Two-metal versus one-metal mechanisms of lysine adenylylation by ATP-dependent and NAD\textsuperscript{+}-dependent polynucleotide ligases

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Polynucleotide ligases comprise a ubiquitous superfamily of nucleic acid repair enzymes that join 3′-OH and 5′-PO\textsubscript{4} DNA or RNA ends. Ligases react with ATP or NAD\textsuperscript{+} and a divalent cation cofactor to form a covalent enzyme-(lysine-N\textsubscript{ε})-adenylate intermediate. Here, we report crystal structures of the founding members of the ATP-dependent RNA ligase family (T4 RNA ligase 1; Rnl1) and the NAD\textsuperscript{+}-dependent DNA ligase family (Escherichia coli LigA), captured as their respective Michaelis complexes, which illuminate distinctive catalytic mechanisms of the lysine adenylylation reaction. The 2.2-Å Rnl1-ATP-(Mg\textsuperscript{2+})\textsubscript{2} structure highlights a two-metal mechanism, whereby: a ligase-bound “catalytic” Mg\textsuperscript{2+} (H\textsubscript{2}O\textsubscript{3})\textsuperscript{2−} complex lowers the pK\textsubscript{a} of the lysine nucleophile and stabilizes the transition state of the ATP α phosphate; a second octahedral Mg\textsuperscript{2+} coordination complex bridges the β and γ phosphates; and protein elements unique to Rnl1 engage the γ phosphate and associated metal complex and orient the pyrophosphate leaving group for in-line catalysis. By contrast, the 1.55-Å LigA-NAD\textsuperscript{+}-Mg\textsuperscript{2+} structure reveals a one-metal mechanism in which a ligase-bound Mg\textsuperscript{2+} (H\textsubscript{2}O\textsubscript{3})\textsuperscript{2−} complex lowers the lysine pK\textsubscript{a} and engages the NAD\textsuperscript{+} α phosphate, but the β phosphate and the nicotinamide nucleoside of the nicotinamide mononucleotide (NMN) leaving group are oriented solely via atomic interactions with protein elements that are unique to the LigA clade. The two-metal versus one-metal dichotomy demarcates a branchpoint in ligase evolution and favors LigA as an antibacterial drug target.

metal catalysis | covalent nucleotidytransferase | lysyl-AMP

Polynucleotide ligases join 3′-OH and 5′-PO\textsubscript{4} RNA termini via a series of three nucleotidyl transfer steps. In step 1, a ligase reacts with ATP or NAD\textsuperscript{+} to form a covalent ligase-(lysyl-N\textsubscript{ε})-AMP intermediate and release pyrophosphate (PP\textsubscript{i}) or nicotinamide mononucleotide (NMN). In step 2, ATP is transferred from ligase-adenylate to the 5′-PO\textsubscript{4} DNA or RNA end to form a DNA-adenylate or RNA-adenylate intermediate (AppDNA or AppRNA). In step 3, ligase catalyzes attack by a DNA or RNA 3′-OH on the polynucleotide-adenylate to seal the two ends via a phosphodiester bond and release AMP. All steps in the ligase pathway require a divalent cation cofactor.

The autoadenylylation reaction of polynucleotide ligases is performed by a nucleotidytransferase (NTase) domain that is conserved in ATP-dependent DNA and RNA ligases and NAD\textsuperscript{+}-dependent DNA ligases (1, 2). The NTase domain includes defining peptide motifs that form the nucleotide-binding pocket. Motif I (Kx(DG)\textsubscript{2}) contains the lysine that becomes covalently attached to the AMP. As Robert Lehman pointed out in 1974 (3), it is unclear how lysine (with a predicted pK\textsubscript{a} value of ~10.5) loses its proton at physiological pH to attain the unprotonated state required for attack on the α phosphorus of ATP or NAD\textsuperscript{+}. In principle, a ligase might use a general base to deprotonate the lysine. Alternatively, the pK\textsubscript{a} could be driven down by positive charge potential of protein amino acids surrounding lysine-\textsubscript{N\epsilon}. Several crystal structures of ligases absent metals provided scant support for either explanation. In these structures, the motif I lysine nucleophile is located next to a motif IV glutamate or aspartate side chain (2, 4–6). The lysine and the motif IV carboxylate form an ion pair, the anticipated effect of which is to increase the pK\textsubscript{a} of lysine by virtue of surrounding negative charge. It is unlikely that a glutamate or aspartate anion could serve as a general base to abstract a proton from the lysine cation. A potential solution to the problem would be if a divalent cation abuts the lysine-\textsubscript{N\epsilon} and drives down its pK\textsubscript{a}.

