Atomic structures of the RNA end-healing 5′-OH kinase and 2′,3′-cyclic phosphodiesterase domains of fungal tRNA ligase: conformational switches in the kinase upon binding of the GTP phosphate donor

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Received September 24, 2019; Revised October 16, 2019; Editorial Decision October 20, 2019; Accepted November 07, 2019

ABSTRACT

Fungal tRNA ligase (Trl1) rectifies RNA breaks with 2′,3′-cyclic-PO₄ and 5′-OH termini. Trl1 consists of three catalytic modules: an N-terminal ligase (LIG) domain; a central polynucleotide kinase (KIN) domain; and a C-terminal cyclic phosphodiesterase (CPD) domain. Trl1 enzymes found in all human fungal pathogens are untapped targets for antifungal drug discovery. Here we report a 1.9 Å crystal structure of Trl1 KIN-CPD from the pathogenic fungus Candida albicans, which adopts an extended conformation in which separate KIN and CPD domains are connected by an unstructured linker. CPD belongs to the 2H phosphotransferase superfamily by dint of its conserved central concave β sheet and interactions of its dual HxT motif histidines and threonines with phosphate in the active site. Additional active site motifs conserved among the fungal CPD clade of 2H enzymes are identified. We present structures of the Candida Trl1 KIN domain at 1.5 to 2.0 Å resolution—as apoenzyme and in complexes with GTP•Mg²⁺, IDP•PO₄, and dGDP•PO₄—that highlight conformational switches in the G-loop (which recognizes the guanine base) and lid-loop (poised over the nucleotide phosphates) that accompany nucleotide binding.

INTRODUCTION

Invasive fungal infections, particularly Candidiasis and Aspergillosis, are a major cause of morbidity and mortality in individuals with neutropenia following cancer chemotherapry or hematopoietic stem cell transplantation. The recent emergence of Candida auris as a multi-drug-resistant invasive fungal pathogen in hospitalized patients is especially worrisome (1). The development of more effective (and less toxic) drugs for treatment of fungal infections hinges on defining new targets for antifungal drug discovery. We regard the tRNA splicing enzymes Trl1 (tRNA ligase) and Tpt1 (tRNA 2′-phosphotransferase) as attractive antifungal targets.

Fungal tRNA ligase Trl1 is an essential agent in the repair of RNA breaks with 2′,3′-cyclic phosphate and 5′-OH ends that are formed during tRNA splicing and the unfolded protein response (2,3). Trl1 executes three RNA repair reactions performed by three autonomous catalytic domains: (i) the 2′,3′-cyclic phosphate (>p) end is converted to a 3′-OH,2′-PO₄ by a C-terminal cyclic phosphodiesterase (CPD) module; (ii) the 5′-OH end is phosphorylated by a central GTP-dependent polynucleotide kinase (KIN) module; and (iii) the 3′-OH,2′-PO₄ and 5′-PO₄ ends are joined by an N-terminal ATP-dependent RNA ligase (LIG) module to form a 2′-PO₄, 3′-5′ phosphodiester splice junction (4–10). To complete the pathway, the essential enzyme Tpt1 transfers the 2′-PO₄ from the splice junction to NAD⁺ to form ADP-ribose 1″–2″ cyclic phosphate and nicotinamide via two unique steps (11–17). First, NAD⁺ reacts with the tRNA 2′-PO₄ to expel nicotinamide and generate a 2′-phospho-ADP-ribosylated RNA intermediate. Then, esterification of the ADP-ribose 2″-O to the tRNA 2′-PO₄ displaces the tRNA product and generates ADP-ribose 1″– 2″ cyclic phosphate.

Fungal Trl1 enzymes are potential therapeutic targets because their domain structures and biochemical mechanisms are unique compared to the RtcB-type tRNA repair systems elaborated by metazoa, archaea, and many bacteria.
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recognizes a 2′-AMP moiety in the RNA site that gave insights to how Tpt1 for LIG inhibition. There is no 5′-OH end, 3′-PO4 ends, a property that could be exploited for LIG inhibition. There is no 5′ kinase step in the RtcB pathway of RNA repair. Whereas polynucleotide kinases are widely distributed in nature, the KIN domains of fungal Trl1 enzymes are unique in that they have a strong preference for GTP as the phosphate donor (6,9,10,27,28). An ortholog of the Trl1 CPD domain is present in mammals in the form of CNP, a 2′,3′-cyclic nucleotide phosphodiesterase that is an abundant constituent of brain myelin (29). Mammalian CNP can substitute for Trl1 CPD as a tRNA-splicing enzyme in vivo in yeast (30), yet its role in mammalian RNA metabolism is uncharted. CNP is unlikely to be essential for mammalian cell function generally, insofar as CNP-null mice is viable to adulthood (31). [Adult CNP-null mice develop a neurodegenerative disorder despite maintaining normal myelin morphology. It is unclear whether this pathology is caused by a lack of CNP protein or CNP enzymatic activity.] A Tpt1 ortholog is also present in mammals, notwithstanding that mammalian RNA ligation by RtcB does not generate a 2′-PO4 end. Whereas it is not obvious what reactions Tpt1 performs in mammals, it is clear that Tpt1 is not essential for mammalian physiology, i.e., genetic ablation of mouse Tpt1 has no phenotypic consequences (32).

The above considerations highlight all four enzymatic activities in fungal tRNA splicing as potentially vulnerable to inhibition with selective toxicity for fungi versus mammalian hosts. To strengthen the case for tRNA splicing as an antifungal drug target, we aim to attain atomic structures of the tRNA splicing enzymes and elucidate the basis for their unique biochemical specificities. To that end, we recently reported the crystal structures of: (i) Trl1 LIG domain–as an apoenzyme and in complex with 2′-PO4 RNA splice junction (34); (ii) Trl1 KIN from the human pathogen Candida albicans in a complex with GDP•Mg2+, thereby revealing a ‘G-loop’ motif as the structural determinant of GTP specificity (28); and (iii) the Tpt1 ortholog from Clostridium thermocellum in a product-mimetic complex with ADP-ribose-1″-phosphate in the NAD” site and a pAp moiety in the RNA site that gave insights to how Tpt1 recognizes a 2′-PO4 RNA splice junction (34).

