**Chlamydia trachomatis CT771 (nudH) is an asymmetric Ap₄A hydrolase**

Michael L. Barta¹, Scott Lovell², Amy N. Sinclair¹, Kevin P. Battaile³, and P. Scott Hefty¹,*

¹Department of Molecular Biosciences, University of Kansas, Lawrence, KS 66045, USA
²Protein Structure Laboratory, Del Shankel Structural Biology Center, University of Kansas, KS 66047, USA
³IMCA-CAT, Hauptman-Woodward Medical Research Institute, Argonne, IL 60439

**Abstract**

Asymmetric diadenosine 5′,5′″-P¹,P⁴-tetraphosphate (Ap₄A) hydrolases are members of the Nudix superfamily that asymmetrically cleave the metabolite Ap₄A into ATP and AMP while facilitating homeostasis. The obligate intracellular mammalian pathogen *Chlamydia trachomatis* possesses a single Nudix family protein, CT771. As pathogens that rely on a host for replication and dissemination typically have one or zero Nudix family proteins, this suggests that CT771 could be critical for chlamydial biology and pathogenesis. We identified orthologs to CT771 within environmental *Chlamydiales* that share active site residues suggesting a common function.

Crystal structures of both apo- and ligand-bound CT771 were determined to 2.6 Å and 1.9 Å resolution, respectively. The structure of CT771 shows a αβα-sandwich motif with many conserved elements lining the putative Nudix active site. Numerous aspects of the ligand-bound CT771 structure mirror those observed in the ligand-bound structure of the Ap₄A hydrolase from *Caenorhabditis elegans*. These structures represent only the second Ap₄A hydrolase enzyme member determined from eubacteria and suggest that mammalian and bacterial Ap₄A hydrolases might be more similar than previously thought. The aforementioned structural similarities, in tandem with molecular docking, guided the enzymatic characterization of CT771. Together, these studies provide the molecular details for substrate binding and specificity, supporting the analysis that CT771 is an Ap₄A hydrolase (nudH).

Found throughout all kingdoms of life, Nudix hydrolases are a superfamily of metal-dependent (typically Mg²⁺) enzymes that catalyze the cleavage of nucleoside diphosphates linked to any other moiety X (Nudix)¹. These proteins serve to maintain physiological homeostasis by modulating levels of signaling molecules and potentially toxic metabolic intermediates². Enzymes of this family are characterized by a structurally conserved metal-and-substrate binding catalytic loop-helix-loop ‘Nudix’ motif (comprising 23 amino acids, GX₃EX₃REUXXEEXGU where U is preferably a hydrophobic residue), which forms one edge...

---

¹To whom correspondence should be addressed: 1200 Sunnyside Avenue, University of Kansas, Lawrence, KS 66045; Phone: 785-864-5392; Fax: 785-864-5294; pshefty@ku.edu.

The atomic coordinates and structure factors (codes 4ILQ and 4MPO) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

**SUPPORTING INFORMATION** Molecular Docking of CT771 with Nudix Substrates, Bound Sulfate in Apo-CT771 Structure and Structural Superposition of Each Chain Within Ligand-Bound CT771 Results sections. Electrostatic surface potential of free and bound CT771 structures (Fig. S1). Interactions Between CT771 and a Bound Sulfate Anion (Figure S2). Structural comparison of bound CT771 and other Ap₄A hydrolase complexes (Figure S3). Structural superposition of each chain within ligand-bound CT771 (Figure S4). Disruption of intrachain disulfide bond upon substrate binding (Figure S5). Bonding distances between CT771 and bound ligands (Table S1). This material is available free of charge via the Internet at http://pubs.acs.org.
of the putative active site. Despite a diverse range of substrates, a conserved $\alpha$-$\beta$-$\alpha$-sandwich scaffold is present within all structurally studied Nudix hydrolases. Substrate specificity is thus determined by side chains outside the Nudix motif, which further subdivides this superfamily, and includes dinucleotide polyphosphatases (e.g. diadenosine tetraphosphate (Ap$_4$A) hydrolases), nucleotide sugar pyrophosphatases, ribo- and deoxyribonucleotide triphosphatases, among others.

Ap$_4$A hydrolase subfamily members are found throughout all kingdoms of life with previous phylogenetic analyses suggesting the subdivision of plant/bacterial and mammalian/archeal enzymes into two distinct classes. Studies detailing structural insights of substrate binding and enzyme mechanics have been revealed within plant, bacterial, and mammalian enzymes. Asymmetric cleavage of the substrate, Ap$_4$A, results in the generation of ATP and AMP (EC 3.6.1.17). While the biological role of Ap$_4$A is not completely understood, it is found at micromolar intracellular concentrations in both eukaryotes and eubacteria and typically accumulates as a result of protein synthesis when aminocyl-tRNA synthetases are charged with their cognate amino acid. Ap$_4$A levels have also been implicated in a variety of signaling pathways including apoptosis, DNA repair, pathogenesis, and as a link between transcription and protein synthesis, among others.

*Chlamydia trachomatis* is an obligate intracellular bacterium that replicates within a parasitophorous vacuole, termed the inclusion. A distinguishing feature of *Chlamydia* is its virulence-defining developmental cycle that involves an infectious, metabolically inactive form (elementary body or EB) and noninfectious, replicative form (reticulate body or RB). After host cell entry and formation of the inclusion by an EB, a burst of nascent bacterial metabolism is initiated that coincides with the EB-to-RB morphological transition. As *Chlamydia* have coevolved with their human host, reductive evolution has resulted in the loss of numerous genes involved in substrate and oxidative phosphorylation, resulting in a limited ability to manufacture ATP and an energetically parasite-like existence. Thus, the ability to salvage and utilize nucleotide intermediates (such as Ap$_4$A) within the inclusion, whether chlamydial-generated or imported from the host cell, is likely crucial to basic bacterial biology as well as pathogenesis. BLAST analysis of the *Chlamydia trachomatis* genome reveals the presence of a single ORF containing a Nudix motif, CT771 (*Chlamydia trachomatis* ORF 771), which is annotated as a 17.4 kDa MutT/Nudix family protein and is conserved across all sequenced *Chlamydiaceae* members. Typically, pathogens that rely on a host for replication and dissemination (e.g. *Rickettsia*, *Borrelia*, mycoplasmas, etc.) lack or have a single Nudix family enzyme, suggesting the singular presence of a Nudix protein could be of critical importance to *Chlamydia*. In order to further characterize CT771 as a Nudix subfamily member, a crystal structure of CT771 was determined to 2.60 Å. Structural homology with Ap$_4$A hydrolase members as well as examination of the putative active site through molecular docking guided the subsequent enzymatic characterization of CT771. Finally, a co-crystal structure of CT771 bound to products of Ap$_4$A hydrolysis was determined to 1.9 Å, which ultimately provided the mechanistic details of substrate binding and specificity for CT771, as well as confirmed its functional role as an Ap$_4$A hydrolase (*nudH*).

