A second NAD\(^+\)-dependent DNA ligase (LigB) in *Escherichia coli*

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**ABSTRACT**

*Escherichia coli* DNA ligase (LigA) is the prototype of the NAD\(^+\)-dependent class of DNA ligases found in all bacteria. Here we report the characterization of *E. coli* LigB, a second NAD\(^+\)-dependent DNA ligase identified by virtue of its sequence similarity to LigA. LigB differs from LigA in that it lacks the BRCA1 C-terminus domain (BRCT) and two of the four Zn-binding cysteines that are present in LigA and all other bacterial NAD\(^+\) ligases. We found that recombinant LigB catalyzed strand joining on a singly-nicked DNA in the presence of a divalent cation and NAD\(^+\), and that LigB reacted with NAD\(^+\) to form a covalent ligase-adenylate intermediate. Alanine substitution for the motif I lysine (\(^{126}\)KxDG) abolished nick joining and ligase-adenylate formation by LigB, thus confirming that the ligase and adenylyltransferase activities are intrinsic to the LigB protein.

**INTRODUCTION**

NAD\(^+\)-dependent DNA ligase was first isolated from *Escherichia coli* in 1967 (1–4) and the *E. coli* enzyme has since served as the prototype for mechanistic studies of the bacterial ligase family (5). *Escherichia coli* DNA ligase (the product of the *ligA* gene) catalyzes the sealing of 5′ phosphate and 3′ hydroxyl termini at nicks in duplex DNA by means of three sequential nucleotidyl transfer reactions. In the first step, attack on the adenylyl α phosphorus of NAD\(^+\) by ligase results in release of nicotinamide mononucleotide and formation of a covalent intermediate (ligase-adenylate) in which AMP is linked through a phosphoamide bond to lysine. In the second step, the AMP is transferred to the 5′ phosphate at the nick to form DNA-adenylate. In the third step, the unadenylated ligase catalyzes attack by the 3′ OH of the nick on DNA-adenylate to join the two polynucleotides and release AMP.

An NAD\(^+\)-dependent DNA ligase is found in many bacterial species. NAD\(^+\)-dependent ligases are of fairly uniform size (656–837 amino acids) and extensive amino acid sequence conservation occurs throughout the entire lengths of the polypeptides. The atomic structures of NAD\(^+\)-dependent ligases from *Bacillus stearothermophilus* and *Thermus filiformis* have been determined by X-ray crystallography (6,7). Although there is scant amino acid sequence similarity between NAD\(^+\)-dependent and ATP-dependent ligases, the tertiary structures of the catalytic cores are conserved and the adenylylate binding pocket of NAD\(^+\) ligases is composed of the five nucleotidyl transferase motifs noted originally in the ATP-dependent enzymes (8–10). The catalytic core of the bacterial NAD\(^+\) ligase consists of nucleotidyl transferase and oligomer-binding fold (OB-fold) domains. The core is flanked by a short N-terminal domain (Ia) and three C-terminal domains: a tetracysteine domain that binds a single Zn atom; a helix–hairpin–helix domain (HhH) and a BRCT domain (named after the C-terminus of the breast cancer gene product BRCA1) (Fig. 1A).

Early genetic studies showed that the *ligA* gene is essential for growth of *E. coli* (11,12). Genes encoding NAD\(^+\)-dependent DNA ligase are also essential in *Salmonella typhimurium*, *Bacillus subtilis* and *Staphylococcus aureus* (13–15). Post-genomic analysis and biochemical studies have revealed that many bacteria contain one or more ATP-dependent DNA ligases in addition to an NAD\(^+\)-dependent enzyme (16,17). This discovery raises the prospect that bacteria, like eukaryotes, may exploit different DNA ligase isozymes for different physiological functions (e.g. replication, repair, homologous recombination and non-homologous end-joining). *Escherichia coli*, however, does not encode an ATP-dependent ligase. Rather, a search of the complete *E. coli* K12 genome with the LigA protein immediately highlights a second DNA ligase-like protein (Fig. 2), encoded by an open reading frame upstream of the gene for guanylate kinase (gmk) (GenBank accession no. AE000442).

