Synthesis of RNA-cofactor conjugates and structural exploration of RNA recognition by an m6A RNA methyltransferase

Vincent Meynier1,†, Laura Iannazzo2,†, Marjorie Catala1, Stephanie Oerum1, Emmanuelle Braud2, Colette Atdjian2, Pierre Barraud3, Matthieu Fonvielle3, Carine Tisné1,*,† and Mélanie Ethève-Quelquejeu2,*,†

1Expression Génétique Microbienne, UMR 8261, CNRS, Université Paris Cité, Institut de Biologie Physico-Chimique (IBPC), 75005, Paris, France, 2Laboratoire de Chimie et Biochimie Pharmacologiques et Toxicologiques, UMR 8601, CNRS, Université Paris Cité, 75006, Paris, France and 3Sorbonne Université, Université Paris Cité, Centre de recherche des Cordeliers, 75006, Paris, France

Received March 20, 2022; Revised April 22, 2022; Editorial Decision April 08, 2022; Accepted May 10, 2022

ABSTRACT

Chemical synthesis of RNA conjugates has opened new strategies to study enzymatic mechanisms in RNA biology. To gain insights into poorly understood RNA nucleotide methylation processes, we developed a new method to synthesize RNA-conjugates for the study of RNA recognition and methyl-transfer mechanisms of SAM-dependent m6A RNA methyltransferases. These RNA conjugates contain a SAM cofactor analogue connected at the N6-atom of an adenosine within dinucleotides, a trinucleotide or a 13mer RNA. Our chemical route is chemo- and regio-selective and allows flexible modification of the RNA length and sequence. These compounds were used in crystallization assays with RlmJ, a bacterial m6A rRNA methyltransferase. Two crystal structures of RlmJ in complex with RNA–SAM conjugates were solved and revealed the RNA-specific recognition elements used by RlmJ to clamp the RNA substrate in its active site. From these structures, a model of a trinucleotide bound in the RlmJ active site could be built and validated by methyltransferase assays on RlmJ mutants. The methyl transfer by RlmJ could also be deduced. This study therefore shows that RNA-cofactor conjugates are potent molecular tools to explore the active site of RNA modification enzymes.

INTRODUCTION

RNAs are heavily co- or post-transcriptionally modified. To date, 152 chemical modifications of bases and ribose have been described, with the largest diversity and highest number of modifications found in transfer RNA (tRNA) (1–4). The mono-methylation of the exocyclic amine of adenine (m6A) is found in all organisms from bacteria to human and in mRNA, tRNA, rRNA, small nucleolar RNA, long non-coding RNA and microRNA. It is the most commonly occurring internal modification in mRNA from eukaryotes. m6A is reversible on mRNA, meaning that distinct sets of proteins introduce (m6A RNA methyltransferases called the writers), recognize (the readers) and remove (the erasers) this epitranscriptomic mark, thus allowing additional levels of regulation of mRNA translation. It has yet to be conclusively determined if such a writer/reader/eraser system exists for m6A on other types of RNA. The regulatory mechanism of the m6A modification is complex (5,6) and its dysregulation is linked to human diseases (7–9) and virus replication (10–12).

m6A RNA methyltransferases (MTases) catalyze the transfer of the methyl group from the cofactor S-adenosyl-L-methionine (SAM, Figure 1A) to the N6-atom of adenines at specific positions in RNA, releasing the cofactor product S-adenosyl-L-homocysteine (SAH, Figure 1A). Six m6A RNA MTases have been structurally characterized so far (3,4), but only one enzyme, METTL16, has been crystallized in complex with a substrate RNA (PDB: 6DU4 and 6DU5) (13). Although the methyltransferase domains of the six MTases superimpose well, the structure of the METTL16/RNA complex does not allow an understanding of RNA recognition by the other m6A RNA MTases.

†The authors wish it to be known that, in their opinion, the first two authors should be regarded as Joint First Authors and the last two authors should be regarded as Joint Last Authors.

*To whom correspondence should be addressed. Tel: +33 1 58 41 50 13; Email: carine.tisne@cnrs.fr
Correspondence may also be addressed to Mélanie Ethève-Quelquejeu. Email: melanie.etheve-quelquejeu@u-paris.fr

© The Author(s) 2022. Published by Oxford University Press on behalf of Nucleic Acids Research.
This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.
Figure 1. (A) RlmJ-catalyzed transfer of the methyl group of SAM to N6-atom of A2030 of the E. coli 23S rRNA. The minimal RNA substrate for RlmJ is drawn in purple. (B) Bisubstrate analogue (A*) previously described (14,15). The inset shows the abbreviations used to differentiate the two adenosines contained in the adenosine-SAM conjugates. (C) SAM cofactor analogue (CA).

In fact, superimposition of RNA-bound METTL16 with other structurally characterized m6A MTases reveals steric clashes between the protein and the RNA for all enzymes, indicating that their specific RNA-substrate recognition differs from that of METTL16. The limited number of RNA-bound m6A MTase structures reflects the difficulty to solve the structure of modification enzymes in complex with RNA. Consequently, the RNA recognition and methylation reaction mechanism of m6A RNA MTases are still poorly understood. In contrast, structures are available for most MTases in complex with either the SAM cofactor or the cofactor product, SAH, allowing for a well-understood cofactor binding pattern within this family of proteins. The cofactor binding guided the initial design of bisubstrate analogues as chemical tools to mimic the state at which both the substrate nucleoside and the SAM cofactor are bound in the catalytic pocket of the enzyme (14–17). In recent years, there have been significant efforts made toward the development of SAM analogues with the aim to study the transfer of moieties other than simple methyl groups (18,19), to obtain protein inhibition (20) or to study SAM-dependent DNA-methyltransferase mechanisms (21,22). Here, the aim is to facilitate protein/RNA crystallization to gain insights into currently poorly understood RNA nucleotide methylation processes.

We previously designed adenosine-SAM bisubstrate analogues that contain the SAM cofactor with the S-atom replaced by a N-atom directly linked to an adenosine at the point of nucleophilic attack to mimic the substrate/cofactor bound state (14–16) (Figure 1B). These molecules were used to study the substrate recognition of the m6A MTase RlmJ, the Ribosomal RNA large subunit methyltransferase J that N6-methylates the adenine A2030 of the E. coli 23S rRNA (23) (Figure 1A). RlmJ is a good example of an m6A MTase that was successfully crystallized with SAM (PDB 4BLV), SAH (PDB 6QE5) and SAM plus AMP as the adenine substrate analogue (PDB 4BLW), but not with an RNA substrate (23,24). In the AMP-bound RlmJ structure, the exocyclic N6-atom of the AMP-adenine was outside hydrogen-bonding distance from potential proton acceptors and was unlikely to mimic the true adenosine substrate (Asub) position. In contrast, we previously showed that the bisubstrate analogue A* (Figure 1B), containing a SAM cofactor analogue CA (Figure 1C) tethered to Asub at the N6-atom by an alkyl linker of three carbons, crystallized in the active site of RlmJ (15). The adenine is positioned in the presumed substrate binding pocket of RlmJ. For the cofactor moiety, the methionine chain is bound like SAM but the adenosine of the SAM analogue (ASA) is rotated out of the canonical binding pocket for SAM. In the present study, we show that CA is able to bind correctly in the active site of RlmJ. CA is therefore used as a universal building block and we develop a new chemical strategy to increase the size of the RNA tethered to CA (Figure 2).