### MATERIALS AND METHODS

#### Cloning, Overexpression, and Purification of Recombinant CT771

A gene fragment encoding the entire open reading frame (residues 1–150) of CT771 was amplified from *C. trachomatis* (serovar 434/Bu) genomic DNA via PCR and subcloned into *BamHI*-digested expression plasmid pTBSG through ligation independent cloning. Upon DNA sequence confirmation, the vector was transformed into BL21 (DE3) *E. coli* cells.
strain was grown to an OD\textsubscript{600} of 0.8 at 37°C in Terrific Broth supplemented with Ampicillin (100 µg/ml), and protein expression was induced overnight at 17 °C by the addition of isopropyl 1-thio-β-D-galactopyranoside (IPTG) to a 1 mM final concentration. Bacterial cells were harvested by centrifugation, resuspended in lysis buffer (20 mM Tris-HCl (pH 8.0), 500 mM NaCl, and 10 mM imidazole), and then lysed by microfluidization. The soluble tagged protein was collected in the supernatant following centrifugation of the cell homogenate and purified on a Ni\textsuperscript{2+}-NTA-Sepharose column according to published protocols \textsuperscript{18}. Recombinant tobacco etch virus (TEV) protease was used to digest the fusion affinity tag from the target protein. After desalting into 20 mM Tris-HCl (pH 8.0), final purification was achieved by ResourceQ anion-exchange and gel filtration chromatography (GE Healthcare). The purified protein was concentrated to 10 mg/ml and buffer exchanged by ultrafiltration into 10 mM Tris-HCl (pH 7.5), 50 mM NaCl, and stored at 4 °C for further use.

**Crystallization**

Recombinant \textit{C. trachomatis} CT771 was crystallized by vapor diffusion in Compact Jr. (Emerald Biosystems) sitting drop plates at 20 °C. Specifically, 0.5 µl of protein solution (10 mg/ml in 10 mM Tris-HCl (pH 7.5), 50 mM NaCl) was mixed with 0.5 µl of reservoir solution containing 100 mM Tris-HCl (pH 8.5) and 1.5 M ammonium sulfate, from the Salt Rx HT screen condition F3 (Hampton Research), and equilibrated against 75 µl of the reservoir solution. Single bipyramidal-shaped crystals appeared after 1 day and continued to grow for ~3 days. Crystals were flash-cooled through serial dilution in a cryoprotectant solution consisting of 2.0 M ammonium sulfate with an increasing concentration of glycerol (5%, 10%, 20% (v/v) final).

Prior to cocrystallization, CT771 (10 mg/ml in 10 mM Tris-HCl (pH 7.5), 50 mM NaCl) was incubated on ice with 10 mM Ap\textsubscript{4}A (Sigma, reconstituted in 10 mM MgCl\textsubscript{2}) for 15 minutes. Upon mixing 0.5 µL of the protein/Ap\textsubscript{4}A solution with 0.5 µL reservoir solution containing 200 mM MgCl\textsubscript{2}, 100 mM Tris-HCl (pH 8.5) and 25% (w/v) PEG 3350, from the Index HT screen condition H1 (Hampton Research), single block shaped crystals appeared after 1 day and continued to grow in size for up to 5 days. Crystals were harvested by flash-cooling in a cryoprotectant solution consisting of 80% mother liquor and 20% (v/v) glycerol and prepared for X-ray diffraction.

**Diffraction Data Collection, Structure Determination, Refinement and Analysis**

Monochromatic X-ray diffraction data were collected at 100K using a Dectris Pilatus 6M pixel array detector at beamline 17ID at the APS IMCA-CAT (Table I). Individual reflections were integrated with XDS \textsuperscript{19} and scaled with Aimless \textsuperscript{20}, which suggested the Laue class was 6/mmm with a likely space group of \textit{P}\textsubscript{6}\textsubscript{1}2\textsubscript{2} or \textit{P}\textsubscript{6}\textsubscript{5}2\textsubscript{2} for the apo-CT771 crystals and \textit{C}\textsubscript{2} for the CT771/AMP-PO\textsubscript{4} cocrystals.

Initial phase information was obtained for the apo-CT771 structure by maximum-likelihood molecular replacement using BALBES \textsuperscript{21}. Both of the likely space groups were analyzed, with only \textit{P}\textsubscript{6}\textsubscript{5}2\textsubscript{2} producing a potential solution. Specifically, 134 residues of chain B from PDB entry 3I7V (\textit{Aquifex aeolicus} Ap\textsubscript{4}A hydrolase) were altered to reflect the sequence of CT771 (residues 13 – 148), and the resulting hypothetical structure was used as a search model. The top solution contained a single copy of CT771 in the asymmetric unit, which corresponded to a Matthew’s coefficient \textsuperscript{22} of 4.66 Å\textsuperscript{3}/Da and a solvent content of 73.6%. The final refined CT771 structure was used as a search model for the CT771/AMP-PO\textsubscript{4} dataset, and the top solution contained eight copies of CT771 in the asymmetric unit with a Matthew’s coefficient of 2.35 Å\textsuperscript{3}/Da and a solvent content of 47.7%.
Structure refinement for the apo-CT771 structure was carried out using Phenix. One round of individual coordinates and isotropic atomic displacement factor refinement was conducted, and the refined model was used to calculate both $2F_o-F_c$ and $F_o-F_c$ difference maps. These maps were used to iteratively improve the model by manual building with Coot followed by subsequent refinement cycles. TLS refinement was incorporated in the final stages to model anisotropic atomic displacement parameters. Hydrogen atoms were included during all rounds of refinement. Ordered solvent molecules were added according to the default criteria of Phenix and inspected manually using FEM electron density maps (contoured at 1.0 $\sigma$) within Coot prior to model completion. Residues 1-3 were not modeled as a result of poor map quality.