In the present study, we report the crystal structure of the two-domain KIN–CPD ‘end-healing’ component of Candida albicans Trl1 with a phosphoryl anion in the CPD active site. This is the first structure available for a Trl1 CPD enzyme and it highlights features shared with other 2H phosphoesterases as well as active site motifs specific to the fungal Trl1 CPD clade. We also report a series of structures of the Candida Trl1 KIN domain – as apoenzyme and in complexes with GTP, IDP, and dGDP – that underscore conformational switches in the G-loop (which recognizes the guanine base) and lid-loop (poised over the nucleotide phosphates) in response to nucleotide binding.

MATERIALS AND METHODS

CalTrl1 domain purifications

Recombinant Candida Trl1[401-832] [KIN-CPD] and Trl1[401-636]-D445N [KIN-D445N] proteins were purified from soluble extracts of IPTG-induced Escherichia coli BL21(DE3) pET28b-His6_bSmt3-Trl1 strains by sequential nickel-affinity, tag cleavage, tag removal by second nickel-affinity, and gel filtration steps (10,28). The peak Superdex 200 fractions were concentrated by centrifugal ultrafiltration to 9.5 mg/ml (for KIN-CPD) or 8.5 mg/ml (for KIN) in buffer containing 20 mM HEPES-NaOH, pH 7.5, 200 mM NaCl, 1 mM DTT, 5% glycerol. Protein concentrations were determined by using the BioRad dye reagent with bovine serum albumin as the standard.

Crystallization and structure determination of KIN-CPD

CalTrl1 KIN-CPD crystals were grown at 22°C by sitting drop vapor diffusion. A 1 μl solution of 0.18 mM KIN-CPD, 10 mM MgCl2, 2 mM GTP and 2 mM 2′-AMP was mixed with an equal volume of precipitant solution containing 0.2 M potassium dihydrogen phosphate, 20% PEG3350. Crystals appeared overnight and increased in size over four days, after which they were transferred to precipitant solution containing 25% glycerol prior to freezing. X-ray diffraction data were collected from a single crystal at the Advanced Photon Source beamline 24ID-C. Indexing and merging of the diffraction data were performed in HKL2000 (35). KIN-CPD crystalized in space group P21, with one protomer in the asymmetric unit. The structure was solved after initial molecular replacement with Phenix Phaser-MR (36) using as a search model the previously reported Candida KIN domain structure (pdb 5U32) (28). The electron density was improved by solvent flattening with Solomon (37) and the CPD domain was traced using Phenix Autobuild (36). Iterative model building into electron density was performed with O (38). Refinement was accomplished with Phenix.Refine (36). Data collection and refinement statistics are summarized in Supplementary Table S1.

Crystallization and structure determination of KIN apoenzyme

KIN crystals were grown at 22°C by hanging drop vapor diffusion after mixing a solution of 250 μM KIN-D445N with an equal volume of precipitant solution containing 0.3 M sodium dihydrogen phosphate, 20% PEG3350. Crystals appearing overnight were harvested and transferred to precipitant solution containing 25% glycerol prior to freezing. X-ray diffraction data were collected from a single crystal at the Advanced Photon Source beamline 24ID-E. KIN apoenzyme crystallized in space group C221, with two protomers in the asymmetric unit. The structure was solved by molecular replacement. There was no nucleotide in the active site; rather a phosphate anion was present at the position occupied by the GDP β-phosphate in pdb 5U32.
Crystallization and structure determination of KIN complexes with various nucleotides

Crystals were grown at 22°C by hanging drop vapor diffusion after mixing a solution of 250 μM KIN-D445N, 10 mM MgCl₂, and 2 mM GTP, ITP or dGTP with an equal volume of precipitant solution containing 0.15 M sodium dihydrogen phosphate, 20% PEG3350. KIN+GTP crystals appearing overnight were harvested immediately and transferred to precipitant solution containing 25% glycerol, 20 mM MgCl₂, and 1 mM GTP prior to freezing. KIN+ITP and KIN+dGTP crystals that were harvested after 4–5 days were transferred to precipitant solutions containing 25% glycerol, 20 mM MgCl₂ and 1 mM ITP or 1 mM dGTP prior to freezing. All of the KIN+NTP crystals were in space group P2₁2₁2₁ with one protomer in the asymmetric unit. The structures were solved by molecular replacement. Refinement statistics are compiled in Supplementary Table S1.