These RNA-cofactor conjugates could be synthetized using two different strategies: (i) a post-synthetic approach, which involves the introduction of a reactive group into the RNA followed by a reaction with the cofactor in a selective manner (Scheme 1, route A) or (ii) a chemo-enzymatic approach called SMILing for ‘Sequence-specific Methyltransferase-Induced Labeling’ (Scheme 1, route B) based on alkylation reactions catalysed by a MTase in presence of cofactor analogues (25,26). This SMILing method has been previously described for modification of proteins (27) and DNA (26,28). In this approach, the nucleophilic attack of the substrate adenosine is expected to open the putative aziridine ring of the SAM analogue and directly provide an adenosine-SAM conjugate tethered by a two-carbon linker. The synthesis of RNA–SAM conjugates with three-carbon linker therefore requires the development of a
Figure 2. RNA–SAM conjugates synthesized in this work to explore the RNA recognition by RlmJ.

Scheme 1. Routes for the synthesis of RNA cofactor conjugates.

post-synthetic approach (Scheme 1, route A) with a chemoselective post-functionalization step to connect the SAM cofactor analogue to the N6-atom of a specific adenosine in the RNA.

Here, we show that nucleophilic aromatic substitution as a post-synthetic reaction can be efficiently used to synthesize four RNA–SAM conjugates, in which the CA SAM analogue is connected to the N6-atom of an adenosine. This modified adenosine is located at the 5′-end or 3′-end of a dinucleotide, in the internal position of a trinucleotide or within a 13-mer RNA with the sequence of the RNA substrate of RlmJ (Figure 2). These compounds were then used in crystallization assays in complex with RlmJ. Two crystal structures were solved with an RNA–SAM conjugate bound in the active site of RlmJ. These structures reveal the specific recognition mode of RNA by RlmJ, which was further confirmed by mutation and MTase activity studies and decipher the mechanism of methyl transfer used by RlmJ.

**MATERIALS AND METHODS**

**General information**

Reactions were carried out under argon atmosphere and performed using freshly distilled solvents. Dimethyl formamide (DMF) and MeOH were dried over calcium hydride. Phosphoramidite nucleosides, tetrazole and solvents for dinucleotides synthesis were purchased from Eurogentec. Progress of the reactions was monitored by thin layer chromatography (TLC). TLC: precoated silica gel thin layer sheets 60 F 254 (Merck, Darmstadt, Germany) and detection by charring with 10% H2SO4 in ethanol followed by heating.

Preparative HPLC were performed using a Reverse-phase HPLC system (Shimadzu, Marne-la-Vallée, France) with a reverse phase C-18 NUCLEOSIL column (250 mm × 21.2 mm, 5 μm) using a solvent system consisting of A: 50 mM aqueous NH4OAc pH 4.5 and B: MeCN (linear gradient from 0% B to 63% B in 30 min) at a flow rate of 15 ml/min and UV detection at 254 nm.

The NMR characterizations for all synthesized compounds are presented in the Supplementary data and the NMR spectra are provided in Supplementary Figures S1–S29. The reverse-phase HPLC chromatograms of the final compounds (A*A, GA*A, GA* and the 13mer RNA–SAM conjugate) are shown in Supplementary Figures S30–S36.

**Chemical synthesis of compound 7 (Scheme 2)**

Inosine (2.0 g, 7.45 mmol) was dissolved in DMF (20 ml) and di-tert-butylsilyl bis(trifluoromethanesulfonate) (2.67 ml, 8.19 mmol) was added dropwise at 0°C. The reaction was stirred at 0°C for 30 min and imidazole (2.53 g, 37.25 mmol) was added. The reaction was allowed to warm to room temperature (rt) and stirred for an additional hour. tert-butyl chloride (2.24 g, 14.9 mmol) and imidazole (1.01 g, 14.9 mmol) were added successively at 0°C and the reaction was stirred at room temperature overnight. The re-
action was diluted with ethyl acetate (EtOAc) and washed five times with brine. The combined organic layers were dried over anhydrous MgSO₄, filtered, and concentrated. The residue was purified by silica gel chromatography (eluent: cyclohexane/EtOAc 5:5) to provide the compound 1 as a white foam (3.9 g, quantitative yield). To a solution of 1 (3.45 g, 6.6 mmol) in THF (60 ml), hydrogen fluoride/pyridine complex (1.48 ml, 16.5 mmol) in pyridine (2.5 ml) was added at 0 °C and the mixture was stirred at room temperature for 15 min. The reaction was diluted with Dichloromethane (DCM) and extracted with saturated solution of NaHCO₃ and brine. The organic layer was dried over anhydrous MgSO₄, filtered, and concentrated under reduced pressure. The residue was purified by silica gel chromatography using DCM as eluent to afford compound 1 as a white foam (1.74 g, 69%). Compound 2 (1.74 g, 4.5 mmol) was dissolved in pyridine (7 ml). 4,4′-Dimethoxytrityl chloride (1.69 g, 4.9 mmol) was added at 0 °C and the reaction was stirred at 0 °C for 16 h. The reaction was diluted with DCM, washed with brine, dried over anhydrous MgSO₄, filtered, and concentrated. The residue was purified by silica gel chromatography (eluent: DCM/MeOH 9/1) as the eluent to afford compound 2 as a white foam (1.74 g, 69%). Compound 2 (1.74 g, 4.5 mmol) was dissolved in pyridine (7 ml). 4,4′-Dimethoxytrityl chloride (1.69 g, 4.9 mmol) was added at 0 °C and the reaction was stirred at 0 °C for 16 h. The reaction was diluted with DCM, washed with brine, dried over anhydrous MgSO₄, filtered, and concentrated. The residue was purified by silica gel chromatography (eluent: DCM/MeOH 96:4) to provide the desired compound 3 as a white foam (1.01 g, 32%). Benzotriazol-1-yloxytripyrrolidinophosphonium hexafluorophosphate (PyBOP) (689 mg, 1.55 mmol) and disopropylamine (DIPEA) (339 μl, 3.19 mmol) were added to a solution of compound 2 (1.74 g, 4.5 mmol) in MeOH (98:2) as eluent to afford compound 3 as a white foam (0.54 g, 31% over three steps).