A metal-driven mechanism was revealed by the recent crystal structure of Naegleria gruberi RNA ligase (NgrRnl) as a step 1 Michaelis complex with ATP and manganese (its preferred metal cofactor) (7). The key to capturing the Michaelis-like complex was the replacement of the lysine nucleophile by an isoteric methionine. The 1.9-Å structure contained ATP and two manganese ions in the active site. The “catalytic” metal was coordinated with octahedral geometry to five waters that were, in turn, coordinated by the carboxylate side chains of conserved residues in motifs I, III, and IV. The sixth ligand site in the catalytic metal complex was occupied by an ATP α phosphate oxygen, indicative of a role for the metal in stabilizing the transition state of the autoadenylylation reaction. A key insight, fortified by superposition of the Michaelis complex on the structure of the covalent NgrRnl-(Lys-\textsubscript{N\epsilon})-AMP intermediate, concerned the role of the catalytic metal complex in stabilizing the unprotonated state of the lysine nucleophile before catalysis, via local positive charge and atomic contact of Lys-\textsubscript{N\epsilon} to one of the metal-bound waters (7).

The NgrRnl Michaelis complex revealed a second metal, coordinated octahedrally to four waters and to ATP β and γ phosphate oxygens. The metal complex and the ATP γ phosphate were

Significance

This season marks the 50th anniversary of the discovery of polynucleotide ligases, the sine qua non enzymes of nucleic acid repair and the enabling reagents of molecular biology, reported in a series of seminal papers in PNAS. Ligases react with ATP or NAD\textsuperscript{+} to form a covalent enzyme–adenylate intermediate in which AMP is linked via a P–N bond to a lysine. This work reports crystal structures of the Michaelis complexes of an exemplary ATP-dependent RNA ligase (bacteriophage T4 Rnl1) and an NAD\textsuperscript{+}-dependent DNA ligase (Escherichia coli LigA) that illuminate the chemical and structural basis for lysine adenylylation, via distinctive two-metal (ATP) and one-metal (NAD\textsuperscript{+}) mechanisms.

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The authors declare no conflict of interest.

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Data deposition: The atomic coordinates have been deposited in the Protein Data Bank, www.pdb.org (PDB ID codes 5TT5 and 5TT6).

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engaged by an ensemble of amino acid side chains (unique to NgrRnl) that collectively orient the PP\_i leaving group apical to the lysine nucleophile. Consistent with a single-step in-line mechanism, the α phosphate was stereochemically inverted during the transition from NgrRnl-ATP Michaelis complex to lysyl–AMP intermediate (7).

NgrRnl exemplifies one of five distinct families of ATP-dependent RNA ligases that arose independently by fusion of an ancestral ATP-using NTase domain to structurally unique flanking domains (7). An important issue for ligase evolution is whether the two-metal mechanism and lysine adenylylation strategy exploited by NgrRnl—an Rnl5 family ligase that seals nicks in duplex RNAs (8)—applies to other ATP-dependent polynucleotide ligases. Here, we address the issue by capturing a structure of the Michaelis complex of bacteriophage T4 RNA ligase 1 (Rnl1). T4 Rnl1 was the first RNA ligase discovered (9, 10), and it spearheads a phage-encoded pathway of tRNA break repair that thwarts a tRNA-damaging host antiviral response (11). We find that T4 Rnl1 adheres to a two-metal mechanism of lysine adenylylation, driven by: (i) a catalytic metal–water complex that engages the lysine nucleophile and the ATP α phosphate and (ii) a second metal that orients the PP\_i leaving group, albeit in a ligase-assisted manner distinct from that seen for NgrRnl.

