Structure of a reaction intermediate mimic in t^6A biosynthesis bound in the active site of the TsaBD heterodimer from Escherichia coli

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ABSTRACT

The tRNA modification N6-threonylcarbamoyladenosine (t^6A) is universally conserved in all organisms. In bacteria, the biosynthesis of t^6A requires four proteins (TsaBCDE) that catalyze the formation of t^6A via the unstable intermediate L-threonylcarbamoyl-adenylate (TC-AMP). While the formation and stability of this intermediate has been studied in detail, the mechanism of its transfer to A37 in tRNA is poorly understood. To investigate this step, the structure of the TsaBD heterodimer from Escherichia coli has been solved bound to a stable phosphonate isosteric mimic of TC-AMP. The phosphonate inhibits t^6A synthesis in vitro with an IC_{50} value of 1.3 μM in the presence of millimolar ATP and L-threonine. The inhibitor binds to TsaBD by coordination to the active site Zn atom via an oxygen atom from both the phosphonate and the carboxylate moieties. The bound conformation of the inhibitor suggests that the catalysis exploits a putative oxanion hole created by a conserved active site loop of TsaD and that the metal essentially serves as a binding scaffold for the intermediate. The phosphonate bound crystal structure should be useful for the rational design of potent, drug-like small molecule inhibitors as mechanistic probes or potentially novel antibiotics.

INTRODUCTION

Transfer RNAs from all forms of life are subject to a large number of enzyme-catalysed post-transcriptional chemical modifications that fine tune both its structure and function in the cell (1–4). The N6-threonylcarbamoyladenosine (t^6A) modification (5) is found at position 37 adjacent to the anticodon in tRNAs that decode 5'-ANN codons (6,7), and is universally conserved throughout life (8). Structurally, t^6A enhances codon recognition by disrupting base pairing with U33 in the anticodon loop and by interacting with both anticodon bases as well as the first base of the mRNA codon (9). Mutants in both bacteria (10,11) and eukaryotes (11,12) that lack t^6A typically have severe growth defects or, as in the case of Escherichia coli, are inviable (13). In many organisms t^6A is further modified to the hydrolytically labile cyclized hydantoin cit^6A, which is formed by ATP-catalysed cyclization of t^6A (14,15) (Figure 1). This has prompted a call for re-evaluation of the structural explanation for its role in codon recognition. The catalytic machinery for the biosynthesis of t^6A differs significantly between bacteria and eukaryotes/archaea. In eukaryotes, t^6A is catalysed by the Tcs1 (SUA5) protein (10) in conjunction with the KEOPS complex (16–19), which contains five proteins. KEOPS was originally identified as a transcription complex (20) involved in telomere regulation (21). In addition to its requirement for t^6A tRNA modification, KEOPS proteins in yeast have t^6A-independent roles in telomere replication via SUA5p (22) and as an auxiliary sensor of DNA double strand breaks (23). KEOPS mutations have been recently implicated in neurodegeneration with nephrotic disease in humans (24–26). In mitochondria, a minimal t^6A system operates, consisting of Tcs1 and the mitochondrial protein Qri7 (27). Expression of Qri7 in the cytoplasm can complement loss of KEOPS proteins in t^6A formation. This result has implicated the KEOPS Qri7 homolog OSGEP (Tcs3) as the main catalytic component responsible for t^6A syn-
In most bacterial tRNAs, t^6A is further cyclized to the hydantoin ct^6A, which is hydrolytically unstable to commonly used isolation procedures. In human mitochondria, t^6A formation was recently found to function as a CO2 sensor, in that low CO2 levels result in low t^6A formation and reduced translation of mitochondrial proteins (28). Based on the dependence of t^6A in bacteria for normal growth, and the apparent divergence of the protein machinery between bacterial and eukaryotic t^6A synthesis, this modification has been identified as a potential antibiotic target.

As shown in Figure 1, the biosynthesis of t^6A in bacteria involves four distinct proteins (29). The first step in the biosynthesis is carried out by TsaC1 (YrdC)/TsaC2 (SUA5), which are short/long bacterial homologs, respectively, of eukaryotic Tcs1/SUA5. These enzymes catalyse an L-threonine-dependent adenylation reaction, using bicarbonate (possibly via CO2), to form a putative L-threonine carbamate that is adenylated with ATP to give the unstable product TC-AMP (30). This mechanism was also shown to operate in vitro for both eukaryotes and archaea (31). There is evidence that although TsaC1 and TsaC2 catalyse the same reaction, they may operate by different mechanisms due to the involvement in TsaC2 of additional hinge and SUA5 domains (32). The TsaBDE proteins together in the presence of ATP can catalyse the addition reaction of exogenously supplied TC-AMP with the tRNA substrate to give t^6A and AMP as products (30). Similarly, in yeast it was shown that SUA5p and Qri7 can catalyse t^6A formation via TC-AMP diffusion without direct protein-protein interaction (33). The bacterial TsaD protein is a homolog to the ancestral (COG0533) family of carbamoyltransferase proteins that include KAE1/OSGEP/TobZ that utilize carbamoyl adenylate substrates or intermediates to carbamoyl nucleophiles (34,35).

Figure 1. Biosynthesis of t^6A in bacteria. In most bacterial tRNAs, t^6A is further cyclized to the hydantoin ct^6A, which is hydrolytically unstable to commonly used isolation procedures.

A comparison of structural studies in bacteria show that the quaternary structure for the TsaBDE complex is species dependent. In E. coli and S. typhimurium, small angle X-ray scattering (SAXS) and isothermal calorimetry (ITC), respectively, were used to determine the ternary complex is a TsaB1D1E1 heterotrimer, the formation of which is ATP-dependent (36,37). In E. coli, there is a novel additional ADP binding site at the TsaB–TsaD interface (37). The T. maritima proteins form a hexameric TsaB2D2E2 complex as a dimer of heterotrimers with TmTsaB as the dimerization domain (38,39). The role of the TsaE protein in both systems appears to be regulatory in nature, in that TmTsaE binding to the TmTsaB2D2E1 complex has been shown to be competitive with tRNA (38). This is consistent with the observation in E. coli that EcTsaBDE does not bind tRNA (37), while EcTsaBD does (38). TsaE-catalysed ATP hydrolysis is required for additional turnover, possibly via a resetting of the active conformation of TsaBD (38).

Missoury et al. (39,40) observed that TmTsaE binding to TmTsaBD induces a partial denaturing of the TsaD active site. Both studies are in agreement that TmTsaE binding precludes the binding of tRNA. Questions remain as to the exact mechanism and timing of the conformational effects elicited by TsaE during the catalytic cycle, as well as the timing of tRNA binding relative to TC-AMP loading onto TsaD and the potential role of TsaC1/C2 in the latter. Each of these steps may be specifically tailored in mesophiles versus thermophiles, due in part to the instability of TC-AMP and the need to avoid wasteful ATP utilization.

was characterized initially as an ATPase with low intrinsic activity (42–44) that is stimulated by the presence of TsaBD (29,37). As shown in Figure 1, the chemical transformation from TC-AMP to t^6A does not formally require ATP. Indeed, bacterial t^6A formation is observed in vitro in the absence of this protein (30); however, as explicitly shown in T. maritima (38), TsaE catalysed ATP-hydrolysis is required for multiple turnover.

A comparison of structural studies in bacteria show that the quaternary structure for the TsaBDE complex is species dependent. In E. coli and S. typhimurium, small angle X-ray scattering (SAXS) and isothermal calorimetry (ITC), respectively, were used to determine the ternary complex is a TsaB1D1E1 heterotrimer, the formation of which is ATP-dependent (36,37). In E. coli, there is a novel additional ADP binding site at the TsaB–TsaD interface (37). The T. maritima proteins form a hexameric TsaB2D2E2 complex as a dimer of heterotrimers with TmTsaB as the dimerization domain (38,39). The role of the TsaE protein in both systems appears to be regulatory in nature, in that TmTsaE binding to the TmTsaB2D2E1 complex has been shown to be competitive with tRNA (38). This is consistent with the observation in E. coli that EcTsaBDE does not bind tRNA (37), while EcTsaBD does (38). TsaE-catalysed ATP hydrolysis is required for additional turnover, possibly via a resetting of the active conformation of TsaBD (38).