**Chemical synthesis of compound 10 (Scheme 3)**

Compound 7 (183 mg, 140 μmol) and Ac-G-PCNE phosphoramidite (250 mg, 266 μmol) were stirred at room temperature for 30 min in MeCN (2 ml). A 0.45 M tetrabenzyl-adenosine solution in MeCN (2.2 ml, 1 mmol) was added. After stirring at room temperature for 20 h, a 0.1 M iodine solution in THF/H₂O/Pyridine (75/20, 3.1 ml) was added. After 1 h, the reaction mixture was diluted with EtOAc, washed with water, a saturated solution of Na₂S₂O₃ and brine. The organic layers were combined, dried over anhydrous MgSO₄, filtered, and evaporated. The residue was then purified by silica gel chromatography using DCM/MeOH (98/2) as eluent to afford compound 7 as a white foam (50 mg, 38% over three steps).

**Chemical synthesis of compound 15 (Scheme 4)**

To a solution of inosine (1.0 g, 3.7 mmol, 1 eq) and imidazole (2.03 g, 29.8 mmol, 8 eq) in DMF (11 ml) was added tetrabenzoyl-adenosine 6 (70 mg, 100 μmol). The reaction mixture was stirred at room temperature for 30 min and a 0.45 M tetrabenzoyl-adenosine solution in MeCN (2.2 ml, 1 mmol) was added. After stirring at room temperature for 20 h, a 0.1 M iodine solution in THF/H₂O/Pyridine (75/20, 3.1 ml) was added. After 1 h, the reaction mixture was diluted with EtOAc, washed with water, a saturated solution of Na₂S₂O₃ and brine. The organic layers were combined, dried over anhydrous MgSO₄, filtered, and evaporated. The residue was then dried at 0.18 M trichloroacetic acid (TCA) solution in DCM (5.7 ml) at room temperature for 30 min. The reaction mixture was diluted with DCM and the organic layer washed with a saturated solution of NaHCO₃ and brine, dried over anhydrous MgSO₄, filtered, and evaporated. The residue was purified by silica gel chromatography using DCM/MeOH (98/2) as eluent to afford compound 7 as a white foam (129 mg, 49% over three steps).
Scheme 3. Synthesis of A* A and GA* A RNA–SAM conjugates. (i) Ac-G-CE-phosphoramidite, tetrazole, MeCN, rt, 20 h, (ii) I2, THF/Pyr/H2O, rt, 1 h, (iii) TCA, DCM, rt, 30 min, (49% over three steps); (iv) DIPEA 12 eq, DMF, rt, 24 h, (v) MeNH2, EtOH/H2O, rt, 24 h, (vi) CsF, MeOH, 60°C, 24 h; (vii) ZnBr2, MeNO2/iPrOH, rt, 24 h.

Scheme 4. Synthesis of the GA* RNA-conjugate. (i) TBDMSI, imidazole, DMF, 50°C, 18 h, 87%, (ii) PyBOP, DIPEA, DMF, rt, 16 h, 80%, (iii) TFA/H2O, 60°C, 1 h, 83%, (iv) I2, THF/Pyr/H2O, rt, 1 h, (v) TCA, DCM, rt, 30 min (56% over three steps), (vi) DIPEA 12 eq, DMF, rt, 24 h, (vii) MeNH2, EtOH/H2O, rt, 24 h, (viii) CsF, MeOH, 60°C, 24 h (4% over three steps), (ix) ZnBr2, MeNO2/iPrOH, rt, 24 h, 30%.

added tert-butyldimethylsilyl chloride (TBDMSI) (2.25 g, 14.9 mmol, 4 eq) at 0°C. The reaction mixture was then stirred at 50°C for 16 h. The residue was partitioned between EtOAc and brine. The organic phase was washed with brine, dried over anhydrous MgSO4, filtered, and concentrated. The residue was purified by silica gel chromatography (cyclohexane/EtOAc 5:5) to provide the compound 12 as a white foam (1.96 g, 87%). PyBOP (618 mg, 1.40 mmol) and DIPEA (304 μl, 1.74 mmol) were added at 0°C to a solution of inosine 12 (711 mg, 1.16 mmol) in DMF (5 ml) and the reaction mixture was stirred at room temperature for 16 h. The residue was partitioned between EtOAc and brine. The organic layer was dried over anhydrous MgSO4, filtered, and concentrated. The residue was purified by silica gel chromatography (cyclohexane/EtOAc 9:1) to provide the compound 13 as a white foam (677 mg, 80%). A 1/1 (v/v) mixture of TFA/H2O (3.5 ml, 46 mmol) was added at 0°C to a solution of compound 13 (677 mg, 0.92 mmol) in THF (20 ml). The reaction mixture was stirred at room temperature for 1 h. EtOAc and a saturated solution of
NaHCO₃ were added and the organic phase was washed with brine. The organic layer was dried over anhydrous MgSO₄, filtered and concentrated. The residue was purified by silica gel chromatography using cyclohexane/ EtOAc (7/3) as eluent to afford compound 14 (361 mg, 63%) as a white foam. Compound 14 (85 mg, 138 µmol) in MeCN (2 ml) was added to a solution of Ac-G-PCNE phosphoramidite (250 mg, 262 µmol) in MeCN (300 µl). The reaction mixture was stirred at room temperature for 30 min and a 0.45 M tetrazole solution in MeCN (3 ml, 1.38 mmol) was added. After stirring at room temperature for 20 h, a 0.1 M iodine solution in THF/H₂O/Pyridine (75/2/20, 4.1 ml) was added. After 1 h, the reaction mixture was diluted with EtOAc, washed with water, a saturated solution of Na₂S₂O₃ and brine, dried over anhydrous MgSO₄, filtered, and evaporated. The residue was purified by silica gel chromatography using DCM/MeOH (96/4) as eluent to afford compound 15 (85 mg, 63% over three steps).

Chemical synthesis of compound 18 (Scheme 5)

To a solution of iPr-Pac-dG-PCNE phosphoramidite (500 mg, 0.53 µmol) in MeCN (600 µl) was added compound 14 (111 mg, 0.18 mmol) in MeCN (4 ml). The reaction mixture was stirred at room temperature for 30 min and a 0.45 M tetrazole solution in MeCN (4 ml, 1.8 mmol) was added. After stirring at room temperature for 20 h, a 0.1 M iodine solution in THF/H₂O/Pyridine (75/2/20, 5.4 ml) was added. After 1 h, the reaction mixture was diluted with EtOAc, washed with water, a saturated solution of Na₂S₂O₃ and brine, dried over anhydrous MgSO₄, filtered, and concentrated under vacuo. The residue was then stirred with a 0.18 M TCA solution in DCM (7.6 ml) at room temperature for 30 min. The reaction mixture was diluted with DCM and the organic layer was washed with a saturated solution of NaHCO₃ and brine, dried over anhydrous MgSO₄, filtered, and evaporated. The residue was purified by silica gel chromatography using DCM/MeOH (96/4) as eluent to afford compound 17 (147 mg, 69% over three steps). Compound 17 (147 mg, 0.12 mmol) was dissolved in a 5 M solution of Na₂S₂O₃ and brine. The organic layers were concentrated, the residue was dissolved in a 5 M solution of NaHCO₃ and brine, dried over anhydrous MgSO₄, filtered, and evaporated. The residue was purified by silica gel chromatography using DCM/MeOH (96/4) as eluent to afford compound 18 (134 mg, 85% over 2 steps).