The CT771/AMP-PO$_4$ cocrystal structure was refined in a similar manner as described above. Regions of poor map quality with the CT771/AMP-PO$_4$ cocrystal structure prevented modeling of residues 1-4 and 149-150 in all chains (with the exception of chain B, where residues 1-4 were modeled and chain C, where residue 4 was modeled). The following residues were not modeled: chain F, 147-148; chain G, 18-21, 84-88. Disordered side chains were truncated to the point where electron density could be observed. Additional information and refinement statistics are presented in Table 1.

Kinetic Characterization of CT771

A luciferase-based bioluminescence assay was used to characterize the kinetic constants of wild-type CT771 enzyme via the detection of liberated ATP. Each 100 $\mu$L assay mixture was comprised of 100 mM Tris-HCl (pH 7.5), 1.5 – 5.0 mM MgCl$_2$ or MnCl$_2$, 10 $\mu$L of reconstituted rL/L Reagent (Promega), 10 $\mu$L of Ap$_4$A at varying concentrations (Sigma) and 10 ng of purified CT771, and was incubated at room temperature for 5 minutes before measurements were taken. Light output was measured on a Biotek Synergy 2 luminometer and converted into units of enzyme activity ($\mu$M/min) using an ATP standard calibration curve. All reactions were measured in triplicate and data interpretation was carried out using GraphPad (GraphPad Software). Additional information is presented in Table 4.

Ligand Docking

A model for diadenosine tetraphosphate (Ap$_4$A), 8-deoxy-dGTP, adenosine diphosphate ribose and GDP-mannose were generated using the PRODRG server. Polar hydrogens were added to ligands prior to docking. Docking calculations were carried out with the refined crystal structure of CT771 using AutoDock Vina, a comprehensive program for molecular docking and virtual screening. As grid maps are calculated automatically in AutoDock Vina, only the size and location of search space is required. The grid box was set to 60 x 60 x 60 Å with grid points spaced every 0.375 Å and centered within the putative Nudix family active site cleft. Docking simulations were performed at physiological conditions (37°C). Theoretical dissociation constants were calculated for the lowest energy site using the following equation: $\Delta G = RT*ln(K_d)$. Only ligands docked within the putative active site cleft were analyzed. Additional information is presented in Table 3.

Multiple Sequence Alignments and Figure Modeling

Multiple sequence alignments were carried out using ClustalW and aligned with secondary structure elements using ESPRIPT. The Ap$_4$A hydrolase sequences used in alignments, along with their respective GenBank accession numbers, are as follows: C. trachomatis serovar 434/Bu (166154113); Parachlamydia acanthamoebae (282890520); Candidatus Protochlamydia (46446871); Simkania negevensis (338733601); Waddlia chondrophila (297621593); Homo sapiens (4502125); Caenorhabditis elegans (17509897); Aquifex aeolicus (15605731). Three-dimensional structures were superimposed using the Local-Global Alignment method (LGA). The Ap$_4$A hydrolase structures obtained from
the PDB are as follows: Homo sapiens (3U53); Caenorhabditis elegans (1KTG); Aquifex aeolicus (3I7V). Representations of all structures were generated using PyMol. Calculations of electrostatic potentials at the molecular surface were carried out using DELPHI.

RESULTS

CT771 Crystal Structure

The structure of CT771 was solved by molecular replacement and refined to 2.60 Å (Table 1). CT771 is characterized by an αβα-sandwich fold (Fig. 1) with dimensions approximately 50 × 25 × 30 Å, typical of Nudix family proteins, with 3 α-helices (residues Pro52-Thr64, Phe120-Arg125 and Arg133-Tyr146, respectively) surrounding 7 β-sheets (residues Lys5-Phe18, Ser24-His33, Lys37-Gly40, Gly67-Phe71, Phe76-Asn83, Phe89-Lys102 and Cys113-Leu118, respectively). The overall topology of CT771 is β1-β2-β3-β4-α1-β5-β6-β7-α2-α3 (Fig. 1). The conserved Nudix motif (GXGE-X-XGL, residues 44-66) is found within α1 and the loop regions surrounding this helix (represented as ball-and-sticks and colored magenta in Fig. 1, right panel). The protein interfaces and assemblies (PISA) server was used to assess potential modes of oligomerization. This analysis failed to identify any interfaces that were judged to be thermodynamically favorable, consistent with the monomeric elution profile of CT771 in analytical gel filtration experiments (data not shown).

Structural Comparison with Ap4A Hydrolases

In order to better understand the function of CT771, the refined crystal structure was used to search for structurally-related enzymes within the PDB via the DALI server, with the top 5 hits listed in Table 2. Not surprisingly, over 100 of the highest scoring matches are enzymes from the Nudix family. Structures displaying the highest homology to CT771 are diadenosine tetraphosphate (adenosine-P1-P2-P3-P4-adenosine or Ap4A) hydrolases. Furthermore, both mammalian and eubacterial enzymes are found within the top 5 hits, suggesting that they are structurally more alike than previously thought. Indeed, structural alignment of CT771 and the top 3 DALI hits, Ap4A hydrolases from H. sapiens, C. elegans and A. aeolicus, (Fig. 2A) further underscores the conserved nature of the αβα-sandwich fold found within this class of enzymes. All 4 structures align with RMSD values better than 2.2 Å, with large-scale variations limited to the loops connecting β1-β2 and β5-β6. Neither of these regions are expected to contribute to substrate binding or catalysis and likely reflect crystallographic differences stabilized by packing.

Kinetic Characterization of Ap4A Hydrolase Activity of CT771

In order to better understand the function of CT771 and provide support for the structural similarities to Ap4A hydrolases revealed by both the DALI search and molecular docking (described in Supporting Information), purified protein was enzymatically characterized using a Luciferase assay as described in the Materials and Methods section. This assay allows the continuous monitoring of cleaved Ap4A via released ATP upon substrate hydrolysis, and has been extensively used to characterize this specific Nudix enzyme class. The results of this analysis support the conclusion that CT771 is a bona fide Ap4A hydrolase (Table 4). Intriguingly, the largest observed kinetic activity was in the presence of 5 mM Mg2+ with \( k_{\text{cat}} = 3.2 \pm 0.1 \) (s\(^{-1}\)), while the lowest observed binding affinity was in the presence of 5 mM Mn2+ with a \( K_m = 5.0 \pm 0.5 \) μM (Fig. 3). \( K_m \) measurements assumed that the rate of product formation was slower than enzyme-substrate...
dissociation. Despite its ability to increase substrate affinity and enzyme efficiency ($k_{\text{cat}}/K_m$), increasing concentrations of Mn$^{2+}$ had an adverse affect on the catalytic rate of CT771 in the presence of Ap$_4$A. These values are in agreement with previously reported values, using a bioluminescent assay, for *H. sapiens* Ap$_4$A, among others.  

