality. Y78, Y194 and D170 are indicated as red sticks.

**Figure 8:** Modeling a N\(_9\)methyladenine in the catalytic pocket of *th*TrmI

(a) Average docked structure of the N\(_9\)methyladenine ring inside the catalytic pocket of *th*TrmI. The enzyme surface is coloured according to the electrostatic surface potential. Positive potential is shown in blue and negative potential in red. The AdoMet ligand and the docked N\(_9\)methyladenine ring are shown as sticks (carbon atoms in light blue, nitrogen atoms in dark blue, oxygen atoms in red, sulphur atom in yellow and hydrogen atoms in white). The Van der Waals surface of the N\(_9\)methyladenine ring is shown as white dots.
(b) Distances between the average structure of the docked N₉-methyladenine ring and key moiety of the enzyme. The enzyme is shown in gray with the cartoon representation. The AdoMet ligand, the docked N₉-methyladenine ring and residues D170, Y78 and Y194 are shown as sticks. Distances are indicated with dashed lines and are given in Å.
Figure 3

T : 115450 ± 5 Da  T + 1 tRNA : 140347 ± 6 Da  T + 2 tRNA : 165455 ± 10 Da
* : tRNA
Figure 7
Table 1 *Data collection and refinement statistics for the determination of TrmI structure.*
Values in parenthesis are for the outer resolution shell.

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Table 2  Kinetic and thermodynamic parameters of wild-type and mutant TrmI enzymes.
The parameters are indicated with a confidence interval of 90 %.

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<th>Mutants</th>
<th>$k_{cat}$ (s$^{-1}$ x 10$^{-3}$)</th>
<th>$K_M$ (µM)</th>
<th>$k_{cat}$ (s$^{-1}$ x 10$^{-3}$)</th>
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<th>$K_D$ (nM)</th>
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<td>WT</td>
<td>37 (30 - 44)</td>
<td>2.1 (1.4-3.0)</td>
<td>36 (28 - 45)</td>
<td>2.7 (1.7-4.8)</td>
<td>13 (5 - 24)</td>
<td>100.0</td>
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<tr>
<td>Y194A</td>
<td>6 (5 – 11)</td>
<td>5.7 (3.9-8.7)</td>
<td>10 (9 - 13)</td>
<td>2.2 (1.9-3.3)</td>
<td>18 (10 - 37)</td>
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<tr>
<td>Y78A</td>
<td>1.4 (1.1 – 1.7)</td>
<td>1.9 (1.2-2.7)</td>
<td>1.6 (1.5 – 1.8)</td>
<td>0.16 (0.11-0.23)</td>
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</tr>
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<td>D170A</td>
<td>0.13 (0.11 - 0.14)</td>
<td>5.4 (3.7-8.0)</td>
<td>0.11 (0.10 - 0.12)</td>
<td>0.17 (0.07-0.39)</td>
<td>14 (6 - 25)</td>
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Figure 1 (Supplementary Material): Determination of the catalytic parameters of wild type thTrmI enzyme.

(a) Time course measurements of incorporated $^3$H-Methyl in tRNA$_i^{Met}$ for different AdoMet concentrations (0.5, 1, 2, 4, 8 µM). (b) Michaelis-Menten fit of initial rates corresponding to several AdoMet concentrations.