Chemical synthesis of A*A, GA*A and GA* (Schemes 3, 4 and 5)

SAM analogue 8 (160 mg, 0.18 mmol) and DIPEA (355 µl, 2.04 mmol) were added at 0°C to a solution of compound 7, 10, 15 or 18 (1 equivalent) in DMF and the reaction mixture was stirred at room temperature for 24 h. After concentration, the residue was dissolved in a 5 M solution of MeNH₂ (EtOH/H₂O, 1/1) (200 equivalent) and the reaction mixture was stirred at room temperature for 24 h and concentrated. The residue was then dissolved in MeOH and CsF (200 equivalent) added. The reaction mixture was stirred at 60°C for 24 h. After concentration under vacuo, the residue was purified using a Reverse-phase HPLC system. The appropriate fractions were collected and lyophilized, to give protected intermediates 9, 11, 16 or 19, respectively. Compound 9, 11, 16 or 19 (1 equivalent) was dissolved in a 5 M solution of ZnBr₂ in a 1/1 (v/v) mixture of iPrOH/MeNO₂ (400 equivalent) and the reaction mixture was stirred at room temperature for 24 h. Water was added, and the mixture was lyophilized. The residue was then purified using a Reverse-phase HPLC system. The appropriate fractions were collected and lyophilized to give compound A*A, GA*A or GA*.

Synthesis of the 13mer RNA–SAM conjugate (Scheme 5)

SAM analogue 8 (160 mg, 0.18 mmol) and DIPEA (355 µl, 2.04 mmol) were added at 0°C to a solution of compound 18 (231 mg, 0.17 mmol) in DMF (3 ml) and the reaction mixture was stirred at 60°C for 24 h. After concentration under vacuo, the residue was purified using a Reverse-phase HPLC system. The appropriate fractions were collected and lyophilized, to give the protected dinucleotide 19 (14 mg, 69% over three steps). Dinucleotide 19 (14 mg, 11 µmol) was dissolved in a 5 M solution of ZnBr₂ in a 1/1 (v/v) mixture of iPrOH/MeNO₂ (400 equivalent) and the reaction mixture was stirred at room temperature for 24 h and concentrated. The residue was then dissolved in MeOH (15 ml) and CsF (5.16 g, 34 mmol) was added. The reaction mixture was stirred at 60°C for 24 h. After concentration under vacuo, the residue was purified using a Reverse-phase HPLC system. The appropriate fractions were collected and lyophilized to give the dinucleotide pdGA* (0.4 mg, 4%).

Synthesis of 11mer RNA (5'-GAACUCGCUGU-3').

The 11mer RNA sequence (5'-GAACUCGCUGU-3') was synthesized automatically (1 µmol scale) with the use of an H-2 GeneWorld DNA/RNA automated synthesizer (K&A, Laborgeraete GbR, Schafheim, Germany). More information is given in the Supplementary data.

Enzymatic ligation. The ligation of pdGA* (200 nmol) to the 11-mer RNA (20 nmol) was performed with purified T4 RNA ligase, in the presence of DMSO (10%), ATP (1 mM) and MgCl₂ (15 mM) in 500 µl of 50 mM HEPES buffer pH 7.5 at 30°C for 12 h (Supplementary data, page 23). The 13mer RNA–SAM conjugate was characterized by mass spectrometry and analyzed by reverse-phase HPLC.
Scheme 5. Synthesis of a 13mer RNA–SAM conjugate by enzymatic ligation. (i) iPr-Pac-dG-PCNE phosphoramidite, tetrazole, MeCN, rt, 20 h, (ii) I₂, THF/Pyr/H₂O, rt, 1 h, (iii) TCA, DCM, rt, 30 min (69% over three steps), (iv) (iPr)₂N(OC₂H₅)₂, Tetrazole, MeCN, rt, 3 h, (v) I₂, THF/Pyr/H₂O, rt, 30 min; (85% over two steps), (vi) DIPEA, DMF, rt, 24 h, (vii) MeNH₂, EtOH/H₂O, rt, 24 h, (viii) CsF, MeOH, 60°C, 24 h (7% over three steps), (ix) ZnBr₂, MeNO₂/iPrOH, 4%. 11-mer RNA (GAACUCGCUGU, the sequence mimicking the substrate RNA of RlmJ, was synthesized by SPS).

Cloning, expression and purification of RlmJ and mutants
RlmJ (residues 1–280) was subcloned into the pET15b vector bearing an N-terminal His₆-tag followed by a thrombin protease cleavage site as previously described (15). Mutations were generated using the QuikChange Site-Directed Mutagenesis Kit (Agilent), and verified by sequencing. Plasmids were transformed into E. coli BL21(DE3) competent cells, cultured in Lysogeny Broth at 37°C, and protein expression was induced with 0.5 mM isopropyl-1-thiogalactopyranoside for 3 h at 37°C. Cell pellets were lysed by sonication in a buffer containing 20 mM Tris–HCl pH 7.0, 200 mM NaCl, 5% (v/v) glycerol and 5 mM β-mercaptoethanol, clarified by centrifugation. The clarified lysate was applied to a 5 ml HisTrap HP column (Cytiva) and eluted with a gradient of 20–500 mM imidazole. RlmJ was treated with thrombin protease for 4 h at 22°C to remove the His₆-tag and applied to a 5 ml HisTrap HP column (Cytiva) to trap the His₆-tag. Further purification of RlmJ was performed by size exclusion chromatography (SEC) on Superdex S75 PG (Cytiva). RlmJ was concentrated and stored at −80°C in the SEC buffer (20 mM Tris–HCl pH 7.0, 150 mM NaCl, 5% (v/v) glycerol and 5 mM β-mercaptoethanol).

Crystallization, data collection and structure determination
Crystallization assays were conducted at 18°C using the sitting-drop vapor-diffusion method and the Hampton Research Index HT™ screen, for which crystals of RlmJ in complex with SAM or SAH are routinely produced. RNA–SAM conjugates were dissolved in H₂O at 10 mM final concentration, except for the 13mer RNA–SAM that was dissolved at 1 mM final concentration. RlmJ and RNA–SAM conjugates were mixed at ratios ranging from 1/2 to 1/10 before crystallization assays with RlmJ concentrated at around 10 mg/ml. For CA, A*A and GA*, all the conditions of the screen were performed at different RlmJ/RNA–SAM ratios. For GA*, only five lines (D to H) of the Hampton Research Index HT™ screen were tested with a RlmJ/GA* ratio of 1/2, because the amount of GA* was limiting (217 nmol). Three conditions, where crystals grew without ligand bound to RlmJ when using the 1/2 ratio, were performed again by increasing the RlmJ/GA* ratio to 1/8. For the 13mer RNA–SAM (3.7 nmol available for crystallogenesis assays), only two lines (D and H) at a RlmJ/13mer RNA–SAM ratio of 1/2 could be tested with a final protein concentration of 5.3 mg/ml.