**Structural Basis for Substrate Recognition of CT771**

Crystals of CT771 grown in the presence of Ap$_4$A diffracted X-rays to 1.90 Å resolution in the space group C2 (Table 1). Upon modeling and refinement of eight polypeptides in the asymmetric unit, two regions of strong contiguous electron density (visible when contoured up to 5.0 $\sigma$) were apparent within each putative active site (Fig. 4A). While the electron density was clearly not large enough to accommodate an entire Ap$_4$A molecule (Fig. 4B), an AMP moiety was modeled near the loop region connecting $\alpha_2$ and $\alpha_3$ (Fig. 4C). This phosphate will be designated the P$_1$-phosphate in further discussions, in accordance with Guranowski *et al.*[39], as it is proximal to the tighter binding adenosine group within Ap$_4$A. Additionally, a large cluster of electron density was apparent near $\alpha_1$, where a single phosphate, three magnesium ions, and a network of waters were modeled (Fig. 4D). As wild-type CT771 was used in the crystallization experiments with Ap$_4$A, it was not unexpected for the substrate to undergo hydrolysis. Likewise, previous structural studies on *C. elegans* Ap$_4$A hydrolase[37] with the substrate analog AppCH$_2$ppA yielded nearly the same interpretable regions of electron density as described here (Fig. S3B). The authors speculated that AppCH$_2$ppA had either been catalytically processed by the hydrolase during crystallization or that flexible regions within the substrate (e.g. P$_2$- and P$_3$-phosphates and the second adenosine group) subsequently lacked interpretable electron density. Given the extreme similarities in bound active site ligands between those studies and ours, despite the use of different substrate molecules, we speculate that both compounds have been processed by the crystallized hydrolase in each structure, resulting in a bound AMP molecule and metal-phosphate complex.

Analysis of the bound AMP molecule within each CT771/AMP-PO$_4$ subunit active site revealed that it resides in a deep groove (~11 Å as measured from the Tyr96 hydroxyl to the N3 atom of the bound AMP molecule) created by the loop regions connecting $\beta_2$-$\beta_3$ and $\beta_5$-$\beta_6$, as well as the $\beta$ sheets themselves (Fig. 5). The adenine ring of AMP adopts the *anti* conformation, where it is stabilized by a hydrogen bond interaction with the carboxylic acid moiety of Glu131 (Fig. 4C). Additional protein contacts with the adenine ring involve an extensive set of $\pi$-$\pi$ stacking interactions created by the aromatic side chains Phe129 and Tyr80. All three side chains are highly conserved across examined Ap$_4$A hydrolases (red circles, Fig. 2B), suggesting a common mechanism of substrate binding and orientation. The ribose sugar hydroxyl groups of the bound AMP molecule adopt the C2$'$-end and C3$'$-exo conformations, and the P$_1$-phosphate is oriented towards the catalytic $\alpha_1$ Nudix helix (Fig. 4C). While the ribose sugar moiety lacks any hydrogen bonding with CT771, the P$_1$-phosphate is stabilized by an extensive network of positively charged side chains (His38, Lys43 and Lys92), along with Tyr80 (Fig. 4C, S1C). As with the adenosine moiety, all four of these residues interacting with the P$_1$-phosphate are highly conserved, with both Lysine residues being invariant (red and blue circles, Fig. 2B). While bond catalysis occurs at the P$_3$.P$_4$-phosphate interface[39, 40], proper orientation of Ap$_4$A clearly begins in this ~11 Å-deep groove.

Consistent with the previously reported Ap$_4$A hydrolase binary complex from *C. elegans*[37], the region of strong electron density near the catalytic $\alpha_1$ Nudix helix (Fig. 4A) was modeled with a single phosphate anion bound to a network of magnesium ions and waters (Fig. 4B,D). Bailey *et al.* proposed that this phosphate molecule occupied the P$_4$-phosphate

*Biochemistry. Author manuscript; available in PMC 2015 January 14.*
position of Ap₄A, primarily due to its proximity to the Nudix helix and distance from the P¹-phosphate. Thus, all further discussions of this site will refer to it as the P⁴-phosphate.

Within the CT771/AMP-PO₄ structure the P⁴-phosphate is maintained in a tightly bound position through numerous solvent-ligand contacts and a single direct protein interaction (His45). Solvent-ligand contacts are primarily mediated by three hexacoordinated Mg²⁺ ions, which themselves form tight metal coordination bonds (average bond length, 2.13 Å) with three Glutamate residues (Glu59, Glu63 and Glu111), 6 water molecules and the aforementioned phosphate (Fig. 4D). A fourth Glutamate residue (Glu62) forms hydrogen bonds with two of the active site waters involved in magnesium hexacoordination. Arg58 appears to stabilize the position of Glu59, which coordinates with two of the three Mg²⁺ ions. As these residues are found within the Nudix motif, it is unsurprising that they are all highly conserved amongst Ap₄A hydrolases (red circles, Fig. 2B). Intriguingly, only three of the oxygens within the P⁴-phosphate participate in extensive contacts, suggesting the fourth oxygen would likely be linked to the second adenosyl moiety of Ap₄A. Thus, as the CT771/AMP-PO₄ structure was crystallized in the presence of a high concentration of magnesium (200 mM MgCl₂), the active site metals appear to properly maintain the P⁴-phosphate in a highly specific orientation, priming the substrate for subsequent nucleophilic attack and resulting catalysis at the P³-P⁴-phosphate bond. While substrates other than Ap₄A were not tested for enzymatic activity in the current study, it has previously been reported that this family of enzymes can hydrolyze adenosine polyphosphates with four, five or six phosphates, but not three, indicating positioning of the P⁴-phosphate is critical to substrate catalysis.

Despite efforts to co-crystallize Ap₄A with CT771, it appears that the substrate was hydrolyzed prior to data collection, with the product AMP and a magnesium-bound phosphate complex being retained in the final structure. Despite the absence of interpretable electron density for P²- and P³-phosphates in the ligand-bound CT771 structure, comparison with the sulfate anion bound in the apo-CT771 structure (described in Supporting Information) suggests that both Lys43 and Lys92 could hydrogen bond with multiple phosphate oxygen atoms of Ap₄A (Fig. 5). In agreement with the ATP-bound A. aeolicus Ap₄A hydrolase structure (Fig. S3A), the P²-phosphate of Ap₄A would likely not make contacts with CT771. Finally, the second adenosyl binding site, if there is one, remains unoccupied, as seen in all previously reported Ap₄A hydrolase binary complexes. Analysis of both structures presented herein reveals that a minimal binding module of Np₄ is required for CT771 and this family of enzymes.