DNA ligases are thought to have evolved separately from RNA ligases, initially by fusion of an ancestral ATP-using NTase domain to a C-terminal OB domain to comprise the minimal catalytic core of a DNA ligase and, subsequently, via the fusion of additional structural modules to the NTase–OB core (7). NAD\(^+\)-dependent DNA ligases (LigA enzymes), which are ubiquitous in bacteria and essential for bacterial viability in all cases tested, acquired their specificity for NAD\(^+\) via fusion of an NMM-binding OB domain module to the N terminus of the NTase domain (4, 12, 13). *Escherichia coli* DNA ligase (EcoLigA) was the first cellular DNA ligase discovered and characterized (14–20), and it remains the premier model for structural and functional studies of the NAD\(^+\)-dependent DNA ligase family (21–27). Interest in LigA mechanism is propelled by the promise of targeting LigA (via its signature NAD\(^+\) substrate specificity and unique structural features vis à vis human DNA ligases) for antibacterial drug discovery (24, 28). Here, we report the structure of a Michaelis complex of EcoLigA with NAD\(^+\) and magnesium, which reveals a one-metal mechanism in which a catalytic metal–water complex engages the lysine nucleophile and the NAD\(^+\) α phosphate, whereas the NMM leaving group is positioned by enzymic contacts without agency of a second metal.

**Results**

A Michaelis Complex of T4 Rnl1 with ATP and Magnesium. The initial structure of T4 Rnl1 with the unreactive analog AMPCPP had a calcium ion in the putative catalytic metal site, coordinated to six waters and one AMPCPP α phosphate oxygen (29). Our aim here was to capture a mimetic of the Michaelis complex of T4 Rnl1 with ATP and its physiological metal cofactor magnesium. To accomplish this goal, we exploited a mutated version, K99M, in which the Lys99 nucleophile was replaced with methionine (effectively isosteric to lysine, minus the ε-aminogroup). Rnl1-K99M was preincubated with 1 mM ATP and 2 mM MgCl\(_2\) before crystallization. The refined 2.2-Å structure (Table S1) comprised the entire T4 Rnl1 polypeptide, composed of an N-terminal NTase domain (amino acids 1–254) and a C-terminal domain (amino acids 255–374) unique to Rnl1 (Fig. 1A). The NTase domain per se is capable of autoadenylylation and of “nonspecific” joining of RNA 3′-OH and 5′-PO\(_4\) ends; the distinctive C domain of T4 Rnl1 confers specificity for the repair of tRNAs with breaks in the anticodon loop (30). Electron density for ATP and two magnesium ions was evident in the active site (Figs. S1A and S2A). Fig. 1B shows a stereoview of the active site of the ATP(Mg\(^{2+}\))\(_2\) complex, highlighting atomic interactions relevant to catalysis. For comparison, we superimposed the Lys99 side chain from the T4 Rnl1-AMPCPP structure (29) on Met99 of the Michaelis complex. The modeled Lys99-N\(_\varepsilon\) is situated 3.1 Å from the ATP α phosphorus, in an apical orientation to the pyrophosphate leaving group (N\(_\varepsilon\)–O–O\(_3\)a angle = 169°).