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While a structure of TC-AMP bound to TsaC2 from *Sulfolobus tokodaii* has been reported (34,45), there are as yet no structures of TC-AMP bound to the TsaBD heterodimer. Luthra et al. (41) have recently reported the structure of *T. maritima* TsaB2D2E2 with the unstable ligand carboxy-AMP bound in the TsaD active site. This structure was used to model TC-AMP binding to the complex. Efforts at solving a structure of TsaBD with TC-AMP itself have been hampered by the known instability of this substrate (30). In an effort to circumvent this barrier, we report here the preparation of chemically stable analogs of TC-AMP and an assessment of their ability to inhibit the formation of t6A in the *E. coli* system. The most potent inhibitor is an isoster of phosphonate analog, in which the adenylate ester oxygen of TC-AMP is replaced by a methylene group. We show here that this stable analog binds to the EcTsBD heterodimer with affinity in the low micromolar range and we report the crystal structure of the bound complex. Interestingly, the active site Zn ion interacts with the inhibitor by chelating oxygens from both the carboxylate of the L-threonine moiety as well as one of the oxygen atoms of the phosphonate. The structure suggests that TsaBD binds TC-AMP inside the catalytic pocket in an orientation that may be initially sterically unreactive but a simple conformational rearrangement may lead to a more reactive bound form upon binding of tRNA.

**MATERIALS AND METHODS**

**Protein purification and preparation of site-directed mutants**

Proteins for kinetic assays and binding analysis were overproduced and purified as previously described (30). Site-directed mutants of *E. coli* TsaD were prepared using Quikchange mutagenesis (Agilent Technologies) with *tsaD* cloned into pET28a as the template (30) and pfuTurbo DNA polymerase. Oligonucleotides for mutagenic PCR are from Integrated DNA Technologies, Inc. and are listed in Supplementary Table S1. Plasmids were purified using the QIAprep spin miniprep kit (QIagen) and both strands of the plasmid sequence were determined (UW Biotechnology Center). For assays, proteins were quantified using the BioRad Bradford Protein Assay.

**Preparation of tRNA transcripts**

Unmodified tRNA was prepared by *in vitro* runoff transcription using overlapping oligonucleotides essentially as described (46,47). T4 DNA polymerase Klenow fragment (Promega) was used to extend overlapping oligonucleotides containing a 12-bp overlapping region and a fused T7 promoter sequence. For *E. coli* tRNA, the oligos utilized were the following: Plus strand: 5′-TAATACGACTCACTATAGGGTGCTTACGTAAGCTGTTAGCTCA GTTGGTAGAGCAGTGGACTTTTAATC and minus strand: 5′-TGGTTGGCTGTCGAGGAATTGACGACC TGCCAGCAATGGATTGAACGATAC (T7 promoter underlined and overlapping bases in italics). The extension reaction used 1 mM each oligo and 0.4 mM dNTPs with 10 units/ml polymerase in the manufacturer supplied buffer at 37°C for 2 h. DNA was isolated by phenol then chloroform extraction and ethanol precipitation. The redissolved DNA was then used directly for *in vitro* transcription. Transcription reactions contained 1 µM template DNA in a reaction mixture consisting of 50 mM Tris, pH 7.8, 35 mM MgCl2, 1 mM spermidine, 5 mM of each NTP, 20 units/ml RNase (Promega) and 1000 units of T7 RNA polymerase (Lucigen). After incubating at 37°C overnight, 10 units of RNase-free DNase was added directly to the mixture and incubated at 37°C for 30 min. After addition of an equal volume of 7.5 M urea with 0.1% bromphenol blue, the RNA was purified by polyacrylamide gel electrophoresis using 8% slab gels and run at 20 W for 1.5 h. RNA bands were visualized by UV shadowing on a fluorescent TLC plate (Merck). The RNA was excised and eluted in 0.5 M NaCl, filtered and then precipitated with 2.5 vol ethanol at −20°C. After centrifugation at 14 000 × g at 4°C (Beckman Avanti-J 25.50 rotor), the RNA pellet was washed with cold 70% ethanol, air-dried briefly and resuspended in deionized water and stored at −80°C. Typical yields were 1.6 mg of tRNA transcript/ml of transcription reaction.

**In vitro assay for t6A formation**

An HPLC assay was used to quantify t6A by UV detection at 260 nm. Full reaction mixtures (50 µl) contained 50 mM MOPS pH 7.5, 20 mM NaHCO3, 25 mM KCl, 50 mM L-threonine, 1 mM ATP, 10 µM unmodified *E. coli* tRNA1–35, and 1 µM each of TsaC, TsaB, TsaD and TsaE. For substrates with apparent *Km* values <10 µM, 0.5 µM enzyme concentrations were used.) For kinetic studies, one of the substrates was varied while all other substrates were kept at the concentrations stated above. Reactions were initiated with enzymes by adding TsaE, TsaB, TsaD and then TsaC as the last component followed by mixing by vortexing for 1 s. Reaction mixtures were incubated at 23°C for 5 min and quenched by adding 25 µl of the reaction mixture to a new tube containing 4 µl of 0.25 M EDTA pH 8 followed by vortexing and placing on dry ice. Once a set of reactions was complete, tubes were thawed and placed in a microcentrifuge (Eppendorf) and spun at 14 000 rpm for 10 s. Small molecules were removed by addition of the quenched reaction mixtures to micro biopan columns (BioRad) containing 0.7 ml (settled volume) Sephadex G-50 (fine), which had been drained by centrifugation at 7000 rpm for 1 min using a microcentrifuge. After careful addition of the reaction mixture to the top of the gel, the column was eluted by centrifugation at 7000 rpm for 1 min. To the resulting RNA eluate was added 0.1 volume of 0.3 M NaOAc, 1 mM Zn(OAc)2, pH 6, and the tRNA was digested by addition of 1.5 units of nuclelease P1 and 0.05 units of alkaline phosphatase for ≥2 h at 37°C. The digested RNA (~45 µl) was analysed directly by HPLC using a Supelco LC-18S column (250 × 4.6 mm) similar to reported methods (48,49). The HPLC analysis used an isocratic buffer system consisting of 5 mM ammonium acetate pH 5.3, containing 4%/v/v acetonitrile. Under these conditions the nuclelease P1 reaction was the rate determining enzyme in the digestion. If insufficient nuclease or digestion time was used for this step, an additional peak is observed at a longer retention time (24 min versus 21 min for t6A). Material isolated from this peak was consistent with the dinucleotide 5′-t6AApA-3′ as determined by.
enzymental analysis. For kinetic analysis of TC-AMP as the substrate, TsaC, l-threonine and bicarbonate were omitted, and the reaction was initiated by addition of TC-AMP.

**Synthesis of TC-AMP mimic BK951 (1)**

Figure 3 shows the overall synthetic scheme. Experimental details for the synthesis of analogs 2–5 can be found in the Supplementary Material.

**N-o-(2-Chloroacetyl)-O-benzyl-l-threonine benzyl ester (7)**

Chloroacetyl chloride (3.19 ml, 0.040 mol) was added drop wise to a solution of K₂CO₃ (5.8 g, 0.041 mol) and MgSO₄, and filtered. The solvent was removed under reduced pressure to give 7 as a white solid (1.23 g, 94%). 1H NMR (400 MHz, CDCl₃) δ 7.74–7.75 (m, 2H), 7.04 (d, J = 8.51 Hz, 1H), 4.52, 4.30 (ABq, JAB = 12.00 Hz, 2H), 4.19 (qd, J = 6.19, 2.32 Hz, 1H), 4.04 (s, 2H), 1.22 (d, J = 6.19 Hz, 3H).

**No-[2-(Dimethoxyphosphinyl)acyetyl]-O-benzyl-l-threonine benzyl ester (8a)**

A round bottom flask equipped with a stir bar was charged with 8a (1.94 g, 0.00432 mol) and CH₂CN (85 ml). After the addition of LiBr (0.750 g, 0.00864 mol), the mixture was brought to 75°C and stirred for 3 h. Upon completion, the solvent was removed under reduced pressure and the crude oil was diluted with H₂O (200 ml) and 1M HCl solution (50 ml). This aqueous layer was extracted with DCM (4 × 75 ml). The organic fractions were combined, washed with brine (1 × 100 ml), dried with MgSO₄, filtered, and the solvent was removed under reduced pressure to give 8b as a hard, pale white film (1.5 g, 80%). 1H NMR (400 MHz, CDCl₃) δ 11.05 (br. s., 1H), 9.96–9.53 (m, 11H), 4.95–5.22 (m, 2H), 4.74 (d, J = 8.89 Hz, 1H), 4.45, 4.24 (ABq, JAB = 11.75 Hz, 2H), 4.11 (s, 1H), 3.65 (m, 3H), 2.85–3.10 (m, 3H), 1.01–1.36 (m, 3H). 13C NMR (101 MHz, CDCl₃) δ 170.2, 166.5, 138.0, 135.5, 128.6, 128.5, 128.4, 127.8, 127.4, 70.8, 67.3, 57.2, 16.2. 31P NMR (162 MHz, CDCl₃) δ 21.48. MS (ESI): m/z 434 (M-H⁻). HRMS calculated for C₂₁H₂₆NO₇P (M⁻H⁻): 434.137412; Found: 434.13863.