RlmJ bound to CA (RlmJ/CA ratio of 1/10) crystallized with a final protein concentration in the drop of 4.75 mg/ml, in a 150 μL drop with a 40 μL reservoir solution containing 0.1 M Bis–Tris pH 6.5, 28% (w/v) PEG MME 2000. RlmJ bound to GA* (RlmJ/GA* ratio of 1/2) crystallized with a final protein concentration of 5.5 mg/ml, in a 400 μL drop with a 40 μL reservoir solution containing 0.1 M Bis–Tris pH 6.5, 0.2 M NaCl, 25% (w/v) PEG 3350. RlmJ bound to GA*A (RlmJ/GA*A ratio of 1/8) crystallized with a final protein concentration of 5.0 mg/ml, in a 200 μL drop (100 + 100) with a 20 μL reservoir solution containing 0.1 M Bis–Tris pH 6.5, 20% (w/v) PEG MME 5000.

Crystals were cryoprotected with reservoir solution supplemented with 19, 23 or 21% (v/v) glycerol for crystals with CA, GA* or GA*A respectively, and flash frozen in liquid nitrogen. Diffraction data were collected at the Synchrotron Soleil beamline PX1 or PX2 and refined to 2.09 Å for RlmJ/CA, 2.29 Å for RlmJ/GA* and 1.59 Å for RlmJ/GA*A. Diffraction data were processed using XDS.
(29). Phases were determined by molecular replacement in
PHASER (30) in the CCP4 Suite of programs (31), using
an existing RlmJ structure as search model (PDB 6QDX).
Modelling and refinement were carried out using Refmac
(32), COOT (33) and Phenix (34). Density was visible for all
residues aside from loop-y parts covering residues 53–57 in
RlmJ/CA, residues 54–58 of RlmJ/GA* and residues 52–57
in chain A and residues 52–56 in chain B of RlmJ/GA*A.

Differential scanning fluorimetry (DSF)

DSF was performed in a 96-well plate using a CFX96 Touch
real-time PCR detection system (Bio-Rad) with excitation
and emission filters of 450–490 and 515–530 nm, respec-
tively. Each 20 μl well reaction was carried out in the DSF
buffer (20 mM Tris-HCl pH 8.0, 150 mM NaCl, 5% (v/v)
glycerol and 5 mM β-mercaptoethanol) and consisted of 2
μl protein at a final concentration of 5 μM, 2 μl SYPRO
ORANGE diluted 500-fold from the manufacturer’s stock (i.e.
5000× in DMSO (Invitrogen)) in the SEC buffer and (if
applicable) 2 μl S-adenosyl-L-methionine (SAM) (Merck)
to a final concentration of 1 mM. Fluorescence intensi-
ties were measured from 25 to 85°C with a ramp rate of
1°C/min. The melting temperature T_m was determined by
curve-fitting using GraphPad Prism v.7.0 software (35).

Methyltransferase activity assays

(GG)H72 RNA (27 nucleotides, sequence 5′-GGGAACUCGCUGAGAAGGAGUGUGUA-3′)
was purchased deprotected and desalted from Dharmacon
(HorizonTM). It was resuspended in the activity buffer (20
mM potassium phosphate pH 6.5, 50 mM NaCl, 5 mM
β-mercaptoethanol) to a 500 μM concentration, heated for
2 min at 95°C then cooled down on ice for at least 15 min.
A reaction contained 2000 pmol (GG)H72 RNA supple-
mented with 3000 pmol unlabeled SAM (Sigma-Aldrich)
doped with 5 pmol S-adenosyl-L-[methyl-3H]-methionine
(80 Ci/mmol, American Radiolabeled Chemicals, Inc.)
and followed by the addition of 200 pmol wild-type or mutant
RlmJ or buffer alone. All reactions were carried out at
37°C for 1 h in a 50 μl final volume of the activity buffer
previously described, then quenched in 1.5 ml of ice cold
15% (w/v) trichloroacetic acid and incubated on ice for at
least 10 min. The precipitates were collected by filtration
using GF filters (Whatman) under vacuum and washed
four times with 5 ml of cold 15% (w/v) trichloroacetic acid.
The washed and dried filters were then placed
in vials containing 5 ml of liquid scintillation cocktail
(PerkinElmer), shaken for 15 min and counted in an Hidex
300 SL scintillation counter (LabLogic ScienceTec). All
assays were done within 1 day of enzyme purifications, as
the enzyme is unstable when stored over time.

RESULTS AND DISCUSSION

Chemical synthesis of RNA–SAM conjugates

The synthesis of RNA–SAM conjugates was achieved by a
chemo-selective nucleophilic aromatic substitution (SnAr)
step to connect CA via a 3-carbon alkyl chain to the N6-
atom of a specific adenosine of the RNA. Here, the convert-
ible nucleoside was introduced at the 5′-end or in an internal
position of the RNA using the O6-(benzotriazolyl)inosine
phosphoramide 5 and at the 3′-end of the RNA using the
O6-(benzotriazolyl)inosine 14 by phosphoramidite chem-
istry.

Synthesis of the di- and tri- nucleotides–SAM conjugates

The sequences AA and GAA (Figure 1A) corresponding
to the RlmJ RNA-substrate sequence were chosen to
synthesize di and tri- nucleotides–SAM conjugates. We
started our synthetic work by the synthesis of compound
7 (Scheme 2). First, the inosine was silylated in 2′-, 3′-
and 5′-positions to afford compound 1. A selective depro-
tection followed by the tritylation of the 5′-OH and ac-
tivation of the O6-position using the BOP reagent (37)
gave compound 4. Then, the phosphitylation step pro-
duced compound 5 in 73% yield. Compound 5 in presence
of the tetra-benzoylated adenosine 6 (36) provide the O6-
(benzotriazolyl)-dinucleotide 7 in 38% yield over three steps
(Scheme 2).

The key step of SnAr reaction studied in this work was
then optimized and efficient deprotection conditions were
set up. The O6-(benzotriazolyl)-dinucleotide 7 was submit-
ted to a SnAr reaction with the compound 8 in the pres-
ence of diisopropylethylamine as a base (Scheme 3). During
this step, the amine of the linker plays the role of the nucle-
ophile in the SnAr reaction but, also, in the deprotection
step of the nucleic base. Addition of methylamine is nec-
essary to complete the deprotection of the phosphotriester
function and the nucleic bases. Cesium fluoride was used to
remove the silyl ether groups and the removal of the Boc
and the tBu protecting groups was performed using ZnBr2
salts in a mixture of isopropanol and nitromethane (37,38).
The corresponding dinucleotide-SAM conjugate, A*A
was obtained after reverse-phase HPLC purification and fully
characterized by 1H,31P NMR (Supplementary Figures S8,
S9) and mass spectrometry. A mass of 4.5 mg of A*A was
obtained with an overall yield of 7% over four steps.