Substrate Induced Structural Changes within CT771

Structural comparison of the apo- and ligand-bound forms of CT771 presents insight into substrate binding and conformational changes required for ligand catalysis. Structural superposition of these two CT771 structures results in the alignment of 140/147 Cα atoms with an RMSD of 1.41 Å (Fig. 6). Notably, the catalytic α₁ Nudix helix and surrounding loops are almost identical (highlighted purple in Fig. 6) with only the side chain of Glu63 reorienting to coordinate a magnesium ion in the ligand-bound structure (Fig. 7). While the aforementioned structures are highly similar from a secondary structure perspective, several key loop regions have constricted inward towards the active site. Glu111 directly coordinates two magnesium ions in the ligand-bound structure (shifting ~2.9 Å from its apo-position, (Fig. 7)), as the loop connecting β6 and β7 shifts ~3.0 Å in order to accommodate this interaction (region two in Fig. 6). Additionally within this region, the sidechain of His38 moves ~6.7 Å as it rotates nearly 120° (Fig. 7), where it hydrogen bonds with the P¹-phosphate of the bound AMP molecule. Minor conformational changes within this region of CT771 also disrupt an intrachain disulfide bond between Cys32 and Cy113 upon ligand...
binding (Fig. S5). An extra turn of α3 is formed (region 3 in Fig. 6, residues Pro130 – Arg133) upon ligand binding, which places Phe129 in the appropriate position (shifting ~12.2 Å from its apo-position) to participate in π-π stacking interactions with the adenine ring of AMP (Fig. 4C). The extra turn of α3 facilitates an ~9.3 Å shift of Glu131 away from the active site cleft (Fig. 7), where it hydrogen bonds with the N6 atom of AMP. The loop connecting β5 and β6 shifts ~4.0 Å (region 4 in Fig. 6), however, it does not contact any of the bound ligands. Rather, this shift appears to facilitate bonding interactions involving Tyr80 (shifting ~4.9 Å from its apo-position) on β5 and Lys92 on β6, as these β-sheets subtly bow towards the bound AMP molecule (Fig. 7). Finally, the loop region connecting β1 and β2 has been significantly reordered in the ligand-bound structure (region 1 in Fig. 6); however, this likely arises from protein-protein contacts in crystal packing and does not appear to affect the CT771 active site.

Bioinformatic Analysis of CT771 and Orthologs within Chlamydia

CT771 is conserved across all sequenced members of Chlamydiaceae (sequence identities ranging from 71-99%). However, BLAST analysis of Chlamydiales, members of which include environmental Chlamydiae isolates, reveals the presence of genes orthologous to CT771 (sequence identities ranging from 35-42%) that were not previously clustered with other Chlamydiaceae members (aligned with CT771 in Fig. 2B). It thus appears that the presence of Ap₄A hydrolase is required in the basic biology of Chlamydiae. As such we recommend that the annotation of CT771, along with its current orthologs and the ones identified herein, be updated to asymmetric diadenosine tetraphosphatase (nudH).

DISCUSSION

BLAST analysis with the Nudix motif identifies a single target (CT771) within the obligate intracellular human pathogen Chlamydia trachomatis. This singular protein is conserved throughout both pathogenic and environmental Chlamydiales (Fig. 2B), suggesting it might play an important role in chlamydial biology. In order to better understand the function of this protein, structural studies were undertaken on CT771 from C. trachomatis, in both apo- and ligand-bound forms. Together, the structural and enzymatic characterization of CT771 suggests that this protein functions as an asymmetric diadenosine tetraphosphate (Ap₄A) hydrolase, with the likely ability to cleave diadenosine polyphosphates (n ≥4). Insights into substrate binding and catalysis have been obtained by efforts to co-crystallize CT771 with Ap₄A, with very similar active site electron density occupancy to the previously determined C. elegans Ap₄A binary complex. Intriguingly, the structure of CT771, only the second Ap₄A hydrolase determined from eubacteria, most closely resembles both bacterial and mammalian Ap₄A hydrolases, further supporting that these two classes are more closely linked than previously thought.

Both the apo- and ligand-bound CT771 crystal structures contained sulfate/phosphate anions bound near the Nudix motif within the putative active site (Fig. 4, S2). Numerous highly conserved side chains interact with these anions either directly or through coordinated metal ions. Based upon similarities to binary complexes from C. elegans and A. aeolicus Ap₄A hydrolases, these anions have been designated P¹-, P³-, and P⁴-phosphates; to our knowledge, this is the first description of all 3 contacts within the same protein. While a bound AMP molecule, which represents the first/primary nucleotide binding pocket, was also found within the putative active site, the absence of any appreciable binding contacts for the second adenosine group is consistent with the ability of this subclass of Nudix hydrolases to cleave diadenosine polyphosphates that are linked by four to six phosphates. Given the use of different substrates (AppCH₂ppA and Ap₄A, respectively), the highly similar ligands bound within the active sites of the C. elegans and CT771 binary
complexes are surprising. Despite the fact that Bailey et al. used a theoretically non-hydrolysable Ap₄A analog, it appears both compounds were cleaved by the crystallized hydrolases. Additionally, the previously discussed A. aeolicus binary complex ⁷ also utilized Ap₄A, which was also apparently hydrolyzed. However, a bound ATP molecule was modeled by Jeyakanthan et al. in the equivalent position of the AMP molecule within both the C. elegans ³⁷ and CT771 binary complexes. We speculate that this difference in bound ligand arises from the common use (and absence in the A. aeolicus binary complex) of high concentrations of MgCl₂ in the crystallization conditions of both the CT771 and C. elegans binary structures that facilitated the presence of a tightly bound Mg²⁺-phosphate complex.