**5’-O-[(Hydroxy)(2-oxo-2-(O-benzyl-l-threoninebenzylester)ethyl)phosphinyl]-6-azido-9-(2’,3’-O-isopropylidene-d-ribofuranosyl)purine sodium salt (11)**

A round bottom flask was charged with a diastereomeric mix of 10 (0.4 g, 0.54 mmol) and stir bar. After addition
of DMF (5.83 ml) and NaN₃ (0.087 g, 1.35 mmol), the reaction was allowed to stir overnight at 60°C. TLC the following morning confirmed the absence of starting material. Following the removal of solvent under reduced pressure, the crude oil product was purified directly by column chromatography on silica gel using 10% H₂O in CH₂CN as a running solvent. 1H was isolated as a single diastereomer, resembling an off white solid (0.3 g, 73%). It should be noted that a small fraction (>10%) of the sample appears to exist as the N1, C6 tetrazole tautomer in CD3OD. 1H NMR (400 MHz, CD3OD) δ 9.81 (s, 1H), 8.81 (s, 1H), 7.10–7.28 (m, 1H), 6.40 (d, J = 2.64 Hz, 1H), 5.39 (br. s., 1H), 5.18 (d, J = 4.70 Hz, 1H), 5.04 (s, 2H), 4.66 (m, 1H) 4.55 (br. s., 1H), 4.47, 4.25 (ABq, JAB = 11.75 Hz, 2H), 4.06–4.18 (m, 3H), 2.68–2.93 (m, 2H), 1.56–1.65 (m, 3H), 1.38 (s, 3H), 1.17 (m, 3H). 10C NMR (126 MHz, CD3OD) δ 147.2, 145.0, 143.7, 139.9, 137.4, 130.0, 129.9, 129.8, 129.9, 129.4, 129.1, 115.7, 93.8, 88.4, 88.3, 87.1, 83.9, 76.2, 75.3, 68.7, 66.8, 28.2, 26.3, 17.2. 31P NMR (202 MHz, CD3OD) δ 14.17. MS (ESI): m/z 735 (M+). HRMS calculated for C33H36N8O10P (M+): 735.22975; Found: 735.22854.

Enzymatic preparation of TC-AMP and SC-AMP

Reaction mixtures (80 μl) contained 50 mM MOPS pH 7.5, 10 mM MgCl2, 25 mM KCl, 2 mM ATP, 50 mM L-threonine, 40 mM NaHCO3 and 42 μg (37 μM) of EcTsaC. After incubating at 23°C for 10 min, the reaction mixture was injected onto a Supelco LC-18T HPLC column (250 mm × 4.6 mm) and eluted with an isocratic buffer of 50 mM potassium phosphate pH 6.0. Typical yields were 25 nmol at a concentration of 20–30 μM for TC-AMP. Solutions could be carefully concentrated by rotary evaporation (water bath < 30°C) and used directly or aliquoted and stored at −80°C. (Approximately 10% degradation was typically observed for each freeze/thaw cycle.) The serine analog of TC-AMP (SC-AMP) was also prepared by substituting L-serine for L-threonine and purifying by an analogous method. Typical yields for SC-AMP were 10 nmol per reaction. The identity of SC-AMP was confirmed by accurate mass determination (HRMS Calcd for MH+ 479.0922; Found 479.09214) and purity was assessed by HPLC (>95%, Supplementary Figure S5).

Data analysis for determining kinetically derived parameters

For kinetic analysis, concentrations for all fixed substrates were the same as described above for the full reaction mixture. Reactions were run in duplicate and values for Kₘ and kₖcat were derived from fitting to the standard Michaelis–Menten model (Equation 1). For substrates with Kₘ values near the enzyme concentrations (<5-fold), kinetic values were derived from Morrison's equation (50) for tight binding substrates (Equation 2). For kinetics that show substrate inhibition, the data were fit to a simple model for complete substrate inhibition in which the complex containing two bound substrates is inactive (Equation 3). Error values reported are the standard error for the fit. All data analysis was performed using Prism 8 (GraphPad) software. Values for kₖcat in Tables 1 and 2 were obtained by dividing V₀ by [E]ₘₐₓ.
(\(V_{\text{max}}\)) values obtained from curve fitting by the known enzyme concentration.

\[
v = \frac{V_0 [S]}{K_M + [S]} \quad (1)
\]

\[
v = \frac{V_0}{(\frac{[E_Y]+[S]+K_M)}{2}) - \sqrt{\frac{([E_Y]+[S]+K_M)^2 - 4[E_Y][S]}}}{2}\quad (2)
\]

\[
v = \frac{V_0 [S]}{K_M + [S] \left(1 + \frac{[S]}{K_i}\right)} \quad (3)
\]

Microscale thermophoresis (MST) binding studies

MST measurements were performed using a Monolith NT115.1 Pico instrument (Nanotemper Technologies). EcTsaD was fluorescently labelled with the Monolith Red-NHS second generation dye (cat. no. MO-L011) using the manufacturer’s protocol. Briefly, purified His6-TsaD (20 μM) in buffer (50 mM sodium phosphate pH 7.5, 300 mM NaCl, 5% glycerol) was mixed with an equal volume (100 μl) of dye reagent (60 μM) and column provided in the kit to give the cyan-colored protein. Fluorescently labelled TsaD was analysed for t6A activity using the full reaction kit to give the cyan-colored protein. Aliquots of the labelled protein were frozen in dry ice acetone and stored at −80°C. Binding measurements were conducted in t6A Buffer (equal to full assay buffer described above) containing PBS with 0.05% Tween 80. Full t6A activity was observed in this buffer. A stock concentration of 1 μM EcTsaD: 2 μM EcTsaB was prepared in binding buffer and dilutions were carried out to give aliquots of added to separate micro (PCR) striptubes containing 10 nM of the labelled TsaBD complex. After an initial addition of an equal volume (10 μl) of 1 mM of the ligand BK951 was made to the first tube, serial dilutions were prepared to span the range of 0.1–500 μM BK951 and 5 nM EcTsaBD complex. A glass capillary was then filled with the binding reaction mixture from each tube and placed in the MST tray. The capillaries were analysed for thermophoretic mobility. The instrument software analyses for fluorescence homogeneity in the samples and artefacts such as ligand induced autobleaching, as well as protein aggregation, which could lead to erroneous results. When suitable conditions are found, binding data is collected and analysed by the software package included with the instrument. Binding measurements were performed in triplicate.

TCAMP stability studies in complex with EcTsaBD and EcTsaC

To study the effects of TsaBD on TC-AMP stability, HPLC-purified TC-AMP in 50 mM potassium phosphate was added to a solution of TsaBD in 50 mM HEPES pH 7.5, 0.3 M NaCl to give 100 μl of 10 μM TC-AMP with or without 40 μM TsaBD. The mixtures were incubated at 37°C and 10 μl timed aliquots were withdrawn and analysed by HPLC for AMP and TC-AMP by measuring peak areas using absorbance at 260 nm. Plots of ln[TC-AMP]/\([\text{TC-AMP} + \text{AMP}]\) vs time were analysed by linear regression (GraphPad Prism 8) to get a series of rate constants for first order TC-AMP decomposition to AMP. To measure the effect of TsaC on TC-AMP stability, EcTsaC was overproduced in BL21(DE3) cells from IPTG induction of the gene cloned in pET15b. The protein was purified using Ni-NTA chromatography and buffer exchanged with a 10 μl Sephadex G-50 column as previously described (30). This procedure typically gives 1 ml of 100–150 μg EcTsaC, which contains 70–80% bound TC-AMP. The experiments described in this work used 130 μM TsaC with 100 μM (77%) bound TC-AMP as determined by HPLC and Bradford protein assay. The stability of this complex at different concentrations at 37°C in the same reaction buffer was analysed using the same HPLC protocol as described for TC-AMP: TsaBD mixtures to get a series of rate constants for each concentration of the complex.

Fluorescence anisotropy binding measurements

The binding of the \(E.\ coli\) TsaBD complex to 3′-Fluorescein-labelled tRNA\(^{53}\text{UUU}\) was monitored with a TECAN INFINITE F200PRO microplate reader by using a Nunc F96 microwell plate (Thermo Scientific). 40 nM of 3′-Fluorescein-labelled tRNA probe was incubated in each well (final volume of 200 μl) with increasing amounts of TsaBD (final concentrations between 0 and 3000 nM) in 20 mM HEPES pH 7.5, 4 mM NaCl and 2 mM DTT supplemented or not with BK951 (final concentration 50 μM). The plate was incubated at 22°C in the dark for 30 min and then fluorescence anisotropy was recorded at 535 nm (λ\(_{\text{exc}}\) = 495 nm). Titration curves have been obtained by fitting experimental values to the following hyperbolic saturation function: \(A = B_{\text{max}} \times [\text{TsaBD}] / (K_D + [\text{TsaBD}])\) using SigmaPlot (Systat) software. Measurements have been performed in triplicate and mean values have been used for fitting.