RESULTS

Structure of Candida albicans Trl1 KIN-CPD

C. albicans Trl1 is an 832-amino acid polypeptide composed of N-terminal ligase (LIG), central kinase (KIN), and C-terminal cyclic phosphodiesterase (CPD) domains. We showed previously that an RNA with 5′-OH and 2′,3′-cyclic phosphate ends could be spliced in vitro by reaction with a mixture of recombinant LIG (aa 1–400) and KIN-CPD (aa 401–832) proteins (28). Here we grew crystals of the Candida KIN-CPD protein by sitting drop vapor diffusion after mixing a solution of the protein solution containing KIN-CPD, GTP (KIN phosphatase donor), MgCl₂ (KIN cofactor), and 2′-AMP (CPD product mimic) with an equal volume of precipitant solution containing 200 mM potassium dihydrogen phosphate, 30% PEG3350. The crystals belonged to space group P2₁ and contained one KIN-CPD protomer in the asymmetric unit. The refined model of KIN-CPD at 1.95 Å resolution (Rwork/Rfree 0.185/0.221; Supplementary Table S1) comprised a KIN domain (aa 409–630) connected by an extended 13-aa linker to a CPD domain (aa 644–832) (Figure 1A and Supplementary Figure S1). The KIN domain had GDP and Mg²⁺ bound in the active site. The CPD domain had a phosphate anion in its active site (Figure 1A and Supplementary Figure S1). Amino acid sequence alignment of the KIN-CPD components of rRNA ligases from diverse fungal taxa indicates that the lengths of this linker segment are variable (13–40 aa) and the linker primary structure is not conserved (8). There were no contacts between the KIN and CPD modules within a single KIN-CPD polypeptide. The interactions of the KIN-CPD protomer with three symmetry related protein molecules in the crystal lattice are shown in Supplementary Figure S2. Analysis of the three interfaces in PISA returned a Complex Formation Significance Score of 0.000, implying that these interfaces are the result of crystal packing only. We regard this crystal packing interface as being of no physiological significance given that KIN-CPD is a monomer in solution (28). The structure of the KIN domain of KIN-CPD is highly similar to the structure of the isolated Candida Trl1 KIN domain in complex with GDP·Mg²⁺ (28).

Overview of the Trl1 CPD domain structure

The core CPD tertiary structure (shown in stereo in Figure 2A) consists of eight β strands that form a central anti-parallel β sheet. Flanking the β sheet are four α helices and two 3₁₀ helices. The secondary structure elements are aligned to the CPD primary structure in Figure 2B. The signature 2H phosphoesterase family motifs – 682HVT684 and 763HIT769 – are located in strands β2 and β5. The two histidines and two threonines coordinate a phosphate anion in the active site. Although we had preincubated KIN-CPD with the CPD product-like molecule 2′-AMP prior to crystallization, it was apparently out-competed for occupancy of the CPD active site by the 200 mM phosphate in the precipitant solution used to grow the crystals).

A DALI search (39) with the CPD structure retrieved other 2H phosphoesterase enzymes with varying extents of structural homology. The top hits were mammalian CNP (pdb 2YDD; Z score 13.8; rmsd of 3.0 Å at 150 Cx positions; 10% amino acid identity) and goldfish CNP (pdb 213E; Z score 12.3; rmsd of 2.6 Å at 149 Cx positions; 10% amino acid identity) (40,41). The next best homolog was E. coli ThpR (42), a biochemically validated RNA 2′,3′-cyclic phosphodiesterase enzyme (pdb 4QAK; Z score 6.1; rmsd of 3.5 Å at 111 Cx positions; 8% amino acid identity). Other hits with lower Z scores included: rotavirus VP3 (43), an...
Figure 1. Structure of Candida Trl1 KIN-CPD. The tertiary structure is shown as a cartoon model with the KIN domain in magenta, CPD domain in cyan, and the interdomain linker in beige. GDP and magnesium in the KIN active site are depicted as a stick model and a green sphere, respectively. Phosphate anion in the CPD active site is shown as a stick model.

Figure 2. Structure of the CPD domain. (A) Stereo view of the CPD tertiary structure, depicted as a cartoon model with magenta β strands, cyan α helices (numbered sequentially), and blue 3_10 helices. The phosphate anion in the active site and the histidine and threonine side chains that coordinate the phosphate are rendered as stick models. (B) Secondary structure elements (colored as in panel A) are displayed above the CPD primary structure. The histidine and threonine residues in the signature HxT motifs are denoted by red dots.
RNA 2′,5′-phosphodiesterase that degrades the innate immune signaling molecule 2′–5′ oligoadenylate (pdb 5AF2; Z score 5.4; rmsd of 3.6 Å at 98 Ca positions; 16% amino acid identity); and Arabidopsis ADP-ribose 1′−2″-cyclic phosphodiesterase (44), an enzyme that hydrolyzes the product of the Tpt1 reaction (pdb 1FSI; Z score 4.9; rmsd of 4.3 Å at 108 Ca positions; 8% amino acid identity). The primary structure conservation between Candida Trl1 CPD and the other 2H phosphoesterases is low and mostly localized to the 2H catalytic motifs.

**CPD active site**

Figure 3A shows a detailed stereo view of the CPD active site. The antiparallel β2 and β5 strands (containing the HxT motifs) and their preceding loops are arranged with pseudo-two-fold symmetry around the phosphate anion in the center of the active site. The enzymatic contacts to the phosphate reflect this two-fold symmetry. The phosphate oxygens receive hydrogen bonds from His682-Ne, Thr684-Oγ, His767-Ne and Thr769-Oγ. The tautomeric state of the two histidines, whereby histidine-Ne engages the phosphate, is ensured by hydrogen bonding of the histidine-Nε to the amino acid two positions upstream of the histidine: to the Glu680 carboxylate in the case of His682 and to the Gln765 main chain carbonyl in the case of His767 (Figure 3A). The phosphate oxygens are also engaged to four water molecules (labeled a, b, c, and d in Figure 3A) that bridge to the enzyme. Waters a and b are coordinated to the side chain amide N6 atoms of the residues four positions upstream of the catalytic histidines: Asn763 for water a and Gln768 for water b (Figure 3A). Water a makes an additional contact to Asn782 in helix α5. Waters c and d are engaged pseudo-symmetrically to the phosphate via bridging interactions with the HxT histidine main chain carbonyls and the threonine main chain amides (Figure 3A).