Two-Metal Mechanism of T4 Rnl1 Lysine Adenylylation. The adenine nucleoside of ATP is in the syn conformation. The ribose 3′-OH and 2′-OH make hydrogen bonds to Arg54 and Glu159, respectively. The ATP α phosphate is engaged by a catalytic magnesium. Waters occupy five of the ligand sites in the octahedral Mg\(^{2+}\) complex. Rnl1 binds the Mg\(^{2+}\)(H\(_2\)O)\(_4\) complex via water-mediated contacts to Asp101 (motif I), Glu159 (motif III), Glu227 (motif IV), and Tyr246. The sixth Mg\(^{2+}\) ligand site is occupied by one of the nonbridging α phosphate oxygens (Fig. 1B). The other nonbridging α phosphate oxygen is contacted by Lys240 and Lys242 (motif V). The structure suggests that Lys240, Lys242, and Mg\(^{2+}\) stabilize a pentavalent transition state of the ATP α phosphate during the lysine adenylylation reaction. The structure also reveals how the catalytic Mg\(^{2+}\)(H\(_2\)O)\(_4\) complex stabilizes the unprotonated state of the Lys99 nucleophile before catalysis, via local positive charge and an atomic contact of Lys99-N\(_\varepsilon\) to one of the metal-bound waters (indicated by a yellow dashed line in Fig. 1B). The modeled

![Fig. 1. Structure of a Michaelis complex of T4 Rnl1 with ATP and magnesium. (A) Tertiary structure of T4 Rnl1-K99M, which consists of an N-terminal nucleotidyltransferase (NTase) domain (blue) and a unique C-terminal domain (beige). The ATP in the active site is rendered as a stick model. Mg\(^{2+}\) ions are depicted as magenta spheres. (B) Stereoview of the active site of Rnl1-(K99M)\_ATP(Mg\(^{2+}\))\(_2\) highlighting the catalytic Mg\(^{2+}\)(H\(_2\)O)\(_4\) coordination complex at the α phosphate, a second Mg\(^{2+}\) bridging the β and γ phosphates, and enzymic contacts to the ATP and metal complexes. Amino acids and ATP are shown as stick models with beige and gray carbons, respectively. Mg\(^{2+}\) ions and associated waters are depicted as magenta and red spheres, respectively. Atomic contacts are indicated by black dashed lines. The superimposed Lys99 side chain from the structure of wild-type Rnl1 (29) is shown with its proximity of N\(_\varepsilon\) to the ATP α phosphorus indicated by a magenta dashed line and to a metal-bound water and Glu227-O\(_3\)a indicated by yellow dashed lines.](image-url)
Lys99-Nε is also proximal to Glu227-Oγ, as seen in the T4 Rnl1-AMPCPP structure (29).

There are no direct enzymic contacts to the ATP β phosphate, only water-bridged interactions of one β phosphate oxygen with the catalytic Mg2+ and of the other β phosphate oxygen with Lys242. A second Mg2+ is coordinated octahedrally to three waters, to ATP β and γ phosphate oxygens, and to the Asp273 carboxylate (Fig. 1B). One of the second Mg2+-bound waters makes a bifurcated bridge to the ATP α phosphate and the Asp273 carboxylate. Direct contacts to the ATP γ phosphate are made by Tyr37, Arg54, Lys75, and Lys240. The Rnl1 residues that bind the γ phosphate and the Mg2+ complex with the β and γ phosphates together achieve a conformation of the ATP triphosphate moiety conducive to expulsion of the PPi-Mg2+ leaving group. The structure of the Michaelis complex accords with extensive mutational studies showing that Arg54, Lys75, Glu159, Glu227, Lys240, Lys242, and Tyr246 are essential for RNA ligase activity (31, 32).

When we align the previous Rnl1-AMPCPP structure (29) to that of our Michaelis complex with ATP (Fig. S3), we see that the AMP moieties and the amino acids (Asp101, Glu159, Glu227, Tyr246) that coordinate the waters in the catalytic metal complex superimpose nicely, but there is a difference in the trajectory of the β and γ phosphates of AMPCPP and ATP, whereby the β phosphorus atoms are offset by 1.8 Å and the γ phosphorus atoms are separated by 2.4 Å. This difference results in a 2.6-Å movement of the Tyr37-Oh that contacts the γ phosphate and a shift in the contact made by Lys75 from the γ phosphate of ATP to the β phosphate of AMPCPP (Fig. S3). We surmise that the bridging α-β methylene carbon in the ATP analog perturbs the geometry of the triphosphate moiety of the nucleotide. There are noteworthy differences in the atomic interactions of the PPi, leaving group and the noncatalytic Mg2+ in the AMPCPP structure versus the ATP structure. For example, in the AMPCPP structure, the noncatalytic Mg2+ makes direct contact with the β phosphate, but not with γ phosphate; rather, two of the waters coordinated by the second metal bridge to the γ phosphate (Fig. S3). Insofar as classic Mg2+ binding to an NTP directly engages the β and γ phosphate oxygens, we regard the ATP structure as reflective of the disposition of the second metal during the lysine adenyllylation reaction.