Cloning, expression, purification and crystallization of TsaD and TsaB from \(E.\ coli\)

Genes coding EcTsaD and EcTsaB have been cloned in pET9a expression plasmids and proteins have been produced and purified as previously described by Zhang et al. (37). TsaBD complex was concentrated to 10.1 mg.ml\(^{-1}\) for crystallization trials. Crystals of heterodimer TsaBD from Escherichia coli were obtained by hanging-drop vapor diffusion after at 18°C by mixing 1 μl of the protein solution pre incubated with 2.5 mM BK951 and 5 mM MgCl\(_2\) with 1 μl of 0.25 mM lithium sulfate 0.11 M HEPES pH 7.5 0.15 mM sodium acetate and 25% PEG 4K. The crystallization drop was equilibrated against a 500 μl reservoir solution containing 0.25 mM lithium sulfate, 0.11 M HEPES pH 7.5 0.15 mM sodium acetate and 25% PEG 4K. Crystals were cryo-protected by quick-soaking in crystallization buffer supplemented with 25% ethylene glycol, 3 mM BK951 and 6 mM MgCl\(_2\) prior to flash freezing in liquid nitrogen.
Table 1. Summary of kinetically-derived parameters for t6A overall and partial reactions

| Ligand         | Overall reaction | TsaBDE reaction | TsaC reaction |
|----------------|------------------|-----------------|--------------|
|                | $K_m$ (µM)       | $k_{cat}$ (min$^{-1}$) | $K_m$ (µM)     | $k_{cat}$ (min$^{-1}$) | $K_m$ (µM) | $k_{cat}$ (min$^{-1}$) |
| L-Thr          | 590 ± 90         | 0.42 ± 0.02     | —             | —                      | 1.77 ± 0.26 | 9.0 ± 0.4  |
| ATP            | 60 ± 6 (K$_m$ = 2200)$^d$ | 0.40 ± 0.4     | 14 ± 3 (K$_m$ = 1800)$^d$ | — | 0.12 ± 0.1 | 1.03 ± 0.01 | 7.9 ± 0.2  |
| tRNA$^{5'},{UUU}$ | 0.5 ± 0.2 (K$_d$ = 87 ± 11) | 0.50 ± 0.03 | nd$^b$  | nd | 1.5 ± 0.4 | 0.31 ± 0.02 |
| TC-AMP         | 1.3 ± 0.5        | 7.5 ± 1.1 (K$_m$ = 2.5)$^d$ | (K$_d$ = 3.4 ± 0.6)$^e$ | — | >180 | — | — |
| 2              | 51.2 ± 1.4       | 123 ± 1.2 (K$_m$ = 41)$^d$ | (K$_d$ = 33)$^d$ | — | 31 ± 1.1 | — | — |
| 3              | 45.5 ± 1.1       | 99 ± 1.1 (K$_m$ = 33)$^d$ | 97 ± 1.4 | — | — | — | — |
| 4              | >1000            | nd$^f$          | nd          | — | nd | — | — |
| 5              | >1000            | Nd              | Nd          | — | Nd | — | — |
| SC-AMP         | 9.3 ± 1.2        | Nd              | Nd          | — | Nd | — | — |
| 5'-ADP         | 160 ± 20         | Nd              | Nd          | — | Nd | — | — |
| 5'-AMP         | 113 ± 10         | Nd              | Nd          | — | Nd | — | — |

$^a$Calculated for observed substrate inhibition.
$^b$Not determined.

Table 2. Summary of kinetically-derived inhibition and thermodynamically-derived binding constants of TCAMP and its analogs to t6A proteins

| Ligand | Overall reaction$^a$ | TsaBDE reaction$^b$ | TsaC reaction$^c$ |
|--------|----------------------|---------------------|--------------------|
| BK951  | 1.3 ± 0.5            | 7.5 ± 1.1 (K$_m$ = 2.5)$^d$ | >180                      |
| 2      | 51.2 ± 1.4           | 123 ± 1.2 (K$_m$ = 41)$^d$ | 31 ± 1.1           |
| 3      | 45.5 ± 1.1           | 99 ± 1.1 (K$_m$ = 33)$^d$ | 97 ± 1.4           |
| 4      | >1000                | nd$^f$              | nd                  |
| 5      | >1000                | Nd                  | Nd                  | — | — |
| SC-AMP | 9.3 ± 1.2            | Nd                  | Nd                  |
| 5'-ADP | 160 ± 20             | Nd                  | Nd                  |
| 5'-AMP | 113 ± 10             | Nd                  | Nd                  |

$^a$ATP = 1 mM, L-Thr = 0.6 mM.
$^b$ATP = 1mM.
$^c$ATP = 1.2 mM, L-Thr = 1.8 mM.
$^d$Calculated using Cheng–Prusoff equation (59).
$^e$Obtained using MST with fluorescently labelled EcTsaBD.
$^f$Not determined.

Data collection and structure determination

X-ray diffraction data collection was carried out on beamline Proxima2 at the SOLEIL Synchrotron (Saint-Aubin, France) at 100 K. Data were processed, integrated and scaled with the XDS program package. TsaBD belonged to space group $P1$ with unit cell parameters of $a = 63.23$ Å, $b = 69.25$ Å, $c = 86.54$ Å; $\alpha = 109.38^\circ$, $\beta = 92.116^\circ$, $\gamma = 116.94^\circ$. The structure was solved by molecular replacement using the PHASER module implemented in the CCP4 software package (search model was EcTsaBD, PDB ID: 4YDU). Two copies of the TsaBD dimer were found into the asymmetric unit of the crystal. The $2F_o - F_c$ electronic density indicated the presence of the BK951 inhibitor bound to a zinc ion. The initial structure was refined using the BUSTER program and completed by interactive and manual model building using COOT. Data collection and refinement statistics are gathered in Table 3. The refined structure was deposited at the Protein Data Bank (code 6Z81). All figures of structures were generated with PyMol (The PyMOL Molecular Graphics System, Version 2.0 Schrödinger, LLC).

Molecular docking of adenine into the active site of TsaD

To find out how A37 of tRNA substrate could be oriented into the active site, molecular docking was performed using AutoDockTools 4.2. Adenine was retrieved from the PDB and converted using the Babel format molecule converter (http://openbabel.org/). We removed TsaB from the present crystal structure (keeping TsaD bound to BK951) and added Kollman charges. The Lamarckian Genetic Algorithm (LGA) implemented in Autodock 4.2 was used for docking experiments. The docking parameters were as fol-
wherein L-threonine and inosine would provide the sources of synthesis (Figure 3), we envisioned a convergent strategy, 

\[ \text{activation of the 5'-hydroxyl group} \]

and reduction of the azide to the amine at N6 was achieved through a reaction catalysed by TsaBDE. Such geometry could be selectively stabilized by TsaBD binding. Compounds 4 and 5 employed malonates as the bridging group between the ribose and threonine and were designed to promote potential Zn binding to the active site of TsaD, via the enol tautomers of the beta-amido ester (4) or amide (5). Final products were purified by HPLC and characterized by NMR and High Resolution MS, which were consistent with the assigned structures (see Supplementary Material).

RESULTS

Rationale and preparation of stable mimics of TC-AMP

Because TC-AMP is unstable, we sought to prepare chemically stable analogs that are close structural mimics in order to obtain inhibitors that could provide structural information on the bound conformation of TC-AMP itself when bound to TsaBD. The structures of the mimics are shown in Figure 2. We began with the isosteric phosphonate BK951 (1). Simple replacement of the ester oxygen of the carbamate in TC-AMP gives a molecule that is stable to all of the observed TC-AMP degradation pathways that result in the formation of AMP. These include cyclization, direct hydrolysis or (potential) elimination pathways (30). For the synthesis (Figure 3), we envisioned a convergent strategy, wherein L-threonine and inosine would provide the sources of stereochemistry in the target molecule. The retrosynthesis employs a key disconnection between one of the phosphonate ester oxygens and the ribose 5'-carbon, which may be linked via Mitsunobu coupling (51). 6-Chloropurine riboside was used instead of an adenosine derivative to minimize the tendency of adenosine to cyclize at N3 during activation of the 5'-hydroxyl group (52). Protecting groups included an isopropylidene for the 2',3'-cis-diol of the riboside, while L-threonine was protected using benzyl groups at both the β-hydroxyl and carboxylic acid groups. The 6-chloro could be converted to a 6-azido functionality using azide, and the benzyl groups could then be removed during azide reduction to regenerate the adenosine moiety in a single step. As shown in Figure 3, the route begins with the starting dibenzyl protected L-threonine derivative 6, which was reacted with chloroacetyl chloride to give 94% of the chloride 7, which itself was converted to the dimethylphosphonate by Arbuzov reaction with trimethylphosphite in 78% yield. Dealkylation with LiBr in acetonitrile gave the phosphonate monoester, which was coupled with the isopropylidene protected 6-chloroinosine 10 in 78% yield. Reaction of 10 with sodium azide in DMF with heating gave the 6-azido purine derivative with concomitant demethylation to the phosphonic acid monoester. Global deprotection and reduction of the azide to the amine at N6 was achieved by hydrogenation using PdCl₂ in MeOH, followed by acetylation with TFA to complete the removal of the isopropylidene group to give crude 1. Purification of 1 by reversed phase HPLC gave the pure material in 36% yield from 10 on 100 mg scale. This material was >99% pure (Supplementary Figure S5) and used for both the enzymatic inhibition assays and structural analysis with TsaBD.