Comparative analyses of the three phosphate binding sites (and the absence of protein mediated contacts at the putative P²⁻-phosphate site) from both the apo- and ligand-bound CT771 structures presented herein are in agreement with previously determined binary complexes; this supports the role of the Nudix motif in metal binding and proper substrate positioning, prior to catalysis of the P³⁻-P⁴⁻ phosphate bond. Previous studies have described the asymmetric Ap₄A hydrolase reaction mechanism as an “in-line” nucleophilic attack, which is mediated by a conserved glutamate functioning as a catalytic base, coordinating a water molecule and up to 3 magnesium ions ⁸. Structural and mutagenic studies have suggested that Glu63 (CT771 numbering) is the most likely candidate for the catalytic base; however the absence of structural evidence with an intact substrate has precluded definitive assignment of this residue. Within the CT771/AMP-PO₄ structure, three glutamate residues (59, 63 and 111) each coordinate with two of the three magnesium ions (Fig. 4D). Furthermore, Glu63, Glu111, and the Lys43 carbonyl coordinate with a common magnesium ion (identified as 1 in Fig. 4D) that is bound to a water molecule in an appropriate position to function as a nucleophile within the phosphoryl-binding groove of CT771 (denoted by a dashed arrow in Fig. 4D).

Enzymatic characterization of CT771 was facilitated by monitoring hydrolyzed Ap₄A via released ATP and cognate luciferase activity. Unfortunately, this method precluded the analysis of substrates other than Ap₄A due to an absence of released ATP. However, both in silico docking and crystallographic studies on CT771 support a preference for Ap₄A as a true substrate. Furthermore, enzymatic analyses did confirm Ap₄A hydrolase activity for CT771 and reveal an intriguing inverse relationship between concentration and the preferred divalent cation. While the majority of Ap₄A hydrolases preferentially utilize Mg²⁺ ⁴¹, instances of maximal activity in the presence of Mn²⁺ have been previously documented ⁴². Furthermore, in the presence of Mn²⁺ the Kₘ value for Ap₄A was roughly two orders of magnitude tighter than in the presence of Mg²⁺ for CT771, yet increasing concentrations of Mn²⁺ resulted in a decreased catalytic rate and efficiency (Table 4). The results presented by Szurmak et al. are in agreement with those reported here, particularly with respect to increasing catalytic rate in the presence of decreasing Mn²⁺ and increasing Mg²⁺ cations. It is difficult to speculate what physiological role this difference in metal ion affinity would serve within Chlamydia, but it could be a reflection of the intracellular environment where the chlamydial developmental cycle occurs.

Chlamydia species are obligate intracellular pathogens that typically rely on the infected host cell for the majority of their energetic needs. Thus, intracellular energy storage pools must be closely monitored in order to prevent the accumulation of toxic byproducts, or alarmones, which could potentially shutdown protein synthesis ⁴³. One such alarmone is Ap₄A, which is typically generated when aminoacyl-tRNA synthetases charge their cognate tRNA molecule ¹⁰. The chlamydial inclusion is not passively permeable to small molecules ≥520 Da ⁴⁴, indicating that Ap₄A (~800 Da) may not be freely diffusible and would need to be processed within the inclusion, providing a clear role for CT771 throughout the developmental cycle. Substrate multispecificity has been documented within a variety of
Nudix hydrolase family members\(^9\) and of particular note with respect to CT771, bacterial Ap\(_4\)A hydrolases have been demonstrated to efficiently remove the 5′ termini of mRNA, suggesting a role in translation regulation\(^{45}\). Clearly, further studies exploring alternative substrate preferences and specificities are needed, but at this time it appears that CT771 functions as a housekeeping gene, preventing Ap\(_4\)A levels from rising during periods of high metabolic activity. In support of this role, quantifiable levels of CT771 protein are detected in EBs\(^{46}\), indicating the organism is primed to handle a rapid burst of nascent transcription (~1-2 hpi)\(^{47}\) and a potential build-up of Ap\(_4\)A through tRNA charging during translation, upon infecting and entering the host cell.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**Acknowledgments**

We thank John Karanicas and Jimmy Budiardjo for technical assistance with the Luciferase assay.

**FUNDING SOURCE STATEMENT**

Use of the Advanced Photon Source was supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences under Contact No. DE-AC02-06CH11357, and use of the IMCA-CAT beamline 17-ID by the companies of the Industrial Macromolecular Crystallography Association through a contract with the Hauptman-Woodward Medical Research Institute. Use of the KU COBRE-PSF Protein Structure Laboratory was supported by grants from the National Center for Research Resources (5P20RR017708-10) and the National Institute of General Medical Sciences (8 P20 GM103420-10) from the National Institutes of Health. PSH, MLB and ANH were supported in part by National Institutes of Health grant AI079083.

**ABBREVIATIONS**

- Ap\(_4\)A: diadenosine 5′,5″′-P\(_{1},P_{4}\)-tetraphosphate
- Nudix: nucleoside diphosphates linked to any other moiety X
- IPTG: isopropyl 1-thio-β-D-galactopyranoside
- PDB: Protein Data Bank
- TEV: tobacco etch virus
- PISA: protein interfaces and assemblies server

**REFERENCES**


Biochemistry. Author manuscript; available in PMC 2015 January 14.


Figure 1. 2.60 Å Crystal Structure of CT771 from Chlamydia trachomatis
Crystal structure of CT771 shown in cartoon ribbon format using common rainbow colors (slowly changing from blue N-terminus to red C-terminus). Image is rotated 90° about the vertical axis on the right and amino acids comprising the Nudix motif are highlighted in purple (ball-and-stick), remainder of CT771 structure is colored gray.
Figure 2. Sequence Conservation of CT771 and Ap₄A Hydrolase Members

A, structural superposition of CT771 (gray) and top 3 structural hits from DALI search (Table II) in ribbon format. Structures correspond to Ap₄A hydrolases from the following organisms: *Homo sapiens* (PDB ID: 3U53, orange); *Caenorhabditis elegans* (PDB ID: 1KTG, purple); *Aquifex aeolicus* (PDB ID: 3I7V, cyan).