Michaelis Complex of E. coli LigA with NAD+ and Magnesium. The 671-aa EcoLigA protein consists of a catalytic core composed of an NTase domain (amino acids 70–316) and an OB domain (amino acids 317–404). These two modules are common to NAD+, and ATP-dependent DNA ligases. The LigA core is flanked by an N-terminal “Ia” domain (amino acids 1–69) that confers NAD+ specificity and by three C-terminal modules: tetra-cysteine Zn-finger (amino acids 405–432), helix-hairpin-helix (HhH) (amino acids 433–586), and BRCT (amino acids 587–671). Each step of the ligation pathway depends on a different subset of the LigA domains, with only the NTase domain being required for all steps. The 2.3-Å crystal structure of EcoLigA bound to a nicked DNA-adenylate intermediate (AppDNA) showed that the ligase encircles the DNA helix as a C-shaped protein clamp (24) (Fig. 2B and D). The LigA-DNA interface entails extensive DNA contacts by the NTase, OB, and HhH domains over a 19-bp segment of duplex DNA centered about the nick, many of which are essential for DNA ligase activity (25, 26).

The Ia and NTase domains suffice for LigA enzymes to react with NAD+ and magnesium to form the covalent ligase–AMP intermediate (13, 33, 34). The 2.7-Å crystal structure of the Ia-NTase segment of Enterococcus faecalis LigA in a complex with NAD+ (generated in crystalllo from AMP and NMN) provided the first insights to a bipartite mode of NAD+ recognition, whereby the AMP moiety of NAD+ is engaged by the NTase domain, whereas the NMP moiety is bound by the Ia domain (4). However, the EfaLigA Ia-NTase structure, devoid of metals, and with the lysine nucleophile making a bifurcated salt bridge to an aspartate (thus
enforcing the protonated state of lysine), left many issues unsettled anent catalysis of the lysine adenyllylation reaction of LigA enzymes.

Here, we aimed to capture a Michaelis complex of EcoLigA with NAD\(^+\) and magnesium, using a mutated version in which the Lys115 nucleophile was replaced by methionine. LigA-K115M was preincubated with 0.3 mM NAD\(^+\) and 5 mM MgCl\(_2\) before crystallization. The refined 1.55-Å structure (R\(_{\text{free}}\) = 0.177/0.205) (Table S1) spanned from Met1 to Pro586 (embracing the Ia, NTase, OB, Zn, and HhH domains; Fig. 2A) with a 5-Å gap between Leu102 and Val1108 in a surface loop of the NTase domain for which there was no interpretable electron density. Electron density for NAD\(^+\) and one magnesium ion was evident in the active site (Figs. S1B and S2B). No electron density was observed for the C-terminal 85-Å BRCT domain.

The arrangement of the component domains of the EcoLigA Michaelis complex is shown in Fig. 2A. The NTase domain forms a binding pocket for the NAD\(^+\) substrate akin to that seen in the EfaLigA Ia-NTase-NAD\(^+\) complex. The C-terminal OB, Zn, and HhH domains of the Eco-LigA Michaelis complex adopt an extended conformation distinct from any previous LigA structure. A superposition of the NTase domains of the EcoLigA-NAD\(^+\)-Mg\(^{2+}\) Michaelis complex (Fig. 2A) and the EcoLigA-DNA-adenylate complex (Fig. 2B; minus the nicked DNA) highlights the large movements of each of the flanking domains in the transition between these two functional states along the nick sealing pathway. For example: Met1 in the Ia domain moves 62 Å; Arg333 in the OB domain and the Zn\(^{2+}\) atom in the Zn-finger each move by 57 Å; and Pro586 at the end of the HhH domain moves 102 Å. Superposition of the HhH domains of the Michaelis complex (Fig. 2C) and the DNA-adenylate complex (Fig. 2D; with the nicked DNA) provides a different view of the massive conformation rearrangement between functional states of EcoLigA, entailing a 106-Å movement of the Met1 residue of the Ia domain.