A series of other bioisosteres of TC-AMP were prepared (see Supplementary Data and Figures S1-S4 for experimental details). These include sulfamate isosteres 2 and 3, in which either the phosphonate only or both phosphonate and the amide of the intermediate were replaced by sulfamoyl and sulfonyleura groups, respectively. Sulfamoyl groups are well described successful substitutions as phosphodiester isosteres for the preparation of adenylate mimics in the inhibition of aminoacyl-tRNA synthetases and many other adenylate intermediates (53,54). For both 2 and 3, the bridging nitrogen is expected to be deprotonated under physiological conditions. The alkylaminosulfonylester moiety of 3 is unprecedented and was found to be chemically stable. We chose this group due to the likelihood that the carbonyl carbon of the TC-AMP carbamate would form a tetrahedral sp³-hybridized geometry during attack of N6 of A37 of the tRNA in the carbamoyl transfer reaction catalysed by TsaBDE. Such geometry could be selectively stabilized by TsaBD binding. Compounds 4 and 5 employed malonates as the bridging group between the ribose and threonine and were designed to promote potential Zn binding to the active site of TsaD, via the enol tautomer of the beta-amido ester (4) or amide (5). Final products were purified by HPLC and characterized by NMR and High Resolution MS, which were consistent with the assigned structures (see Supplementary Material).

Kinetic analysis of E. coli t6A reaction using in vitro assays

Before the TC-AMP mimics were tested for inhibition of t6A formation, we carried out an initial kinetic investigation of the overall reaction, as well as the half reaction for TC-AMP conversion to t6A, using a previously described HPLC assay (30). The assay is based on isolation and enzymatic digestion of the modified tRNA product and HPLC quantitation of t6A directly. This assay shows linear reaction rates (Supplementary Figure S6) and allowed us to approximate initial rates for fitting data to the Michaelis-Menten (MM) model (see Table 1). Because of the relatively weak affinity of the TsaBDE ternary complex, protein concentrations of 0.5 μM or above were required for the assays. We are thus reporting apparent kinetic parameters, due to the likely heterogeneity in protein complex formation and its effects on the initial rates for some substrate and inhibitor concentrations. For L-threonine in the overall reaction, fitting the kinetic data to the classical MM model gave an apparent value for \( K_m = 0.59 \pm 0.09 \text{ mM} \) (Figure 4A, Table 1). The kinetic data for ATP as the variable substrate show a good fit to MM kinetics up to a concentration of 1 mM with a value for \( K_m = 60 \pm 6 \text{ μM} \) (Figure 4B). However, severe substrate inhibition was observed for higher ATP concentrations (Figure 4C). This inhibition shows a 50% reduction in the \( V_{max} \) at 5 mM ATP (90% at 10 mM). The physiological relevance of this inhibition is not clear. Although average intracellular ATP concentrations in E. coli are in the millimolar range (55,56), free ATP is unknown. Single cell analysis in E. coli has also shown a wide distribution of intracellular concentrations from 0.1 to 2 mM (57). The substrate inhibition observed is consistent with a similar observation recently reported by Swinehart et al. (58). Kinetic data with tRNA as the variable substrate was somewhat limited by the apparent low values of \( K_m \) for most tRNA transcripts. We were able to measure an apparent \( K_m = 0.5 \pm 0.2 \text{ μM} \) for unmodified tRNA (59) (Figure 4D) by fitting data to the Morrison quadratic equation (50), which is used to account for substrate depletion in reactions in which the tRNA substrate is close to the enzyme concentration.

In addition to measuring t6A formation with the full reaction mixture, we could also isolate the second step of the TC-AMP conversion to t6A, using a previously described assay (4). The substrate inhibition observed is consistent with a similar observation recently reported by Swinehart et al. (58). Kinetic data with tRNA as the variable substrate was somewhat limited by the apparent low values of \( K_m \) for most tRNA transcripts. We were able to measure an apparent \( K_m = 0.5 \pm 0.2 \text{ μM} \) for unmodified tRNA (59) (Figure 4D) by fitting data to the Morrison quadratic equation (50), which is used to account for substrate depletion in reactions in which the tRNA substrate is close to the enzyme concentration.
modification reaction by using HPLC-purified TC-AMP as the substrate with TsaBDE in the presence of ATP (30). Using TsaBDE at a concentration of 0.5 μM, we used the Morrison equation to fit kinetic data for reaction rates with TC-AMP in the concentration range of 0.5–10 μM. Under these conditions, with ATP = 1 mM, we observe an apparent $K_m = 1.5 \pm 0.4$ μM for TC-AMP (Table 1, Figure 4E). Furthermore, with ATP as the variable substrate and [TC-AMP] = 10 μM, we see similar substrate inhibition as observed in the overall reaction (Figure 4F). This suggests that the substrate inhibition observed in the overall reaction is due to ATP interference with the TsaBDE-catalysed transfer of the carbamoyl group to tRNA. Although ATP binding was not observed by ITC with EcTsaBD, likely due to hydrolysis, isosteric ATP analogs have been observed to bind to the heterodimer with low micromolar $K_d$ values (37). By fitting the data to a substrate inhibition model (Equation 3 above), we measured a $K_m = 14$ μM and a $K_{is} = 1.8$ mM for ATP in the second step with TC-AMP as substrate (Table 1). Figure 4G shows more clearly the fit of the data for variable ATP concentrations ≤1 mM. Interestingly, this $K_m$ value is 40-fold lower than the $K_m$ value (0.64 mM) for TsaE-catalyzed ATPase activity by the EcTsaBDE complex (37). Thus, ATP is utilized more efficiently for t6A formation at low ATP concentrations, albeit with a turnover that is ~80-fold lower than the ATPase activity. It is not yet clear what the implications of these data are for the mechanistic role of TsaE in t6A formation or its physiological relevance.

### Inhibition of t6A formation by TCAMP analogs

In testing the TC-AMP mimics for inhibition of the t6A reaction, we first used the full reaction assay to measure IC50 values with concentrations of ATP and L-threonine at 1 mM each. These substrate concentrations were necessary to achieve an optimum rate of t6A formation in our in vitro assay and are close to the estimated physiological concentrations in E. coli (55). The ATP concentration was also chosen to avoid substrate inhibition. The IC50 values were obtained by varying the inhibitor concentration and measuring as close to the initial rates as possible, given limitations on the lower range of enzyme concentration that could be used. The IC50 values for the inhibitors are listed in Table 2. The most potent inhibitor was the phosphonate BK951, which had an IC50 value of 1.3 ± 0.5 μM (Figure 5A). The sulfamate derivative 2 (IC50 = 51 μM) showed 40-fold weaker inhibition under these conditions, as did the aminosulfonysulfamate 3 (IC50 = 45.5 μM). We also prepared the serine analog of TC-AMP, SC-AMP, which is a poor substrate, and gives an IC50 = 9.3 μM (Supplementary Figure S7A–C). This shows the relative importance of the methyl group of the threonine moiety in binding. Both the malonate ester derivative 4 and the malonate amide 5...
Figure 3. Synthesis of phosphonate inhibitor BK951. See Materials and Methods for experimental details.

showed no observable inhibition at concentrations as high as 1 mM (data not shown). The ester analog 4 was also particularly unstable during storage at –20°C or in solution.