The synthesis of the trinucleotide GA*A started from
dinucleotide 7 which was submitted to a phosphoramide
coupling in presence of the commercially available Ac-G-
CE phosphoramidite leading to the formation of the O6-
(benzotriazolyl)-trinucleotide 10 in 49% yield. The conju-
gate GA*A was then obtained after the addition of compound
8 followed by the three deprotection steps showing the
possibility to functionalize an RNA at an internal posi-
tion in the RNA sequence with the A*SAM linked to a nucleo-
side. The yields of the two last steps could not be determined
since a pure GA*A compound has never been obtained.
However, the synthesized quantity (0.7 mg) of GA*A and
the purity were sufficient to obtain crystals and to solve
the crystal structure of GA*A in complex with RlmJ.

To introduce the SAM-cofactor at the 3′-end of the RNA,
we first prepared the convertible nucleoside 14. Inosine was
silylated in 2′-, 3′- and 5′-positions to afford compound
12, which was then activated using the PyBOP reagent to give
compound 13. The selective removal of the TBS group at the
5′-position was achieved in the presence of aqueous TFA to
give compound 14 in an overall yield of 43% (Scheme 4).
The O6-(benzotriazolyl)-dinucleotide 15 was then obtained
in 56% yield using the strategy described for compound 7.
Use of RNA–SAM conjugates to solve crystal structures with RlmJ

Crystallization assays of RlmJ were performed in the presence of A*Ag, GA*, GA*A or the 13mer RNA–SAM conjugates in various RlmJ/compound ratios. Crystals were obtained for GA* and GA*A bound to RlmJ at a RlmJ/GA* ratio of 1/2 and RlmJ/GA*A ratio of 1/8. The GA*A conjugate density was not well-defined with RlmJ/GA*A ratios < 1/8. The structures of RlmJ bound with GA* or GA*A were refined to 2.3 Å and 1.6 Å, respectively (Supplementary Table S1). Both structures displayed well-defined ligand density in the active site, which facilitated fitting of both conjugates (Supplementary Figure S37). RlmJ displays a class I MTase Rossmann-fold with a helical subdomain inserted at residues 47–98, similar to what was previously described (15,24).

In the RlmJ/GA* structure, the A^SAM moiety is superimposable with the RlmJ-bound conformation of SAH, A_sub (corresponding to A2030 in the E. coli 23S rRNA) is bound in the presumed substrate binding site, and the neighbouring guanosine (corresponding to G2029 in the E. coli 23S rRNA) binds in a pocket on the surface of RlmJ next to the catalytic site (Figure 4A). The N6-atom of A_sub is positioned, through the alkyl chain of the linker, at 3.0 Å away from the carbon corresponding to the Ce-atom of the methionine moiety in the SAM cofactor. Such a distance, in two non-linked moieties, would allow for an Sn2 methyl transfer to the N6-atom. A_sub of GA* is involved in an intricate network of interactions with strictly conserved residues of RlmJ (Supplementary Figure S38) that reveals the RlmJ active site. A_sub is indeed π-stacked between the aromatic residues, H9 and W195 (Figure 4B), and is stabilised by hydrogen bonds from its N1- and N3-atoms to the side chains of the K18 and N12 residues, respectively (Figure 4C). The classical m^6A MTase catalytic 164DPPY^167 motif (D/N-PP-Y/F/W) also participates to the binding of GA* (Figure 4D). D164 forms a hydrogen bond of 2.7 Å to A^SAM and we propose that a hydrogen bond between D164 and the N6-atom of A2030 would form in the case of the non-alkylated RNA substrate (Figure 4F). This hydrogen bond cannot be formed in this structure due to the N6-alkylation of A_sub in GA*. Mutations of K18 and D164 to alanine were previously shown to abolish RlmJ activity, confirming their involvement in RNA binding and catalysis. P165 in the 164DPPY^167 motif, through its carbonyl group, is also likely to form a second hydrogen bond to the N6-atom of A_sub (Figure 4D). In addition, P166 makes hydrophobic contacts with A^SAM (Figure 4D) and Y167 is engaged in hydrophobic interactions with the guanine ring of GA* (Figure 4E). Y167 is not stacked with A_sub in contrast to what was previously observed for F187 of the catalytic NPPF motif.
Figure 4. Crystal structure of RlmJ/GA* complex. (A) GA* on the surface of RlmJ. CA in the RlmJ/CA complex is represented as yellow sticks. (B) Stacking of the A$^{\text{sub}}$ between the strictly conserved aromatic residues W195 and H9 of RlmJ. (C) A$^{\text{sub}}$ is involved in hydrogen bonds with the strictly conserved residues N12 and K18 of RlmJ. (D) Interactions between the catalytic DPPY$_{167}$ motif of RlmJ and A$^{\text{sub}}$. (E) Interactions of the guanosine (G) of GA* with RlmJ. (F) Outline of RlmJ/GA* interactions from a LigPlot analysis (40), dashed lines in black indicate residues in RlmJ that form hydrogen bonds with GA* in the crystal structure, dashed lines in green indicate residues in RlmJ that could form hydrogen bonds with the ligand if the N6-nitrogen is not alkylated (which is the case in the natural substrate), small dark blue lightnings highlight residues in RlmJ involved in strong hydrophobic contacts with GA* and residues involved in formation of the binding pocket environment are shown in grey.

The N-terminal tail of RlmJ is flexible and includes the strictly conserved aromatic residues Y4, H6, F8 and H9. F8$^{\text{F8/H9}}$-stacks with A$_{3'}$ in the GA$^*$-bound RlmJ structure (Figure 5B) whereas H9$^{\text{H9/H9}}$-stacks with Asub in the GA*-bound RlmJ structure (Figure 4B). Interestingly, H9 swaps conformation from solvent exposed in the RlmJ/CA structure, to buried in the RlmJ/GA* structure (Figure 5C), from where it can $\pi$-stack with A$^{\text{sub}}$. The positions of Y4 and H6 across RlmJ structures presented in this paper (Figure 5C) suggest an involvement in RNA nucleotide stacking, probably in the loop of the RNA substrate. Indeed, mutation of Y4 for an alanine or H6 for an aspartate abolished MTase activity (24). The N-terminal tail of RlmJ displays vastly different conformations in various crystal structures. In the apo form (PDB 4BLU), in the structures with SAM-adenosine bisubstrates (PDB: 6QDX and 6QE0) and in the three structures presented here, the tail is located on the surface of the protein and is devoid of any secondary structure. In structures of RlmJ bound to SAH (PDB: 6QE5) or SAH and AMP (PDB: 4BLW), a short 3$_{10}$ structural motif of METTL16 (13). The same ‘non-stacking’ position of Y167 is found in all RlmJ structures and appears important for stabilisation of the catalytic loop conformation, rather than for the substrate adenine position. Y167 also—along with residues V199, Y173 and P197—provides a hydrophobic surface near the catalytic pocket where the guanosine is positioned (Figure 4E). Taken together, each nucleic base of the GA* conjugate is involved in hydrophobic interactions with RlmJ residues, and both adenines, A$^{\text{SAM}}$ and A$^{\text{sub}}$, form hydrogen bonds to conserved residues in RlmJ (Figure 4F).
helix is formed in the tail that packs against SAM and SAH (Figure 5D). In this ‘closed’ conformation of the tail, H9 is unable to π-stack with the RNA-substrate adenine due to steric clashes with H6 (Figure 5D). This conformation, if it exists in solution, is unlikely to be active. The crystal structure of RlmJ/SAM (PDB 4BLV), contains two monomers of RlmJ in the asymmetric unit. One monomer of SAM-bound RlmJ displays the ‘closed’ conformation, while the second one exhibits the open conformer with residues 1–8 adopting an extended conformation with no helix present. This suggests that, in the SAM-bound state, RlmJ is capable of populating these different conformations, at least in the crystal, with the binding of the RNA substrate requiring or selecting the open conformation of the N-terminal tail to clamp the RNA in the RlmJ active site.