Whereas our CPD structure does not show the position and trajectory of the substrate RNA 3′ end, we can glean some insights via comparison to the structure of E. coli ThpR in complex with 2′-AMP (42). The ThpR active site aligns very well with that of Candida Trl1 CPD, notwithstanding that the overall folds of the two proteins have significant differences. The HxT histidines and threonines in ThpR superimpose on the CPD HxT motifs. The upstream HxT motif in ThpR contacts the AMP 3′-OH (the leaving group in the CPD reaction), suggesting that the upstream catalytic histidine acts as a general acid; we envision that His682 plays a similar role in Trl1 CPD. Of note, the ThpR HxT histidine main chain carbonyls and threonine main chain amides coordinate a pair of phosphate-neighboring water molecules equivalent to waters c and d in our CPD structure. A key distinction is that CPD lacks a counterpart of ThpR Arg130, which makes a bidentate contact to the 2′-AMP phosphate and is proposed to stabilize the transition-state of the ThpR CPD reaction (42). Mutating this arginine to alanine abolished ThpR CPD activity (42). In the case of Trl1 CPD, there is no positively charged functional group near the active site phosphate other than the two histidines.

We presume that HxT interactions with the scissile terminal phosphate suffice for catalysis of 3′ end healing during fun-gal tRNA splicing. The ThpR structure taught us that two aromatic amino acids – Phe8 and Phe48—pack against the nucleoside and nucleobase of 2′-AMP. The equivalent (and superimposable) residues in Trl1 CPD are Tyr647 (in strand β1) and His687 (in strand β2) (Figure 3A).

**Distinctive fungal CPD motifs**

The Candida albicans Trl1 CPD structure highlights active site constituents clustered in three motifs that are broadly conserved among other human fungal pathogens that are prevalent in the US, including Candida auris, Aspergillus fumigatus, Coccidioides immitis (the agent of Valley Fever), Histoplasma capsulatum, and Blastomyces dermatitidis (Figure 3B). In addition to the active site histidines and threonines (shaded gold in Figure 3B), the conserved functional residues include: the equivalents of Gln678, Asn763 and Asn782 (shaded green in Figure 3B) that make water-bridged phosphate contacts; and the Tyr/Phe and His equivalents of Tyr647 and His687 that we suggest interact with the terminal RNA nucleoside (shaded gray in Figure 3B). The shaded amino acids are conserved in Trl1 from the non-pathogenic yeast Saccharomyces cerevisiae (Figure 3B). Such conservation within the fungal clade may bode well for Trl1 CPD inhibitor discovery.

**New structures of Candida Trl1 KIN in GTP-bound and nucleotide-free states**

Our initial structure of the isolated KIN domain obtained from crystals grown from a mixture of KIN, GTP, and magnesium had GDPs⋅Mg2+ in the kinase active site (28), as did the KIN domain of KIN-CPD crystallized here in the presence of GTP and magnesium. This suggested that the γ phosphate of the added GTP was hydrolyzed during crystal growth. Alternatively, the KIN donor site was pre-filled with GDP during enzyme production in E. coli and GDP remained tightly bound during the KIN purification. To explore this issue, we grew crystals of a mutant kinase (KIN-D445N) from a mixture of KIN-D445N, GTP and magnesium. The mutant is inactive as a kinase because the Asp445 general base that deprotonates the 5′-OH RNA end for its attack on the GTP γ phosphate cannot perform this function when changed to an isosteric asparagine (28). The KIN+GTP crystals were in space group P2121210.193 and had one KIN protomer in the asymmetric unit. The structure was refined to a resolution of 1.85 Å (Rwork/Rfree 0.170/0.215) (Supplementary Table S1). The electron density clearly revealed the presence of GTP and magnesium in the kinase active site (Supplementary Figure S3). The tertiary structure of the KIN-GTP complex is shown in Figure 4A. We also grew crystals of KIN-D445N in the absence of added nucleotide. The apoenzyme crystallized in space group C2221 and had two KIN protomers in the asymmetric unit. The apoenzyme structure was refined to a resolution of 1.98 Å (Rwork/Rfree 0.193/0.236). Whereas neither apoenzyme protomer had a nucleotide in the active site, both protomers had a phosphate anion in the active site (derived from the 300 mM phosphate in the precipitant solution used for crystallization) at the position corresponding to the β phosphate of GTP or GDP. Figure 4A shows a superposition of the KIN apoenzyme B protomer
Figure 3. CPD active site. (A) Stereo view of the CPD active site highlighting CPD interactions with the phosphate anion. Amino acids are shown as stick models with beige carbons. Waters are denoted by red spheres. Atomic contacts are indicated by dashed lines. (B) Conservation of three active site motifs among Trl1 CPD domains from six species of human pathogenic fungi: *Candida albicans*, *Candida auris*, *Aspergillus fumigatus*, *Coccidioides immitis*, *Histoplasma capsulatum*, and *Blastomyces dermatitidis*. Positions of amino acid side chain identity or similarity in all six proteins are indicated by dots above the *C. albicans* sequence. The phosphate-binding histidines and threonines are highlighted in gold shading. Amino acids making water-bridged contacts to the phosphate are in green shading. Two aromatic residues imputed to interact with the terminal nucleoside are shaded gray. The corresponding motifs in *S. cerevisiae* Trl1 CPD are shown at bottom.

on the KIN•GTP structure. Pairwise alignment of the apo and GTP-bound KIN domains in DALI returned a Z score of 31.8 with a rmsd of 1.5 Å at 208 Cα positions.

The KIN fold comprises a central 5-strand β sheet surrounded by seven α helices (Figure 4A). The GTP phosphates are nestled into the P-loop (422GCGKTT427) (Figure 4A and B). The α4 and α5 helices form a ‘lid’ over the GTP phosphates, wherein the lid helices is disordered, in both the KIN apoenzyme and KIN•GTP•Mg2+ structures (Figure 4A) as it was in the KIN•GDP•Mg2+ structure reported previously (28). Comparison of the KIN apoenzyme and KIN•GTP•Mg2+ structures reveals shifts in the positions of the lid helices (α4 and α5) and of α7 that accompany nucleotide binding (Figure 4A). Most striking, however, is the complete remodeling of the conformation of the G-loop element in apoenzyme versus KIN•GTP. The upshot of this difference (described below) is that the apoenzyme structure is incompatible with GTP binding and thus a conformational switch is necessary to mobilize and properly position the NTP specificity determinants resident in the G-loop.