Since TC-AMP is a product of TsaC in the first step and a substrate for TsaBDE in the second step of t^6A formation, we tested the analogs as inhibitors for each step of the reaction. For the phosphonate BK951, we found an IC\textsubscript{50} = 7.5 ± 1.2 \mu M for the second reaction step with 10 \mu M TC-AMP and 1 mM ATP (Figure 5B). By assuming competitive inhibition with TC-AMP, the \( K_i \) for BK951 can be estimated using the Cheng–Prusoff equation (59) to give a calculated value of 2.5 \mu M. This \( K_i \) calculation assumes that ATP concentration does not affect inhibitor binding. We did observe that the ATP concentration does have a measurable effect on the IC\textsubscript{50} value for BK951 in the overall reaction (Supplementary Figure S8A). At 0.1 mM ATP, the IC\textsubscript{50} for BK951 is reduced to 0.4 \mu M, while increasing ATP to 5 mM raises the IC\textsubscript{50} of BK951 to 5.6 \mu M. Thus, the \( K_i \) calculation is only an estimate due to ATP competition with TC-AMP in the active site. Similar measurements at 1mM ATP with analogs 2 and 3 gave IC\textsubscript{50} values of 123 and 99 \mu M, respectively, showing that the phosphonate is a better inhibitor of the TsaBDE catalyzed second step in t^6A formation.

To provide an additional, direct measurement of BK951 binding to EcTsaBD, we used a recently described technique, microscale thermophoresis (MST), which measures the difference of thermal motion of fluorescently labelled targets as a function of ligand binding (60). The technique is highly versatile, and can be used to measure protein-protein interactions, as well as small molecule and ion binding to proteins or nucleic acids (61). For these experiments, we used fluorescently labelled EcTsaD, prepared using a probe supplied by the manufacturer that covalently reacts with accessible amines on the protein surface. We found that labelled EcTsaD showed 95% of the activity of the nonlabelled protein (data not shown) in our t^6A assay. In addition, measurement of the affinity of fluorescently tagged EcTsaD to EcTsaB by MST gave a value of 45 ± 6 nM (Figure 5C), which is in good agreement with the previously reported measurement of 120 nM by isothermal calorime-
Figure 4. Kinetics of E. coli t6A formation in the full reaction and isolated second step with purified TC-AMP as substrate. (A) Full reaction mixture with L-threonine as variable substrate fit to MM kinetics (Equation 1). (B) Full reaction mixture with ATP as variable substrate ≤1 mM. (C) Full reaction with ATP as variable substrate showing substrate inhibition at high concentrations (fit to Equation 3). (D) Full reaction with tRNA^{Lys}_{UUU} as the variable substrate fit to Morrison equation (2). (E) Isolated second step of reaction with TC-AMP as variable substrate and 1 mM ATP fit to Morrison equation (Equation 2). (F) Second step of the reaction with TC-AMP (10 μM) as substrate and variable ATP displaying substrate inhibition as in (C). (G) Second step of the reaction with constant TCAMP (10 μM) and ATP as the variable substrate (≤1 mM) fit to MM Equation (1).

We also looked at the effect of high concentrations of TsaE on the binding of Bk951 in the kinetic assay of the overall reaction. We find that high relative concentrations of TsaE (>2.5 μM with the other enzymes at 0.5 μM) strongly inhibit t6A formation (Supplementary Figure S9A). However, this has little effect on Bk951 binding (IC_{50} = 2.4 ± 0.8 μM at 10 μM TsaE, Supplementary Figure S9B). On the other hand, 10 μM TsaE increases the K_{d} of tRNA^{Lys} over 20-fold to 12.0 μM (Supplementary Figure S9C). This latter observation is consistent with the lack of observed tRNA binding by the EcTsaBDE ternary complex (37) and with the rapid loss of tRNA binding upon increasing TsaE to give the TsaB_{2}D_{2}E_{2} complex of T. maritima. These results suggest that TsaE competes with tRNA binding to EcTsaBD but not for Bk951 binding. Because of the differential effects of TsaE on the binding of tRNA and BK951, we also asked if the binding of BK951 affects tRNA binding to EcTsaBD. Preparation of a fluorescently labelled tRNA allowed us to use fluorescence anisotropy to measure the binding of tRNA to the EcTsaBD heterodimer. By titrating a solution of labelled tRNA with increasing con-
centrations of EcTsaBD, either alone or in the presence of 50 μM BK951, we measured $K_d$ values of 87 ± 11 and 112 ± 13 nM, respectively (Supplementary Figure S10). This shows that the presence of bound BK951 has no significant effect on the binding affinity of the tRNA substrate.

Finally, since TC-AMP has been observed as a bound product to TsaC during isolation (34, vide infra), we tested the most potent inhibitors of the overall reaction for the ability to inhibit EcTsaC as product analogs. Table 2 shows that BK951 surprisingly does not bind with significant affinity to this enzyme. This contrasts with the sulfamates 2 and 3, which bind with IC$_{50}$ values of 31 and 97 μM, respectively (Supplementary Figure S7D, E). Thus, the inhibition of the overall reaction by phosphonate BK951 is due solely to its binding to TsaBD, whereas for the sulfamate derivatives, it is a combination of inhibition of both enzymatic steps.

**Structure of BK951 bound to EcTsaBD**

Knowledge of the interactions between the TC-AMP substrate and TsaD is essential for the understanding of the $t^6$A reaction mechanism. We therefore determined the crystal structure of EcTsaBD in complex with our highest affinity inhibitor BK951. Crystallization of EcTsaBD in presence of 2 mM BK951 yielded crystals that diffracted to a resolution of 2.4 Å. The structure was solved by molecular replacement, revealing two identical copies of the EcTsaBD complex in the asymmetric unit. The structure of EcTsaBD is nearly identical to those of apo-form or nucleotide complexes of StTsaBD and EcTsaBD (36,37) (RMSD of about 0.39 Å for 337 superposed C$_\alpha$ positions). The presence of the inhibitor bound to EcTsaD was supported by an extended cloud of residual electron density near the metal sites of TsaD for the two copies in the asymmetric unit (Figure 6A). A complete BK951 molecule could be easily fitted into this density. At full occupancy, the refined B-factors of the BK951 atoms are on average comparable to those of the surrounding side chains. BK951 binds as expected in the active site groove between the N- and C-terminal domains of EcTsaD (Figure 6B). The inhibitor is bound by an extensive network of hydrogen bonding and van der Waals interactions between active site residues, the bound metal ion and the inhibitor (Figure 6C). The purine and ribose engage in identical interactions as those observed for the corresponding moieties of the ATPyS complex with StTsaD (36): the ribose oxygens are H-bonding with the Asp167 carboxy-
Figure 6. Structural characteristics of the BK951–EcTsaBD complex. (A) Stereoview of the residual 2Fo−Fc electron density contoured at 1σ levels for the bound BK951 (green sticks) with metal ion (gray sphere) and surrounding side chains (red sticks). (B) Cartoon presentation of the TsaBD heterodimer bound to BK951; TsaB in blue, TsaD in red, BK951 in green sticks and Zinc as a grey sphere. (C) BK951 (green sticks) bound in the active site of TsaD, Zinc ion as grey sphere. H-bonds and ligand-metal interactions: black dotted lines. (D) LigPlot+ (69) representation of the inhibitor BK951 bound into the active site of EcTsaD (E) Superposition of ATPγS bound to StTsaD (orange sticks) onto BK951 bound to EcTsaD (green sticks). Only residues from EcTsaD are labelled.
late and the N1 of adenine interacts with the Asn272 side chain (Figure 6D). Figure 6E superposes ATPγS (StTsaD) onto BK951, showing that (i) their α-PO₄ moieties superpose very well and (ii) the carbamoyl carbon of BK951 superposes onto the β-P position. The active site Zn coordination sphere is composed of a phosphonate-oxygen and the carboxylate of BK951, His111, His115 and Asp300. The interactions of the threonylcarbamoyl moiety of BK951 are discussed below.

Binding pocket of the L-threonine moiety bound to EcTsaBD

For the first time, the BK951-EcTsaBD structure allows a detailed look at the binding pocket for the threonine moiety. Asp300 binds to the Zn and is conformationally restricted by hydrogen bonding with Ser9, while also making a weak hydrogen bond with the carbamoyl NH of the phosphonate inhibitor. The threonyl side chain specificity for the bound inhibitor appears to be the result of hydrogen bonding to the β-hydroxyl by Ser9 and a pocket formed by van der Waals interactions between the β-methyl group of the inhibitor with His111, Cys10, and a small loop containing residues Pro82, Gly83 and Leu84 (Figure 6C and D). Our kinetic studies show that (±)-3-hydroxynorvaline (hNV) is a good substrate with a Kₘ = 0.5 mM to give the methyl homolog h₄A (Supplementary Figure S11), which has been observed in the tRNA of both bacteria (62) and archaea (63). In vitro substrate activity of hNV with SUA5 and KEOPS has been shown in the archaean system by Perrochia, et al. (31) and also recently shown for E. coli and T. maritima by Swinehart et al. (58). In the structure, the binding pocket for the threonyl side chain has an open face into the ‘tunnel’ formed in TsaD such that extended alky chains may be accommodated if they can project into the tunnel but not if they are restricted in the direction of Cys10. The insertion of a methyl group for one of the β-methyl hydrogens is likely tolerated by projecting the methyl into this open space.