Deciphering the RNA recognition mode and methyl transfer used by RlmJ

From the structures of RlmJ bound to GA* and GA*A, we constructed a model of RlmJ bound to the SAM cofactor and a trinucleotide RNA covering 2029GAA2031 of the E. coli 23S rRNA sequence (Figure 6A, B). This model kept the structures of RlmJ and the GA dinucleotides of the RlmJ/GA* complex and that of A5SAM and the methionine chain of GA* to build the SAM. The third nucleotide, A2031, was added and positioned as the A3' of GA*A in the RlmJ/GA*A structure. The phosphodiester backbone conformation of A2031 was adjusted to enable its linkage to A2030. To challenge the model, RlmJ residues in the GAA-binding interface (Figure 6A) were mutated and the mutant proteins were tested for MTase activity with the E. coli 23S rRNA hairpin (Figure 1A, Figure 6C). Mutated residues were selected according to the model as: (i) the two conserved aromatic residues in the N-terminal tail of RlmJ (F8, H9) that π-stack with A2031 and A2030, respectively, (ii) the three conserved residues (N12, K18 and W195) interacting with the target adenine A2030 and iii) the three residues (Y167, Y173 and V199) interacting with G2029. All mutant proteins are still folded and able to bind the SAM cofactor (Supplementary Figure S39). They all show a reduced MTase activity compared to the wild-type RlmJ, particularly for the mutations K18A and W195 that abolish the MTase activity. This confirms the contribution of F8 in the RNA-substrate binding, the involvement of H9, N12, K18 and W195 in the binding of the target adenine A2030 and the presence of a hydrophobic cavity made by the side chains of Y167, Y173 and V199 to bind G2029. Mutation of D164 to alanine was previously shown to abolish RlmJ MTase activity (24). Altogether, these data demonstrate that the SAM-RNA conjugates have been efficient to reliably investigate the RNA-binding determinants used by RlmJ to specifically recognize and modify its RNA substrate.

Based on this model, we propose here a mechanism for the methyl transfer reaction catalyzed by RlmJ (Figure 6D). The proper placement of adenosine 2030 is conditioned by π−π stacking with the aromatic side chains of H9 and W195 and by hydrogen bonds with N12 and K18. The N-terminal flexible tail is used to bind nucleosides in the loop of the 23S rRNA, F8 for A2031, H9 for A2030 and Y4 and H6 for two potential other nucleosides, explaining how the loop is bound by RlmJ. The N6 amino group of the A2030 is engaged in two hydrogen bonds, which enhance the nucle-
Figure 6. Model of a trinucleotide bound to RlmJ in position to be N6-methylated on the central adenosine. (A) RlmJ surface shown in pale cyan. (B) RlmJ secondary structure shown in pale cyan and side chains of residues interacting with the trinucleotide (in orange) shown as sticks. The model was built from the crystal structures of RlmJ/GA* and RlmJ/GA*A. (C) Methyltransferase assays of RlmJ variants mutated in the RNA binding interface. (D) Proposed mechanism of methyl transfer catalyzed by RlmJ.

The ophilicity of the N6 and triggers the SN2 reaction into the electrophilic methyl group (Figure 6D).

CONCLUSION

We have developed a method to synthesize RNA–SAM conjugates based on a convertible nucleoside strategy starting from O6-(benzotriazolyl)inosine. The O6-(benzotriazolyl)inosine can be introduced by phosphoramidite chemistry either at the 5′-end, 3′-end or an internal position of an oligonucleotide. The SNAr reaction in the presence of a SAM analogue containing an alkyl linker was optimized and efficient deprotection conditions to provide fully deprotected dinucleotide- or trinucleotide-SAM conjugates were developed. We were also able to synthesize a 13mer RNA–SAM conjugate using an enzymatic ligation. The strategy, devised here, is chemoselective and enables us to modify the size and the sequence of the RNA and to connect specifically the SAM cofactor to the adenine, which is methylated by m6A RNA MTases. Crystal structures of RlmJ/GA* and RlmJ/GA*A complexes have been solved and allowed us to construct a model of RlmJ bound to the SAM cofactor and a trinucleotide. This model is corroborated by site-directed mutagenesis and MTase activity assays and allows us to propose a mechanism for the methyl transfer reaction catalyzed by RlmJ. We have thus demonstrated here that RNA–SAM conjugates can be used to decipher RNA recognition by m6A RNA MTases. These structures were critical to obtain structural data impossible to obtain to date using fragments of the substrate RNA.

The conjugates described here can be used for any m6A MTases to probe their active site, by adapting the RNA sequence to the RNA substrate of the MTase of interest. In the future, chemistry that allows coupling of the cofactor to different positions in the RNA base or sugar, would facilitate the production of a full co-crystallization library of RNA-cofactor conjugates, matching each family of proteins catalyzing post-transcriptional modifications of RNA. Such a library would be a major advancement in the fields dealing with substrate recognition and enzymatic mechanisms of these protein families.

DATA AVAILABILITY

Atomic coordinates and structure factors for the reported crystal structures have been deposited with the Protein Data bank under accession number 7P90, 7P91, 7P8Q.
SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS

The authors are grateful to Jacqueline Plumbridge (IBPC) for careful reading of the manuscript.

FUNDING

CNRS; ANR ARNTools [ANR-19-CE07-0028]; Labex DYNAMO [ANR-11-LABX-0011]; Equipex CAC-SICE [ANR-11-EQPX-0008]. Funding for open access charge: ANR, RNATools [ANR-19-CE07-0028].

Conflict of interest statement. None declared.