One-Metal Mechanism of LigA Lysine Adenylylation. Fig. 3 shows a stereoview of the active site of the EcoLigA Michaelis complex, featuring atomic interactions relevant to NAD\(^+\) recognition and catalysis. For comparison, we superimposed on Met115 of the Michaelis complex the Lys115 side chain from the LigA-AppDNA structure. The Lys115-N\(_{\text{ε}}\) is poised 2.9 Å from the NAD\(^+\) α phosphor, in an apical orientation to the NMM leaving group (N\(_{\text{ε}}\)-Pα-Oα angle = 175°). The adenosine nucleoside of NAD\(^+\) is in the syn conformation. The ribose 3'-OH and 2'-OH make hydrogen bonds to Arg136 and Glu173, respectively. The NAD\(^+\) α phosphor is engaged by a catalytic Mg\(^{2+}\)(H\(_2\)O)\(_5\) complex. LigA binds the Mg\(^{2+}\)(H\(_2\)O)\(_5\) complex via water-mediated contacts to Asp117 (motif I), Glu173 (motif III), and Asp285 (motif IV). The sixth Mg\(^{2+}\) ligand site is filled by one of the nonbridging α phosphate oxygens (Fig. 1B). The other nonbridging α phosphate oxygen is contacted by Lys314 (motif V). The structure suggests that Lys314 and Mg\(^{2+}\) stabilize the pentavalent transition state of the NAD\(^+\) α phosphorus during the lysine adenyllylation reaction. The catalytic Mg\(^{2+}\)(H\(_2\)O)\(_5\) complex also stabilizes the unprotonated state of the Lys115 nucleophile before catalysis, via local positive charge and atomic contact of Lys115-N\(_{\text{ε}}\) to one of the metal-bound waters (indicated by the yellow dashed line in Fig. 3). The modeled Lys115-N\(_{\text{ε}}\) is also proximal to Asp285-O\(_{\text{ε}}\), as seen in the LigA-AppDNA structure (24).

The NAD\(^+\) β phosphor (i.e., the phosphor of the NN M leaving group) is coordinated directly on the same nonbridging oxygen by the Ser81 and Arg136 side chains of the EcoLigA NTase domain (Fig. 3). The other nonbridging β phosphate oxygen coordinates one of the Mg\(^{2+}\)-bound waters.

The nicotinamide nucleoside of NAD\(^+\) is in the anticonformation and is held in place solely by enzymic contacts to the Ia domain. Tyr22 and Tyr35 make a π sandwich above and below the nicotinamide ring (Fig. 3). The amide nitrogen of nicotinamide donates a hydrogen bond to Asp32. Asp36 accepts a hydrogen bond from the ribose 2'-OH of NMN, whereas the ribose 3'-OH coordinates one of the Mg\(^{2+}\)-bound waters (Fig. 3).

The atomic interactions of EcoLigA with NAD\(^+\) and Mg\(^{2+}\) in the Michaelis complex accord with mutational studies showing that domain Ia residues Tyr22, Asp32, Tyr35, and Asp36 and NTase domains residues Asp117, Arg136, Glu173, Asp285, and Lys314 are essential for DNA ligase activity (12, 22, 23, 26). Of the NTase residues, Arg136, which interacts with the β phosphor and adenosine ribose of NAD\(^+\), stands out as essential for the reaction of EcoLigA with NAD\(^+\) to form the covalent LigA-AMP intermediate (step 1), but dispensable for phosphodiester formation at a pre-adenylated nick (step 3) (26).