KIN•GTP active site

Figure 4B shows a detailed stereo view of the atomic interactions with the GTP nucleotide in the KIN active site. As in the previous KIN•GDP•Mg2+ structure (28), the guanine nucleobase is engaged by a network of hydrogen bonds from G-loop amino acids Asp583 (to guanine N1 and N2 from the Asp carboxylate) and Ser589 (to guanine O6 and N7, from the main chain amide and Oγ, respectively). The Arg528 side chain of the lid α4 helix makes a π-cation stack on the guanine base. Arg528 also makes a hydrogen bond to the G-loop Ser589 side chain. There are no enzymic contacts to the ribose sugar in the KIN•GTP•Mg2+ complex. Main chain amide nitrogens of the 422GCGKTT427 P-loop donate hydrogen bonds to the α and β phosphates of GTP. The Thr427 hydroxyl donates a hydrogen bond to the GTP α phosphate. The Lys425 side chain makes a bifurcated ionic interaction with the β and γ phosphates. Arg528 of the lid-loop also engages the GTP γ phosphate (Figure 4B). Arg528 was disordered in the previous KIN•GDP structure (28) and is also not visualized in the present KIN apoenzyme structure (Figure 4C), suggesting that this arginine at
Figure 4. Structures of the KIN•GTP•Mg\(^{2+}\) complex and KIN apoenzyme. (A) Stereo view of the superimposed tertiary structures of the KIN•GTP•Mg\(^{2+}\) complex (colored gold) and the KIN apoenzyme (colored cyan). The seven α helices flanking the central β sheet are numbered sequentially. GDP is depicted as a stick model with yellow phosphorus atoms. Magnesium is depicted as a magenta sphere. The phosphate anion in the KIN apoenzyme is shown as a stick model with a green phosphorus atom. (B) Stereo view of the KIN active site highlighting interactions with GTP•Mg\(^{2+}\). Amino acids are shown as stick models with beige carbons. Magnesium and waters are denoted by magenta and red spheres, respectively. Atomic contacts are indicated by dashed lines. (C) Stereo view of the equivalent active site region of the KIN apoenzyme, highlighting a conformational switch in the G-loop segment.
the start of the lid-loop becomes ordered when it can engage the γ phosphate. Two of the sites in the octahedral coordination complex of the Mg\(^{2+}\) ion are occupied by nonbridging β and γ phosphate oxygens of GTP. The other sites are filled by the Thr426 Oγ atom and three waters (Figure 4B). These features of the GTP•Mg\(^{2+}\) substrate complex in the phosphate donor site suggest a mechanism whereby Lys425, Arg532, and the metal complex orient the γ phosphate for nucleophilic attack by the polynucleotide 5′-OH and stabilize the extra negative charge developed in the pentacoordinate transition state during an associative in-line phosphoreryl transfer.

Conformational switch in the G-loop

A stereo view of the phosphate donor site fold in the KIN apoenzyme structure is shown in Figure 4C, onto which the guanine base from the KIN\(\bullet\)GTP\(\bullet\)Mg\(^{2+}\) structure is superimposed. The 579 KLSKDENSLKSS580 segment that comprises the G-loop in the nucleotide-bound KIN structures includes only two amino acids that adopt a secondary structure: Ser589 and Ser590 at the outset of the α6 helix. By contrast, in the KIN apoenzyme, the α6 helix is extended proximally into the erstwhile G-loop so that the helix initiates with 585 NS—i.e. four amino acids upstream of its onset in the GTP/GDP-bound state—and the conformation/trajectory of the residual preceding loop is completely different (Figure 4A). Thereby, the Asp583 side chain that confers GTP specificity is pointed away from the guanosine binding site and the extended α6 helix occludes the site occupied by the guanine base in the KIN\(\bullet\)GTP\(\bullet\)Mg\(^{2+}\) structure (Figure 4C). We surmise that a conformational switching entailing unfolding of the proximal α6 helix segment accompanies, and is necessary for, engagement of the GTP phosphate donor.