The threonyl carboxylate is both coordinated to the active site Zn atom and also bound by hydrogen bonds from Ser136 and His139. One of the possible strategies to catalyze the reaction would be an activation of the carbamoyl group by the bound metal ion. In the present configuration, the carbamoyl is ideally positioned for nucleophilic attack on the carbonyl oxygen of the TC-AMP carbamoyl group to replace the carboxylate inside the coordination sphere of the Zn. This would allow Lewis acid activation of the carbonyl and facilitate acylation of the N6 of A37 of the tRNA by TC-AMP. The plausibility of such a conformational rotation is suggested by the comparison of the structure of BK951/EcTsaBD and ATPγS/StTsaBD (Figure 6E). The α and β-phosphates of ATPγS superpose onto the αP and carbamoyl carbon of BK951, but the dihedral configuration of the bonds is different. One can easily imagine a rotation of the α–N bonds that leads to an orientation comparable to the P–O configurations of ATPγS to bring the carbamoyl into a catalytic competent orientation. Alternatively, the metal could impose a reactive conformation of the TC-AMP without actively participating in chemical catalysis of the reaction pathway. Indeed, as discussed further, catalytic assistance is offered by active site residues other than the metal ion. By initially binding TC-AMP in a stabilizing conformation the enzyme could protect it from decomposition until the tRNA is bound. Further kinetic studies would be useful to determine if the binding of TC-AMP precedes tRNA binding which would lend support to this hypothesis.

Functional analysis of residues in the TsaD active site

We used site-directed mutagenesis to explore the functional significance of EcTsaD amino acid side chains in the active site binding region of the threonyl moiety of the inhibitor (Figure 7). As in the previously reported structure of EcTsaBD bound to ADP, the Zn atom is coordinated by three universally conserved residues His111, His113 and Asp300 (37). These residues were each mutated to alanine and found to give purified proteins that were inactive in the in vitro t⁶A assay. We also mutated His139, because of its conservation in bacterial TsaD and its position as a Zn ligand in TmTsaDE (41). In the EcTsaBD-ADP structure, His139 is not liganded to the Zn, and in the BK951 bound structure reported here, it is also not chelated to the Zn but it is making a relatively short hydrogen bond to the non-ligated oxygen of the carboxylate of L-threonine (Figure 6C). Surprisingly, both the His139Ala and His139Asn mutants retained activity, with the alanine mutant indistinguishable from wild type. It is possible that a compensatory interaction in the mutants makes up for the loss of this hydrogen bond. In the T. maritima carboxy-AMP bound TsB₂D₂E₂ structure, the analogous His137 is a ligand for the active site Zn ion and found to be required for protein stability. These results likely reflect subtle differences in the organization of the active site in these quite different organisms and may have mechanistic implications.

We also investigated possible specificity determinants for the threonyl moiety of BK951. Interactions between EcTsaD and the threonyl moiety of BK951 are largely made of weak hydrogen bonding interactions (>3.5 Å) with Ser9, His111 and the amide nitrogen of Gly83, as well as van der Waals interactions with the β-carbon of Cys10 and
Modeling A37 of the tRNA substrate into the active site of EcTsaD

Assuming the BK951 adopts the position of the TC-AMP in the active site just before transfer of the threonylcarbamoyl group onto acceptor tRNA, we wondered how the incoming A37 of the latter could be positioned. We therefore modeled an adenine into the active site occupied by BK951 using the Autodock 2 software (64). The protein was kept rigid during the docking process and the distance between the N6 atom of the adenine and the carbamoyl carbon was restrained to 2.5 Å (Supplementary Figure S12). In this model the adenine ring is held between the loops encompassing residues 9–12, 40–44, 137–138 and 176–179. The adenine N9 atom points towards an empty space above the active site, leaving room for an attached tRNA molecule. Because tRNAs that are modified in their stem loops often undergo important conformational changes when bound in the active site of the corresponding enzymes, we did not attempt to model a complete tRNA into the active site of TsaD. Residues Cys10, Asp11, Val42 and Tyr177 are surrounding the adenine. Mutation of Cys10 into Ala causes a more than 50% drop in tA activity. To provide full access of a flipped-out adenine connected to a complete tRNA substrate, the 40–44 loop probably would have to move from its present position in the TsaBD–BK951 complex. Interestingly, this loop was shown to be very flexible and completely disordered in the context of the TmTsaBDE complex (39,40).

Stabilization of TCAMP bound by EcTsaBD and EcTsaC

The inhibitor BK951 is bound to TsaBD in an extended conformation that likely precludes certain decomposition pathways. However, in this structure, the molecule appears to be readily exposed to solvent water. To understand more about the potential stabilizing effect of TsaBD on bound TC-AMP, we carried out a stability study. A mixture of 10 μM purified TC-AMP was incubated with or without 20 μM EcTsaBD at 37°C and timed aliquots were removed for HPLC analysis of TC-AMP conversion to AMP. As shown in Figure 8A, there is only a small protective effect (k<sub>obs</sub> = 6.0 ± 0.2 × 10<sup>−2</sup> versus 7.3 ± 0.2 × 10<sup>−2</sup> min<sup>−1</sup>) at a concentration of EcTsaBD where TC-AMP should be mostly bound. For comparison, the stabilising effect of 13 μM TsaC is nearly an order of magnitude higher (k<sub>obs</sub> = 4.3 ± 0.1 × 10<sup>−3</sup> min<sup>−1</sup>). The rate constants correspond to a t<sub>1/2</sub> = 9.4 min without enzyme, compared to t<sub>1/2</sub> = 11.5 min in the presence of TsaBD and t<sub>1/2</sub> = 160 min in the presence of TsaC. These data suggest that either TC-AMP is transferred to tRNA rapidly after binding to TsaBD, or additional binding events involving TsaC, or tRNA may help to protect bound TC-AMP from hydrolysis before it is transferred by shielding it from solvent water.

For the TsaC experiments, we utilized the TC-AMP bound form of EcTsaC that can be isolated from the purification of highly overexpressed protein. We find that the amount of TC-AMP bound to TsaC can be quantified directly by HPLC and typically approaches 70% occupancy after affinity and size exclusion chromatography (Supplementary Figure S13A). Bound TC-AMP can also be converted to ATP quantitatively by the addition of 1.5 equivalents of pyrophosphate in the presence of Mg ion (Supplementary Figure S13B) and subsequently removed by size exclusion chromatography to provide the unliganded form of the enzyme (Supplementary Figure S13C). Further investigation of TsaC stabilization of TC-AMP in these complexes shows a significant reduction in protection as the concentration of the complex is lowered (Figure 8B). This concentration dependence is likely due to an increased fraction of unbound TC-AMP. A plot of the t<sub>1/2</sub> values for TC-AMP stability versus TC-AMP concentration gives a crude estimated half-maximal value of ~5 μM (data not shown), which may reflect a similar magnitude for the K<sub>d</sub> of TC-AMP binding to EcTsaC. If this were true, it would suggest that unless the concentration of TsaC or TC-AMP in E. coli...
is greater than 5 μM, the majority of TsaC in the cell will be free of bound TC-AMP.

DISCUSSION

The phosphonate BK951 is a potent inhibitor of \( ^6 \text{A} \) biosynthesis in vitro

The phosphonate BK951 was the most potent analog in this study, with an IC\(_{50}\) value of 1.3 μM for the overall \( ^6 \text{A} \) reaction. We measured inhibitor IC\(_{50}\) values in the presence of L-threonine and ATP concentrations of 1 mM each. For \textit{E. coli}, 1 mM ATP is in the lower part of the physiological range of up to 10 mM. The intracellular L-threonine concentration has been recently reported to be 1.3 mM (55).

Substitution of the phosphonate with a sulfamoyl group in 2 results in a loss of inhibition in the kinetic assay by 40-fold, which is possibly due to the weaker binding of the sulfonyl oxygen with Zn compared to the fully ionized phosphonate. Further substitution of the carbamoyl group of TC-AMP with a tetrahedral sulfonyl gives an alkylaminosulfonyl-N-sulfamate 3, which is an intended transition state mimic of the attack by N6 of A37. The unique pharmacophore structure of this inhibitor has surprisingly not been previously described in the literature. Similar to a few known alkylaminosulfonamides, this structure likely forms an ‘inner salt’, with the bridging nitrogen deprotonated and the amine of the L-threonine protonated to give a zwitterion structure. Despite the novelty of its structure, compound 3 has virtually the same IC\(_{50}\) as sulfamate analog 2. Since the carbonyl oxygen of BK951 is making only one hydrogen bond in the EcTsaBD bound structure, there does not appear to be amino acid side chains in place to distinguish the sp\(^3\) orientation of the sulfonyl group in this area of the active site in this conformation of the protein. Finally, both the malonate ester and the malonate amide bridge between the adenosine ribose and threonine (4 and 5, Figure 2) failed to give molecules that showed any observable binding to EcTsaBD. Both molecules were designed to examine the hypothesis that the Zn could chelate the phosphonate and the carbamate carbonyl of TC-AMP, which in this structure is not observed. Hence, the lack of observed binding of the malonates supports the overall significance of the current structure.