REFERENCES

1. Boccaletto, P., Machnicka, M.A., Purta, E., Piatkowski, P., Baginski, B., Wirecki, T.K., de Crecy-Lagard, V., Ross, R., Limbach, P.A., Kotter, A. et al. (2018) MODOMICS: a database of RNA modification pathways. Nucleic Acids Res. 46, D303–D307.
2. Carell, T., Brandmayr, C., Hienzsch, A., Müller, M., Pearson, D., Reiter, V., Thomass, P. and Wagner, M. (2012) Structure and function of noncanonical nucleobases. Angew. Chem. Int. Ed. 51, 7110–7131.
3. Wiener, D. and Schwartz, S. (2017) The RNA modification N(6)-methyladenosine and its implications in human disease. Genomics Proteomics Bioinformatics, 15, 154–163.

16. Adtijan, C., Coelho, D., Iannazzo, L., Ethene-Quelquejeu, M. and Braud, E. (2020) Synthesis of triazole-linked SAM-Adenosine conjugates: functionalization of adenosine at N-1 or N-6 position without protecting groups. Molecules, 25, 3241.
17. Ahmed-Belkaecem, R., Surto-Oritz, P., Guiraud, M., Canard, B., Vasseur, J.J., Decroly, E. and Debart, F. (2020) Synthesis of adenosine dimethylsides SAM analogs as specific inhibitors of SARS-CoV-2 nsp14 RNA cap guanine-N7-methyltransferase. Eur. J. Med. Chem., 201, 112572.
18. Deen, J., Vranken, C., Leen, V., Neely, R.K., Janssen, K.P.F. and Hofkens, J. (2017) Methyltransferase-Directed labeling of biomolecules and its applications. Angew. Chem. Int. Ed. Engl., 56, 5182–5200.
19. Mattay, J., Dietmar, M. and Rentmeister, A. (2020) Chemoenzymatic strategies for RNA modification and labeling. Curr. Opin. Chem. Biol., 63, 46–56.
20. Kaniskan, H.U. and Jin, J. (2015) Chemical probes of histone lysine methylationtransferases. ACS Chem. Biol., 10, 40–50.
21. Hobley, G., McKelvie, J.C., Harmer, J.E., Howe, J., Oyston, P.C. and Rouch, P.L. (2012) Development of rationally designed DNA N6 adenine methyltransferase inhibitors. Bioorg. Med. Chem. Lett., 22, 3079–3082.
22. Halby, L., Menon, Y., Rilova, E., Pechalriou, D., Masson, V., Faux, C., Bouhlel, M.A., David-Cordonnier, M.H., Novosad, N., Aussagues, Y. et al. (2017) Rational design of bisubstrate-type analogues as inhibitors of DNA methyltransferases in cancer cells. J. Med. Chem., 60, 4665–4679.
23. Golovina, A.Y., Dzama, M.M., Osterman, I.A., Sergeiv, P.V., Serebryakova, M.V., Bogdanov, A.A. and Dontsova, O.A. (2012) The first rRNA methyltransferase of e. coli revealed: the yhiR gene encodes adenosine-N6 methyltransferase specific for modification of A2030 of 23S ribosomal RNA. RNA, 18, 1725–1734.
24. Pungekar, A.S., Liljerholm, J., Shepherd, T.R., Forster, A.C. and Selmer, M. (2013) Structural and functional insights into the molecular mechanism of rRNA m^6A methyltransferase rml3. Nucleic Acids Res., 41, 9537–9548.
25. Pignot, M., Stethoff, C., Linscheid, M. and Weinhold, E. (1998) Coupling of a nucleoside with DNA by a methyltransferase. Angew. Chem. Int. Ed. Engl., 37, 2888–2891.
26. Pjevaljcić, G., Schmidt, F. and Weinhold, E. (2004) Sequence-specific methyltransferase-induced labeling of DNA (SMILing DNA). Chem Bio Chem, 5, 265–269.
27. Osborne, T., Weller Roska, R.L., Rajski, S.R. and Thompson, P.R. (2008) In Situ generation of a bisubstrate analogue for protein arginine methyltransferase 1. J. Am. Chem. Soc., 130, 4574–4575.
28. Yu, H., Hendrick, C.E., Frye, K.S. and Comstock, L.R. (2012) Fluorescent DNA labeling by N-mustard analogues of S-adenosyl-L-methionine. ChemBioChem, 13, 2225–2233.
29. Kabsch, W. (2010) xds. Acta Crystallogr. D, 66, 125–132.
30. McCoy, A.J., Grosse-Kunstleve, R.W., Adams, P.D., Winn, M.D., Storoni, L.C. and Read, R.J. (2007) Phaser crystallographic software. Appl. Crystallogr. 40, 658–674.
31. Winn, M.D., Ballard, C.C., Cowtan, K.D., Dodson, E.J., Emsley, P., Evans, P.R., Keegan, R.M., Krissinel, E.B., Leslie, A.G.W., McCoy, A. et al. (2011) Overview of the CCP4 suite and current developments. Acta Crystallogr. Sect. D Biol. Crystallogr., 67, 235–242.
32. Yagin, 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. Sect. D Biol. Crystallogr., 60, 2184–2195.
33. Emsley, P. and Cowtan, K. (2004) Coot: model-building tools for molecular graphics. Acta Crystallogr. D Biol. Crystallogr., 60, 2126–2132.
34. Afonine, P.V., Grosse-Kunstleve, R.W., Echols, N., Headd, J.I., Moriarty, N.W., Mustyakimov, M., Terwilliger, T.C., Zwart, P.H. and Adams, P.D. (2012) Towards automated crystallographic structure refinement with phenix.refine. Acta Crystallogr. Sect. D Biol. Crystallogr., 68, 352–367.
35. Niesen, F.H., Berglund, H. and Vedadi, M. (2007) The use of differential scanning fluorimetry to detect ligand interactions that promote protein stability. Nat. Protoc., 2, 2212–2221.
36. Zhu, X.F. and Scott, A.I. (2008) Novel synthetic approach to multibenzoylated nucleosides. Synth. Commun., 38, 1346–1354.
37. Kaul, K., Brouillette, Y., Sajjadi, Z., Hansford, K. A. and Lubell, W. D. (2004) Selective tert-butyl ester deprotection in the presence of acid labile protecting groups with use of ZnBr₂. *J. Org. Chem.* **69**, 6131–6133.

38. Ferreira, F. and Morvan, F. (2005) Silyl protecting groups for oligonucleotide synthesis removed by a ZnBr₂ treatment. *Nucleoside Nucleotides Nucleic Acids* **24**, 1009–1013.

39. Chemama, M., Fonvielle, M., Villet, R., Arthur, M., Valery, J. M. and Etheve-Quelquejeu, M. (2007) Stable analogues of Aminoacyl-tRNA for inhibition of an essential step of bacterial cell-wall synthesis. *J. Am. Chem. Soc.*, **129**, 12642–12643.

40. Laskowski, R. A. and Swindells, M. B. (2011) LigPlot+: multiple ligand-protein interaction diagrams for drug discovery. *J. Chem. Inf. Model.*, **51**, 2778–2786.