B, limited structure-based alignment of CT771 and Ap₄A Hydrolase Members was generated using ClustalW and rendered with ESPRIPT. Numbers above the sequences correspond to *C. trachomatis* CT771. The secondary structure of CT771 is shown above the alignment. Residues are colored according to conservation (cyan = identical and purple = similar) as judged by the BLOSUM62 matrix. Red circles below the sequences correspond to ligand interacting CT771 amino acid side chains in the binary complex, while blue circles correspond to sulfate-bound side chains in the apo-CT771 structure. Sequences used within the alignment are comprised of Ap₄A hydrolases with significant structural similarity to CT771 and BLAST-identified orthologs from environmental *Chlamydiae*. Accession numbers are detailed in *Materials and Methods*. 
Figure 3. Affect of Divalent Cations on Ap₄A Hydrolysis by CT771
A. Plot of CT771 activity versus substrate (Ap₄A) concentration in the presence of increasing Mg²⁺ concentration (●, 1.5mM; ∙, 3.0mM; and ▲, 5.0mM). B. Plot of CT771 activity versus substrate (Ap₄A) concentration in the presence of increasing Mn²⁺ concentration (●, 1.5mM; ∙, 3.0mM; and ▲, 5.0mM). Experimental details are defined in Materials and Methods.
Figure 4. Crystal Structure of CT771 Bound to Hydrolyzed Ap₄A Product

A, \( F_o \times F_c \) map (green mesh at 3.0 \( \sigma \) contour) of the refined structure in the absence of modeled ligands. Active site side chains within hydrogen bonding distance (2.5 – 3.5 Å) of bound ligands are depicted as cylinders (cyan) in all panels. CT771 backbone is depicted in cartoon ribbon format (orange). B, \( 2F_o - F_c \) map (blue mesh at 1.0 \( \sigma \) contour) of the refined structure with one AMP and one \( \text{PO}_4\text{Mg}^{2+} \) complex modeled per subunit. Color scheme is the same as panel A. C, Active site side chains within hydrogen bonding distance (2.5 – 3.5 Å) of AMP (yellow). CT771 backbone is depicted in cartoon ribbon format (lime). D, Active site side chains within hydrogen bonding (2.5 – 3.5 Å) and metal coordination (1.7 – 2.2 Å) distances of \( \text{PO}_4\text{Mg}^{2+} \) complex. Magnesium ions and water molecules are represented as spheres and colored purple and red, respectively. Dashed arrow identifies proposed nucleophilic water. Magnesium ions are numbered according to Table S1. The phosphate is colored orange. Color scheme is the same as panel C. Further information on the interaction distances within panels C and D can be found in Table S1.
Figure 5. Bound Sulfate Anion in apo-CT771 Structure Highlights P³-Phosphate Position

Structural superposition of apo- and ligand-bound CT771 (gray, only ligand-bound CT771 surface rendering is represented for clarity). CT771 side chains interacting with AMP (yellow) and the PO₄-Mg²⁺ complex (cyan) or sulfate (purple) are depicted in ball-and-stick format. Residues within appropriate hydrogen bonding distance (2.5 – 3.5 Å) of the sulfate anion are labeled and colored purple.
Figure 6. Structural Comparison of CT771 in Bound and Free Forms
Structural superposition of CT771 in free (orange) and bound (green) forms depicted in cartoon ribbon format. Catalytic Nudix motif is highlighted in purple within both structures. Several regions containing structural differences are highlighted with dashed lines, including: (1) the loop connecting β1 and β2, which likely arises from crystal packing in the bound form; (2) the loop connecting β6 and β7 constricts ~3.0 Å (as measured from the Glu111 Cα for each structure) towards the active site; (3) an extra turn of α3 is formed in the bound structure, resulting in the loop connecting α2 and α3 comprising different residues; and (4) the loop connecting β5 and β6 shifts ~4.0 Å (as measured from the Lys85 Cα for each structure) towards the active site.
Figure 7. CT771 Side Chain Conformational Changes upon Ligand Binding
Structural superposition of CT771 in free (orange) and bound (green) forms depicted as cylinders. Active site side chains within hydrogen bonding (2.5 – 3.5 Å) and metal coordination (1.7 – 2.2 Å) distances of AMP (yellow) and PO\(_4\)-Mg\(^{2+}\) complex (ball-and-sticks, purple spheres) are shown. The side chain of Glu111 in the free CT771 structure (orange) was truncated as described in the Materials and Methods section. Arrows indicate conformational changes greater than 2.0 Å, as described in the Results section.
## Table 1

<table>
<thead>
<tr>
<th></th>
<th>CT771-apo</th>
<th>CT771-AMP-PO₄</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Data Collection</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unit-cell parameters (Å)</td>
<td>$a=116.4, c=84.1$</td>
<td>$a=182.5, c=65.4, b=102.8, \beta=94.0$</td>
</tr>
<tr>
<td>Space group</td>
<td>$P6_522$</td>
<td>$C2$</td>
</tr>
<tr>
<td>Resolution (Å)$^f$</td>
<td>64.57-2.60 (2.72-2.60)</td>
<td>89.53-1.90 (1.93-1.90)</td>
</tr>
<tr>
<td>Wavelength (Å)</td>
<td>1.0000</td>
<td>1.0000</td>
</tr>
<tr>
<td>Temperature (K)</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Observed reflections</td>
<td>206,611</td>
<td>317,054</td>
</tr>
<tr>
<td>Unique reflections</td>
<td>10,818</td>
<td>96,781</td>
</tr>
<tr>
<td>$&lt;I_{\text{tot}}&gt;/\sigma(I)$</td>
<td>30.9 (2.2)</td>
<td>15.7 (2.1)</td>
</tr>
<tr>
<td>Completeness (%)$^f$</td>
<td>100 (100)</td>
<td>99.0 (99.6)</td>
</tr>
<tr>
<td>Multiplicity$^f$</td>
<td>19.1 (20.4)</td>
<td>3.4 (3.4)</td>
</tr>
<tr>
<td>$R_{\text{merge}}$ (%)$^f$</td>
<td>7.4 (151.6)</td>
<td>4.0 (61.7)</td>
</tr>
<tr>
<td>$R_{\text{meas}}$ (%)$^f$</td>
<td>7.8 (159.0)</td>
<td>4.7 (73.4)</td>
</tr>
<tr>
<td>$R_{\text{fin}}$ (%)$^f$</td>
<td>2.4 (47.9)</td>
<td>2.5 (39.3)</td>
</tr>
<tr>
<td>$CC_{1/2}$</td>
<td>100.0 (76.7)</td>
<td>99.9 (74.3)</td>
</tr>
<tr>
<td><strong>Refinement</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Resolution (Å)</td>
<td>38.1 - 2.6</td>
<td>39.6 - 1.9</td>
</tr>
<tr>
<td>Reflections (working/test)</td>
<td>10,788/520</td>
<td>93,746/4,696</td>
</tr>
<tr>
<td>$R_{\text{factor}}$ / $R_{\text{free}}$ (%)$^3$</td>
<td>18.9/21.7</td>
<td>20.4/25.0</td>
</tr>
<tr>
<td>No. of atoms (Protein/Ligand/Water)</td>
<td>1162/11/9</td>
<td>9024/256/327</td>
</tr>
<tr>
<td><strong>Model Quality</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>R.m.s deviations</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bond lengths (Å)</td>
<td>0.013</td>
<td>0.015</td>
</tr>
<tr>
<td>Bond angles (°)</td>
<td>1.139</td>
<td>0.874</td>
</tr>
<tr>
<td>Average $B$-factor (Å$^2$)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Atoms</td>
<td>71.7</td>
<td>56.8</td>
</tr>
<tr>
<td>Protein</td>
<td>71.8</td>
<td>57.2</td>
</tr>
<tr>
<td>Ligand</td>
<td>n/a</td>
<td>52.7</td>
</tr>
<tr>
<td>Solvent</td>
<td>69.3</td>
<td>51.3</td>
</tr>
<tr>
<td>Coordinate error (Å), maximum likelihood (Å)</td>
<td>0.28</td>
<td>0.27</td>
</tr>
<tr>
<td><strong>Ramachandran Plot</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Most favored (%)</td>
<td>93.8</td>
<td>96.3</td>
</tr>
<tr>
<td>Additionally allowed (%)</td>
<td>5.5</td>
<td>3.6</td>
</tr>
<tr>
<td>Outliers (%)</td>
<td>0.7</td>
<td>0.1</td>
</tr>
<tr>
<td>PDB ID</td>
<td>4ILQ</td>
<td>4MPO</td>
</tr>
</tbody>
</table>
1 Values in parenthesis are for the highest resolution shell.