KIN structures in complex with inosine and deoxyguanosine nucleotides

By testing a series of purine ribonucleoside triphosphate analogs as substrates for \(\text{RNA}^\theta\text{P}\) phosphorylation by Candida T\(\text{R}\) KIN (28), it was found that inosine triphosphate (ITP) was the most effective substrate, implying that the exocyclic 2-NH\(_2\) moiety of GTP is not an essential determinant of nucleobase recognition per se. By contrast, 2-aminopurine triphosphate was a much less effective substrate for the kinase reaction. These results highlight the importance of the 6-oxo atom of guanine for phosphate donor function. To better understand base recognition by Candida KIN, we grew crystals from a mixture of KIN-D445N, ITP, and magnesium. The KIN + ITP crystals were in space group \(P2_1_2_1_2_1\) and had one KIN protomer in the asymmetric unit. The structure was refined to a resolution of 1.53 Å (R\(_\text{work}\)/R\(_\text{free}\) 0.174/0.208) (Supplementary Table S1). The active site contained inosine diphosphate (IDP) and a phosphate anion but did not include a magnesium ion (Figure 5A). We surmise that the input ITP ligand was hydrolyzed to IDP in crystallo [see Discussion for why this might be] and that the metal ion dissociated from the enzyme; the retention of a phosphate anion adjacent to, but clearly distinct from, the IDP β phosphate is presumably driven by the presence of 150 mM phosphate in the precipitant solution used to grow the crystals. In the KIN\(\bullet\)IDP structure, Arg528 of the lid α4 helix makes a r-carboxyl stack on the inosine base and Ser590 of the G-loop makes hydrogen bonds to the inosine N7 and O6 atoms (Figure 5A), just as in the KIN\(\bullet\)GTP\(\bullet\)Mg\(^{2+}\) structure (Figure 3B). However, the G-loop has shifted slightly (by 3.5 Å at the Asp583 Ca atom and 1.7 Å at the Asp583 Cα atom) and the Asp583 side chain now makes a single hydrogen bond to the inosine N1 atom (Figure 5A). The P-loop Lys425 side chain makes a bifurcated contact to the IDP β phosphate and the phosphate anion, analogous to the β-γ phosphate-bridging contact seen in the KIN\(\bullet\)GTP\(\bullet\)Mg\(^{2+}\) complex. Absent a metal, the P-loop Thr426 side chain donates a hydrogen bond to the IDP β phosphate. A notable feature of the KIN\(\bullet\)IDP structure is that the 532 RGDNHQSIK540 lid-loop peptide is visible and in contact with the IDP nucleotide and phosphate anion in a surprising fashion (Figure 5A). In particular, Arg532, which contacts the γ phosphate in the GTP\(\bullet\)Mg\(^{2+}\) complex, is now oriented in the opposite direction and makes no contacts. The main chain carbonyls of Arg532 and Gly533 make hydrogen bonds to the IDP ribose 3′-OH. The Asp534 carboxylate is within close hydrogen-bonding distance of the IDP β phosphate (Figure 5A), counter to...
Arg511 and His515 are essential for Trl1 function in vivo (6,8). Replacing Arg511 with lysine or glutamine was lethal. The structure was refined to a resolution of 1.7 Å (Supplementary Table S1). The active site contained dGTP and a phosphate anion, but no magnesium (Figure 5B). The G-loop conformation and nucleobase contacts in the KIN•dGDP structure were identical to that of the KIN•GTP•Mg²⁺ structure. The lid-loop was visible and made contacts to dGDP similar to those seen in the KIN•IDP structure. The phosphate anion in the KIN•dGDP complex was coordinated to His536, Asn547, and Asn444 (Figure 5B).

Mutational analysis of Candida Trl1 KIN

The C. albicans SGCGKTT427 P-loop and S. cerevisiae RGDHNQS1540 lid-loop sequences in the KIN domain are well conserved among fungal Trl1 proteins (8). The corresponding sequences in S. cerevisiae Trl1 are 401 GCGKTT406 and 511 RGNHQS1519, respectively. Previous mutational analysis of the S. cerevisiae Trl1 P-loop showed that single-alanine substitutions for Lys425 and Thr405 had no effect on vegetative growth and little effect on GTP-dependent RNA kinase activity in vitro, whereas a K404A-T405A double-mutation was lethal and abolished GTP-dependent RNA kinase activity (6). By contrast a single alanine mutation of the P-loop Lys425 residue of C. albicans Trl1 was lethal in vivo (as gauged by trl1Δ complementation in S. cerevisiae) and eliminated GTP-dependent RNA kinase activity in vitro (8). By contrast a single alanine mutation of the P-loop Lys425 residue of C. albicans Trl1 was lethal in vivo (as gauged by trl1Δ complementation in S. cerevisiae) and eliminated GTP-dependent RNA kinase activity in vitro (8). Alanine scanning of the S. cerevisiae Trl1 lid-loop showed that Arg511 and His515 are essential for Trl1 function in vivo (6,8). Replacing Arg511 with lysine or glutamine was lethal in vivo, as was replacing His515 with either glutamine or asparagine (8). By contrast, alanine substitutions for S. cerevisiae lid-loop residues Asn513, Asn514, Gln516 and Ser517 had no apparent effect on cell growth (8). In light of the divergent effects of the P-loop lysine mutation on Saccharomyces versus Candida Trl1 and the conformational variability of the lid-loop in the available suite of Candida Trl1 KIN structures, we mutated the Candida P-loop threonines adjacent to Lys425 and we mutated components of the Candida lid loop. We changed P-loop Thr426 (which coordinates the magnesium cofactor in the GTP•Mg²⁺ complex) and Thr427 (which engages the α phosphate in all of the nucleotide-bound KIN structures) to alanine. We introduced alanine in lieu of lid-loop residues Arg532 (which contacts the γ phosphate in the GTP•Mg²⁺ complex), Asp534 (which contacts the β phosphate in the IDP and dGDP complexes), Asn535, His536 (which contacts the phosphate anion in the IDP and dGDP complexes), and Gln537. These alanine changes were made in the context of the full-length Candida Trl1 protein encoded on a CEN LEU2 plasmid under the transcriptional control of the S. cerevisiae TPI1 promoter (28). We then tested the CalTRL1-Ala alleles by plasmid shuffle for complementation of an S. cerevisiae trl1Δ p[CEN URA3 SceTRL1] strain (6).

The Candida T426A allele was lethal in vivo, i.e., Leu+ trl1Δ CalTRL1-T426A transformants were unable to form colonies after incubation for 8 days at 20, 30, or 37°C on agar medium containing 5-fluoroorotic acid (FOA), a drug that selects against the p[CEN URA3 SceTRL1] plasmid. Thus, the metal coordinating P-loop threonine is essential for Candida Trl1 kinase activity in vivo. The other seven alleles yielded viable FOA-resistant trl1Δ CalTRL1-Ala isolates. These strains were tested by serial dilution for growth on YPD agar medium at 30°C and 37°C in parallel with a trl1Δ Δ CALTRL1 control. The P-loop T427A and lid-loop N476A, D534A, N535A, H536A and Q537A mutant strains grew as well as wild-type at both temperatures, as gauged by colony size (Figure 6). By contrast, the R532A strain was severely defective for growth, forming pinpoint colonies at 30°C and 37°C (Figure 6). The importance of Arg532 for Candida Trl1 activity is in keeping with our inferences from the KIN•GTP•Mg²⁺ structure that the contacts of Arg532 with the GTP γ phosphate are on-pathway in the kinase reaction. Conversely, the dispensability of Asp534 for Trl1 function suggests that its β phosphate contacts in the IDP and dGDP complexes are off-pathway.
Mutational analysis of *Candida* Trl1 CPD