Discussion

Metal Catalysis of Lysine Adenylylation. The present study illuminates a conserved mechanism of lysine adenyllylation by ATP-dependent RNA ligases and NAD\(^+\)-dependent DNA ligases via a catalytic metal–(H\(_2\)O)\(_5\) complex that favors the unprotonated form of the lysine nucleophile and stabilizes the transition state of the ATP or NAD\(^+\) α phosphor. The shared structural basis for catalysis was gleaned from crystal structures of the Michaelis-like complexes of T4 Rnl1 (the exemplary Rnl1-family ligase), NgrRnl (Rnl5-family), and EcoLigA (the prototypical NAD\(^+\)-dependent DNA ligase), each of which was captured by replacing the motif I lysine nucleophile by an isosteric methionine. Modeling the lysine in Rnl1 and LigA (based on structure of the wild-type enzymes) confirmed mimesis of the Michaelis complex of the adenyllylation reaction in which the Lys-N\(_{\text{ε}}\) nucleophile and the PP\(_{\text{δ}}\) or NMM leaving groups are oriented apically for in-line catalysis. Superposition of the available Michaelis complexes shows that the positions and enzymic interactions of catalytic Me\(^{2+}\)(H\(_2\)O)\(_5\) complexes that engage the lysine nucleophile and α phosphor are virtually identical in the active sites of Rnl1, NgrRnl, and LigA (Fig. 4). Three carboxylate side chains of the signature nucleotideyltransferase motifs I, III, and IV bind the pentahydrated metal cofactor. A motif IV (EGYVA in Rnl1; DGVVI in LigA; EGLVF in NgrRnl) acidoic side chain coordinates two of the metal-bound waters, including the water that interacts with the lysine nucleophile of motif I (KEDG in Rnl1; KLDG in LigA; KLDG in NgrRnl). A motif I aspartate coordinates a third metal-bound water. A motif III glutamate makes bidentate
leaving group, but with each family of ATP-dependent ligase evolving its own set of enzymic contacts to the PP;Mg\(^{2+}\) complex. Although there are no structures available for a step 1 Michaelis complex of an ATP-dependent DNA ligase, functional studies of *Chlorella* virus DNA ligase suggest that conserved elements of the C-terminal OB domain are likely candidates to orient the PP, leaving group and coordinate a second metal (38, 39).

**Mechanistic Divergence in NAD\(^{+}\)-Dependent DNA Ligase.** The salient insights from the structure of the EcoLigA Michaelis complex are that whereas the catalytic metal is shared with ATP-dependent RNA ligases, the NAD\(^{+}\)-dependent DNA ligases eschew a second metal for leaving group orientation. Indeed, the nicotinamide nucleoside of NAD\(^{+}\) clashes sterically with the second metal site of ATP-dependent RNA ligases (Fig. 4). The metal-independent leaving group strategy of LigA is completely diverged from the ATP-dependent enzymes in two key respects. First, LigA makes essential contacts to the NAD\(^{+}\) β phosphate via an arginine of the (putatively ancestral) ATP-dependent DNA ligase lineage (7) with an NMM leaving group apical to the motif I lysine nucleophile. Thus, replacement of an PP;Mg\(^{2+}\) leaving group in the (putatively ancestral) ATP-dependent DNA ligase lineage (7) was presumably coupled during ligase evolution in bacteria to the abandonment of a second metal and the gain of a novel NMM-binding module.

NAD\(^{+}\)-dependent DNA ligases are promising targets for antibacterial drug discovery efforts, most of which have identified LigA inhibitors that occupy the adenosine site of the NTase domain and the nearby hydrophobic “C2 drug tunnel” unique to the LigA family (24, 28, 40–45). The structure of the LigA Michaelis complex, together with previous LigA structures (4, 24), makes a strong case for targeting small molecule inhibitors to the NMM site (composed of domain Ia and residues in the NTase domain). Such compounds might be especially effective when chemically linked to molecules that occupy the adenosine site. In addition, there would be value in identifying molecules that interfere with the various LigA domain movements that are coupled to progression through the LigA adenylylation and DNA binding steps of the ligation pathway.