2 \[ R_{\text{merge}} = \frac{\sum_{hkl} \sum_i |I_i(hkl) - <I(hkl)>|}{\sum_{hkl} \sum_i I_i(hkl)} \]

where \( I_i(hkl) \) is the intensity measured for the ith reflection and \(<I(hkl)>\) is the average intensity of all reflections with indices hkl.

3 \[ R_{\text{factor}} = \frac{\sum_{hkl} |F_{\text{obs}}(hkl) - |F_{\text{calc}}(hkl)|}{\sum_{hkl} |F_{\text{obs}}(hkl)|} \]

\( R_{\text{free}} \) is calculated in an identical manner using 5% of randomly selected reflections that were not included in the refinement.

4 \( R_{\text{meas}} \) = redundancy-independent (multiplicity-weighted) \( R_{\text{merge}} \)\(^2\), \( R_{\text{pim}} \) = precision-indicating (multiplicity-weighted) \( R_{\text{merge}} \)\(^4\).

5 \( C_{1/2} \) is the correlation coefficient of the mean intensities between two random half-sets of data\(^5\).
Table 2
CT771 DALI Search Statistics

<table>
<thead>
<tr>
<th>Protein name</th>
<th>PDB code</th>
<th>Z-scorea</th>
<th>RMSDb</th>
<th>Cα rangec</th>
<th>% idc</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Hs</em>Ap4A Hydrolase</td>
<td>3U53</td>
<td>18.5</td>
<td>2.2</td>
<td>134/144</td>
<td>31</td>
</tr>
<tr>
<td><em>Ce</em>Ap4A Hydrolase</td>
<td>1KTG</td>
<td>18.4</td>
<td>1.8</td>
<td>130/137</td>
<td>20</td>
</tr>
<tr>
<td><em>Aa</em>Ap4A Hydrolase</td>
<td>3I7V</td>
<td>18.1</td>
<td>1.8</td>
<td>130/134</td>
<td>28</td>
</tr>
<tr>
<td><em>Tt</em>Ap4A Hydrolase</td>
<td>1VC8</td>
<td>17.3</td>
<td>2.2</td>
<td>125/126</td>
<td>26</td>
</tr>
<tr>
<td><em>B. pseudomallei</em> MutT/Nudix family protein</td>
<td>1XSB</td>
<td>16.7</td>
<td>2.5</td>
<td>138/153</td>
<td>30</td>
</tr>
</tbody>
</table>

a Similarity score representing a function that evaluates the overall level of similarity between two structures. Z-scores higher than 8.0 indicate that the two structures are most likely homologous.
b Denotes the number of residues from the query structure that superimpose within an explicit distance cutoff of an equivalent position in the aligned structure.
c Denotes the percent sequence identity across the region of structural homology.
Table 3

AutoDock Vina Results

<table>
<thead>
<tr>
<th>Ligand</th>
<th>ΔG (kcal/mol)</th>
<th>Kd (μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diadenosine tetraphosphate (Ap4A)</td>
<td>-6.3</td>
<td>36.2</td>
</tr>
<tr>
<td>Adenosine diphosphate Ribose</td>
<td>-5.8</td>
<td>81.4</td>
</tr>
<tr>
<td>GDP-Mannose</td>
<td>-5.6</td>
<td>113</td>
</tr>
<tr>
<td>8-deoxy-dGTP</td>
<td>-4.5</td>
<td>672</td>
</tr>
</tbody>
</table>
### Table 4

Kinetic Constants for Ap4A Hydrolysis by CT771

<table>
<thead>
<tr>
<th>[MnCl₂] (mM)</th>
<th>$K_m$ (μM)</th>
<th>$k_{cat}$ (s⁻¹)</th>
<th>$k_{cat}/K_m$ (M⁻¹ s⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.5</td>
<td>7.28 ± 0.65</td>
<td>1.60 ± 1.75</td>
<td>2.2 × 10⁵</td>
</tr>
<tr>
<td>3.0</td>
<td>9.35 ± 0.36</td>
<td>1.51 ± 0.85</td>
<td>1.6 × 10⁵</td>
</tr>
<tr>
<td>5.0</td>
<td>4.98 ± 0.45</td>
<td>0.68 ± 0.90</td>
<td>1.4 × 10⁵</td>
</tr>
<tr>
<td>[MgCl₂] (mM)</td>
<td>$K_m$ (μM)</td>
<td>$k_{cat}$ (s⁻¹)</td>
<td>$k_{cat}/K_m$ (M⁻¹ s⁻¹)</td>
</tr>
<tr>
<td>1.5</td>
<td>407 ± 21</td>
<td>1.78 ± 0.04</td>
<td>4.4 × 10³</td>
</tr>
<tr>
<td>3.0</td>
<td>326 ± 42</td>
<td>2.24 ± 0.11</td>
<td>6.9 × 10³</td>
</tr>
<tr>
<td>5.0</td>
<td>333 ± 27</td>
<td>3.25 ± 0.10</td>
<td>9.8 × 10³</td>
</tr>
</tbody>
</table>