The His682, Thr684, His767 and Thr769 residues of the signature HxT motifs that engage the active site phosphate were replaced individually by alanine in the context of full-length *Candida* Trl1 and the *CalTrl1-Ala* mutants were tested for trl1Δ complementation in *S. cerevisiae*. The H767A mutation was lethal. The H682A strain was sick at 30°C and unable to grow at 37°C (Figure 6). The T684A and T769A strains grew fairly well at 30°C but displayed ts growth defects at 37°C, with T769A being less fit than T684A (Figure 6). The effects of mutating the *Candida* HxT histidines were in accord with previous findings for *S. cerevisiae* Trl1 and the plant tRNA ligase AtRLN (a Trl1 ortholog that complements *S. cerevisiae* trl1Δ), i.e. changing the proximal HxT histidine to alanine elicited a ts growth phenotype whereas alanine mutation of the distal histidine was lethal (6,8). Different phenotypes were seen for the SceTrl1 and AtRLN HxT threonine-to-alanine mutations *vis-à-vis* the *Candida* enzyme: the proximal threonine mutation in SceTrl1 and AtRLN was lethal while the distal threonine SceTrl1 and AtRLN mutant strains grew as well as wild-type at 30°C and 37°C (8).

**DISCUSSION**

The KIN-CPD structure presented here is, to our knowledge, the first instance of an atomic structure of the RNA 2′,3′-cyclic phosphodiesterase component of fungal tRNA ligase. With this accomplished, we now have exemplary structures in hand for all four enzymes that catalyze RNA repair during fungal tRNA splicing (28,33,34). The present study also extends our knowledge of the NTP donor specificity of the RNA kinase module of fungal tRNA ligase and reveals conformational changes in the G-loop and lid-loop of the KIN domain triggered by guanine nucleotide binding. Our findings bolster the case for tRNA splicing as an antifungal drug target.

**CPD structure**

It has been thought for some time, on the basis of its tandem HxT motifs, that the CPD domains of fungal and plant tRNA ligases belong to the 2H phosphoesterase superfamily (45). This classification was fortified by the findings that certain mutations of the signature HxT motif drastically reduce CPD activity *in vitro* and are lethal *in vivo* when tested for complementation of *S. cerevisiae* trl1Δ (6,8,9,46). The crystal structure of *Candida* Trl1 CPD confirms its membership in the 2H superfamily by virtue of the conservation of the fold of the central concave β sheet and the atomic interactions of the HxT motif histidines and threonines with a phosphate anion in the active site, which we regard as a mimic of the scissile RNA 2′,3′-cyclic phosphate. That said, the degree of amino acid identity between Trl1 CPD and other 2H family members is low and the Trl1 CPD diverges from other 2H enzymes with respect to structural elements outside the active site. On the other hand, we discern three active site motifs in the *C. albicans* CPD that are well conserved in the CPD domains of tRNA ligases from multiple other human fungal pathogens. Conserved residues in these motifs make water-mediated contacts to the phosphate in the CPD crystal structure or are imputed to interact with the terminal nucleoside of the RNA substrate. The array of hydrogen-bonding residues and several aromatic residues in the CPD active site (and the relative paucity of charged basic and acid residues) could provide a pocket for the binding of a small-molecule CPD inhibitor.

A striking feature of the KIN-CPD structure is that it adopts an extended conformation in which the KIN and CPD domains are far apart and connected by an unstructured peptide linker. It is possible that this particular extended conformation is a matter of crystal packing and that the domain arrangement of KIN-CPD in solution is flexible about the linker peptide. Although the KIN-CPD structure does not suggest a specific and stable interface between KIN and CPD, we do not exclude the prospect that an interdomain interface exists in Trl1 that is dependent on the presence of the LIG domain. A previous study employing bifunctional lysine chemical crosslinking agents to probe at low resolution the fold of the CPD domain of *Coccidioides immitis* Trl1 (10) identified a network of four lysine-lysine crosslinks involving four amino acids (Lys635, Lys693, Lys703 and Lys707 in *C. immitis* CPD) that are conserved in *C. albicans* (as Lys642 in the linker; Lys692 in the second 310 helix; Lys702 in helix α3; and Lys706 in helix α3). These residues are indeed in physical proximity in the *Candida* CPD tertiary structure (Supplementary Figure S4) at atomic distances (9–12 Å) consistent with the chemical crosslinking results.

**KIN substrate recognition and conformational switches**

We suggested previously (28) that the peculiar G-loop and guanine-binding pocket of *Candida* KIN present a plausible target for a small molecule that could selectively inhibit the fungal Trl1 kinase without affecting the many other P-loop superfamily phosphotransferases that do not share the guanine specificity of the fungal Trl1 kinase domains. The structures solved presently of KIN in complexes with deoxyguanosine and inosine nucleotides underscore that: (i) the enzyme does not discriminate between ribose and deoxyribose sugars in the phosphate donor; (ii) the exocyclic –NH2 of guanine is not essential for occupancy of the G-loop by inosine, which still makes a hydrogen bond from N1 to G-loop Asp583; and (iii) the purine O6 atom is the key determinant of substrate recognition via its hydrogen bonding interactions with Ser590, which is essential for KIN activity (28). The notable finding here was that in the structure of the nucleotide-free KIN apoenzyme, the conformation of the erstwhile G-loop is rearranged such that the guanine-binding pocket is occluded. We surmise that the ordered N-terminal segment of the α6 helix of the apoenzyme must become disordered upon GTP binding and the preceding loop is reshaped accordingly to form the G-loop pocket. This situation raises the prospect that a small molecule that binds to the apo state of the G-loop segment might act as conformational trap to inhibit the kinase.