If we assume the affinity of TC-AMP to EcTsaBD is similar in magnitude to the \( K_m \) value of 1.5 μM, the binding data for the phosphonate analog suggests it is a good mimic that is taking advantage of most of the protein binding interactions. This conclusion is supported by the similarity of the conformation of atoms of the adenosyl \( 5' \)-phosphate/phosphonate group in the active site of TsaD of both BK951 and ATP. In the recently modelled binding of TC-AMP to TmTsaD based on the structure of bound carboxy-AMP, Luthra \textit{et al.} proposed a similar binding of the threonine carboxylate moiety to the active site Zn (41). Their model differs from our structure in the lack of chelation between the phosphonate and the Zn. It is possible that there are species-specific differences in the orientation of some amino acids, as well as rearrangements of the substrates along the reaction coordinate. Thus, many of the various structures may each be relevant for a distinct part of the transformation.

The sulfamate analogs are better mimics of TC-AMP binding to TsaC

Although BK951 is more potent in inhibiting the second step of \( ^6 \text{A} \) synthesis, it surprisingly shows no inhibition of the first step of the reaction catalysed by TsaC. The sulfamates 2 and 3, on the other hand, show significant inhibition of both enzymatic steps in \( ^6 \text{A} \) formation. A reexamination of the crystal structure of TC-AMP bound to TsaC2 from \textit{S. tokodaii} (34) shows a hydrogen bond between the backbone amide of conserved Ser144 to the adenylate ester oxygen of TC-AMP that is replaced by a methylene in BK951. There are also close van der Waals interactions likely between the backbone carbonyl and alpha carbons of Pro143 (usually a Thr in TsaC1). A possible steric clash between these carbon atoms and the methylene of the phosphonate analog, in addition to the loss of hydrogen bond would be sufficient to explain its lack of binding to TsaC. Conversely, the amide nitrogen of the sulfamate group in analogs 2 and 3 is acidic enough to be deprotonated, which makes it an excellent mimic of the adenylate ester oxygen as previously documented (53, 54). Therefore, TsaC appears to use these interactions to closely select for an ester at this position and analogs that preserve the integrity of these interactions will be generally more potent as inhibitors.

The binding conformation of BK951 suggests a precatalytic complex

The binding interactions between BK951 and the active site of the TsaD protein suggests that at one point in the reaction coordinate, the Zn interacts directly with the phosphonate oxygen and carboxylate oxygens of TC-AMP. The carbamoyl oxygen does not form a ligand to the metal ion, but establishes two H-bonds with mainchain NH groups of Ser\(^{136}\)Gly\(^{138}\), the \( \beta \)-hairpin connection between strands \( \beta\)6–\( \beta\)7. This is a similar binding mode as the carbamoyl adenylate bound to the homologous TobZ protein, which catalyses the carbamoylation of tobramycin (34). This position of the carbonyl suggests that the amide main chain NH groups of Ser\(^{136}\)Gly\(^{138}\) could provide an oxyanion hole during the nucleophilic attack of the A37 6-amino group during the transfer reaction. As shown in Figure 9, this loop is extremely well conserved in the TsaD/Kae1/Qri7 family, suggesting its importance for catalytic activity. This loop is also present in the related carbamoylating enzymes TobZ and HypF, reinforcing the hypothesis of its role in the stabilization of the intermediate. This means that the main role of the metal is to provide a scaffold for productive binding of the substrate. In the TobZ active site, a histidine is positioned near the carbamoyl group that could act as a general base to subtract a proton from the incoming tobramycin OH nucleophile. This is not the case for TsaD, probably because the A37 NH\(_2\) nucleophile is deprotonated. Alternatively, the carbamoyl group of the TC-AMP could slightly rotate and engage in a Lewis acid interaction with the metal to increase its reactivity. The latter scenario seems less likely to us because it would render the carbon less accessible for an incoming nucleophile. Clearly, structural data in presence of the tRNA substrate are needed for a better understanding of the mechanism. Assuming that the binding mode of the BK951 carbonyl mimics that of the TC-AMP.
Figure 9. Multiple sequence alignment of the Tsad/Kael/OSGEP family members was realized using the blosum 45 with MAFFT (online version 7) (70). Structure-based alignment was performed using the ESPRIT web-server (71). The SGG peptide conserved in all homologs is highlighted in pink.
substrate, its access by the bulky tRNA substrate would likely be hampered by some of the surrounding active site loops and helices, especially the region contained between residues 32 and 45. Interestingly this region becomes completely disordered upon TsaE binding (40), suggesting that flexibility plays a major role in the catalytic cycle.

The ability of EcTsaBD and EcTsaC to stabilize bound TC-AMP differ significantly

Previous work showed that TC-AMP is unstable with a half-life of 3.5 minutes at pH 7.5 in the presence of 2 mM Mg ion (30). Both the instability and the magnitude of the $K_m$ value for TC-AMP (1.5 µM) for the isolated TsaBDE-catalysed $t^6$A reaction are similar in magnitude for the structurally analogous primary metabolite carbamoyl phosphate (CP). The $K_m$ values for CP for the enzymes aspartate and ornithine transcarbamoylases are ~4–10 µM. Wang et al. (65) have shown that CP is stabilized by binding to these enzymes in the absence of the other substrates, and the effect is more pronounced for hyperthermophiles to the point of requiring substrate channelling (66). Such binding prevents the major mechanism of degradation of CP which involves intramolecular transfer of the carbamoyl NH proton to the phosphate and subsequent rapid formation of cyanate. In our previous analysis of TC-AMP decomposition, NMR data showed L-threonine and the cyclic oxazolidinone as the only two detectable final products produced in an equimolar ratio (30). The oxazolidinone forms via attack by the β-hydroxyl oxygen on the carbamoyl carbonyl carbon (or isocyanate intermediate) to release AMP and give the cyclic product. The structure reported here suggests that TC-AMP is bound by TsaBD in an extended conformation which would minimize the cyclization pathway. Although we did find that EcTsaBD stabilizes TC-AMP in solution, the effect is small. This is likely due to TC-AMP remaining accessible to solvent water when bound to TsaBD, which allows for direct hydrolysis. The fact that the half-life of TC-AMP bound to EcTsaBD is more similar to free TC-AMP in the absence of metal ion (16 min) than in its presence (3.5 min) is consistent with the conformation of bound BK951, in which the carbamoyl carbonyl is not chelated to the Zn. Retaining stability of TC-AMP may be another reason for avoiding such coordination. In other words, if the carbamoyl carbonyl were coordinated to the Zn, the enzyme could catalyze wasteful transfer of the carbamoyl group to water.

In contrast to EcTsaBD, we find an order of magnitude greater stabilizing effect on TC-AMP by binding to EcTsaC. Previous work has shown that overproduced EcTsaC retains bound TC-AMP after purification (30) and that crystallized TsaC2 from S. tokodaii (45) contained bound TC-AMP (34). In this structure, the threonylcarbamoyl group of TC-AMP is sequestered from solvent in a long, narrow binding pocket. The observed greater stabilization by TsaC may be used to prevent wasteful hydrolysis of TC-AMP in the cell (34). However, we find that as the concentration of EcTsaC is lowered into the low micromolar range, the stabilizing effect of the enzyme on TC-AMP is diminished significantly. It thus may be less of an imperative for mesophilic organisms to stabilize this metabolite which could be a relic of enforced stabilization in hyperthermophilic organisms, considering the ancient nature of this enzyme system (34).

In summary, we report here the synthesis, biochemical characterization and cocrystal structure of a potent inhibitor of bacterial $t^6$A formation. The inhibitor appears to bind in the active site of the heterodimeric TsaBD complex in a manner similar to the natural substrate TC-AMP and could therefore be useful as a mechanistic probe for the reaction in a variety of biological contexts. The crystal structure should be useful for the rational design of more potent, cell permeable inhibitors of $t^6$A modification that may find application as biochemical probes in a variety of organisms given the universal importance of this modification. Inhibitors optimized for the bacterial enzymes may find use as potentially novel antibiotics.

DATA AVAILABILITY

Atomic coordinates and structure factors for the reported crystal structures have been deposited with the Protein Data bank under accession number 6Z81.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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