**Methods**

**Crystallization of T4 Rnl1.** Mutant Rnl1-(K99M) was produced in *E. coli* BL21(DE3) as His\(_{6}\)Rnl1 fusion and isolated from a soluble bacterial extract by Ni-affinity chromatography as described (31). Rnl1-(K99M) was purified further by gel filtration through a column of Superdex-200 equilibrated in 20 mM Tris-HCl (pH 8.0), 100 mM NaCl. The peak fractions were pooled, concentrated by centrifugal ultrafiltration, and stored at −80 °C. A solution of 6.5 mg/mL Rnl1-(K99M), 1 mM ATP, and 2 mM MgCl\(_2\) was preincubated for 30 min on ice before mixture with an equal volume of reservoir solution containing 0.2 M MgCl\(_2\), 16% (vol/vol) PEG 3350. Crystals were grown at 22 °C by hanging drop vapor diffusion. Crystals appeared after 1–2 d. Single crystals were harvested, cryoprotected with 0.2 M MgCl\(_2\), 20% PEG 3350, 15% PEG 400, and then flash-frozen in liquid nitrogen.

**Crystallization of EcoLigA.** Mutant LigA-(K115M) was produced in *E. coli* BL21(DE3) as His\(_{6}\)EcoLigA fusion and isolated from a soluble bacterial extract by Ni-affinity chromatography as described (22). LigA-(K115M) was purified further by Superdex-200 gel filtration. The peak fractions were pooled, concentrated by centrifugal ultrafiltration, and stored at −80 °C. A solution of 20 mg/mL LigA-(K115M), 0.3 mM NAD\(^{+}\), and 5 mM MgCl\(_2\) was preincubated for 30 min on ice before mixture with an equal volume of reservoir solution containing 0.2 M Na acetate, 20%PEG 3350. Crystals were grown at 22 °C by sitting drop vapor diffusion. Crystals appeared after 10 d. Single crystals were harvested, cryoprotected with 0.2 M Na acetate, 25% PEG 3350, 15% PEG 400, and then flash-frozen in liquid nitrogen.

**Diffraction Data Collection and Structure Determination.** Diffraction data were collected continuously from single crystals at Argonne National Laboratory Advanced Photon Source beamlines 24ID-E for Rnl1-(K99M) and 24ID-C for Unculea et al.
LigA-K115M) at 100 K and processed with HKL-2000 (46). The structures were
solved by molecular replacement. The search probe for R1 loser (K99M) was 2.5 Å,
and phasing was performed with Phenix (47). For LigA-K115M, the search was
done successively with separate domains of E. coli LigA (Z沃O). NTase domain
(residues 72–315), HHN domain (residues 429–586), and OB together with Zn-
finger (residues 316–428) were found by using MOLREP (48). The N-terminal la
domain was built into difference electron density maps. Interactive model
building was done by using O (49). Both structures were refined with Phenix
(47). Data collection and refinement statistics are presented in Table S1.

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reaction and demonstration of an enzyme-adenylate complex with T4 bacteriophage-
LigA-K115M) at 100 K and processed with HKL-2000 (46). The structures were
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Characterization of an adenosine triphosphate-inorganic pyrophosphate exchange
reaction and demonstration of an enzyme-adenylate complex with T4 bacteriophage-
LigA-K115M) at 100 K and processed with HKL-2000 (46). The structures were
solved by molecular replacement. The search probe for R1 loser (K99M) was 2.5 Å,
and phasing was performed with Phenix (47). For LigA-K115M, the search was
done successively with separate domains of E. coli LigA (Z沃O). NTase domain
(residues 72–315), HHN domain (residues 429–586), and OB together with Zn-
finger (residues 316–428) were found by using MOLREP (48). The N-terminal la
domain was built into difference electron density maps. Interactive model
building was done by using O (49). Both structures were refined with Phenix
(47). Data collection and refinement statistics are presented in Table S1.

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