The present suite of nucleotide-bound KIN structures also highlights the conformational flexibility of the lid including: (i) a ‘closing’ movement of the lid α4 and α5 helices over the nucleotide in the transition from apoenzyme to GTP•Mg2+ complex; and (ii) conformational variability of
the lid-loop ranging from: entirely disordered (apoenzyme and GDP•Mg\(^{2+}\) complexes); to a state in which Arg532 at the margin of the disordered loop interacts with the GTP γ phosphate (GTP•Mg\(^{2+}\) complex); to a situation in which the loop is visible in the magnesium-free dGDP and IDP complexes while making contacts to these nucleotides that we suspect are off-pathway (owing to the acidic crystallization conditions that favor the atypical Asp534 hydrogen bond to the NDP β phosphate). Mutagenesis of the KIN domain lid-loop fortifies our view that the interactions of Arg532 with the GTP γ phosphate in the GTP•Mg\(^{2+}\) complex are indeed on-pathway, i.e. an R532A mutation compromises (but does not abolish) Trl1 function *in vivo*. The corresponding P-loop arginine in the kinase domain of the bacterial RNA repair enzyme Pnkp makes analogous contact to the γ phosphate of its NTP substrate; mutating this Pnkp arginine (Arg120) to alanine reduced the kinase specific activity to 42% of the wild-type level (47). Comparing the effects of mutations at equivalent positions in the KIN domains of *S. cerevisiae* and *C. albicans* Trl1 enzymes indicates that the contacts of the individual loop and lid-loop amino acids can vary in their importance. For example, whereas the P-loop lysine and metal-binding threonine are both individually essential in *C. albicans* Trl1 (ref. 28 and Figure 6), equivalent single-alanine P-loop mutants in *S. cerevisiae* Trl1 are viable – though a double-mutation of Lys-Thr to Ala-Ala is lethal in *S. cerevisiae* (6). Alanines in lieu of the lid-loop arginine and histidine side chains are lethal in *S. cerevisiae*; the equivalent arginine mutant in *C. albicans* Trl1 is sick but alive and the histidine mutation is apparently benign. The take-home lesson is that structure-activity relations are not precisely transferable between fungal Trl1 kinase orthologs.

Our use of the KIN general acid mutant D445N in the present structural studies was prompted by a desire to avoid hydrolysis of the input NTP substrate that was evident in the initial crystallization of wild-type KIN in the presence of GTP and magnesium (28). This maneuver clearly did not prevent NTP hydrolysis, insofar as several of the structures determined here had NDP in the KIN active site. It is conceivable that NTP hydrolysis by the KIN in the presence of magnesium is influenced by the acidic condition that favored successful crystallization. However the key parameter anent NTP hydrolysis seems to be the duration of the crystallization experiment: to wit, all of the crystals that contained NDP in the active site were harvested after 4–5 days, whereas the GTP•Mg\(^{2+}\) structure was obtained from a crystal that was harvested immediately after overnight growth. Notwithstanding that the electron density map clearly reveals the GTP•Mg\(^{2+}\) complex, we suspect that a fraction of the KIN enzymes in the crystal may have hydrolyzed GTP to GDP, insofar as the B factor of the GTP γ phosphorus atom (=51) was higher than the B factors of the α and β phosphorus atoms (=25).

The available LIG, KIN, and CPD structures of fungal tRNA ligase each have mechanistically instructive ligands in their respective active sites. The outstanding questions at this point concern how each catalytic module recognizes the specific RNA substrate on which it acts: RNA2−p for CPD; \(^{5′}OH\) RNA for KIN, and pRNA2−p for LIG. Our attempts to date to crystallize KIN-D445N in the presence of GTP, magnesium, and a \(^{5′}OH\) oligonucleotide to capture a Michaelis complex of the kinase reaction (a strategy that was successful with *Clostridium thermocellum* polynucleotide kinase; 48) have not been fruitful, insofar as the conditions that led to crystals (none of which contained the oligonucleotide) were the same as those that yielded KIN•nucleotide complexes, i.e., it is likely that high concentrations of phosphate and/or acidic pH were inimical to binding of the nucleic acid substrate to KIN (or of 2′-AMP to CPD). Still, the available Trl1 domain structures do suggest that it might be possible to inhibit multiple enzymatic steps in the fungal tRNA splicing pathway, which, if achieved, would reduce the risk of the emergence of drug resistance. The case for Trl1 as a drug target for Candidiasis is bolstered by the recent identification of *TRL1* as an essential gene in *C. albicans* (49).

### DATA AVAILABILITY

Structural coordinates have been deposited in Protein Data Bank under accession codes 6U05 (KIN-CPD), 6U00 (KIN-D455N-apo), 6U03 (KIN-D445N•GTP•Mg), 6TZX (KIN-D445N-IDP), 6TZ0 (KIN-D445N•dGDP) and 6TZM (KIN-D445N•GDP•Mg).

### SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

### FUNDING

National Institutes of Health [R35-GM126945 to S.S.]; Geoffrey Beene Cancer Research Center (to S.S.); Deutsche Forschungsgemeinschaft [39432028 to A.B.]; MSKCC structural biology core laboratory is supported by National Cancer Institute [P30-CA008748]; X-ray diffraction data were collected at synchrotron facilities supported by grants and contracts from the National Institutes of Health [P41GM103403, HEI-S10RR029205]; Department of Energy [DE-AC02-06CH11357]. Funding for open access charge: National Institutes of Health [R35-GM126945]. Conflict of interest statement. None declared.

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