The *E. coli* ligase-like protein (henceforth named LigB) is a 562 amino acid polypeptide. A protein of the same size with 97% sequence identity to LigB is encoded by the enterohemorrhagic *E. coli* strain O157:H7 (GenBank accession no. AE005592). A manual alignment of the LigB amino acid sequence to *E. coli* LigA and to the Tji and Bst ligases (guided by the crystal structure of the *Tji* ligase) reveals conservation of domain Ia, the nucleotidyl transferase domain (including the five catalytic motifs) and the OB-fold (Fig. 2) as well as the HhH domain (not shown). However, the LigB protein lacks the C-terminal BRCT domain and two of the four Zn-binding cysteines that are present in all other bacterial NAD\(^+\) ligases. Given that individual cysteines of the Zn finger have been shown to be essential for the nick-joining activity of bacterial ligases (18,19), and the hypothesis that the BRCT domain plays an important role in DNA binding (20), it is of interest to evaluate whether LigB is a DNA ligase and whether it uses NAD\(^+\) as a cofactor.

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Here we show that this is indeed the case, although LigB is considerably less active in nick joining than is LigA.

MATERIALS AND METHODS

Expression vectors for LigB

Oligodeoxyribonucleotide primers complementary to the 5′ and 3′ ends of the *E. coli* K12 *ligB* gene were used to amplify by PCR the equivalent open reading frame from *E. coli* DH5α. The primers were designed to introduce NdeI and *Bam*HI restriction sites at the 5′ and 3′ ends of the *ligB* gene. The PCR product was digested with NdeI and *Bam*HI, then cloned into the NdeI and *Bam*HI sites of pET16b (Novagen) to yield pET-LigB. Dideoxy sequencing of the entire insert of pET-LigB confirmed that no alterations of the DNA sequence were introduced during PCR amplification and cloning of the *ligB* gene. The K126A mutation was introduced into the *ligB* gene using the two stage PCR-based overlap extension method (21). An NdeI–*Bam*HI restriction fragment of the mutated second stage PCR product was inserted into pET16b. The insert of the resulting pET-LigB-K126A plasmid was sequenced to confirm the presence of the desired alanine mutation and the absence of any unwanted coding changes.

Purification of LigB

The pET-LigB expression plasmid was transformed into *E. coli* BL21(DE3). A single ampicillin-resistant colony was inoculated into LB medium containing 0.1 mg/ml ampicillin and a 100 ml culture was grown at 37°C until the A₆₀₀ reached 0.8. The culture was placed on ice for 30 min, then adjusted to 0.4 °C–D-thiogalactopyranoside (IPTG), and subsequently incubated at 17°C for 16 h with continuous shaking. Cells were harvested by centrifugation and the pellets were stored at –80°C. All subsequent procedures were performed at 4°C. Cell lysis was achieved by treatment of thawed, resuspended cells with 0.2 mg/ml lysozyme and 0.1% Triton X-100 in 10 ml of lysis buffer containing 50 mM Tris–HCl (pH 7.5), 0.5 M NaCl and 10% sucrose. The lysate was sonicated to reduce viscosity and insoluble material was removed by centrifugation at 37 000 g for 20 min. The supernatant was mixed with...
1 ml of Ni-NTA-agarose resin (Qiagen) for 30 min with constant rotation. The slurry was poured into columns and the packed resin was washed with IMAC buffer (50 mM Tris–HCl (pH 7.5), 50 mM NaCl, 10% glycerol) containing 5 mM imidazole. The column was step-eluted with 50 and 500 mM imidazole in IMAC buffer. The polypeptide compositions of the column fractions were monitored by SDS–PAGE. The Histagged LigB was recovered in the 500 mM imidazole eluate (0.9 mg of protein). The preparation was diluted 1:1 with buffer P containing 600 mM NaCl in buffer P (50 mM Tris–HCl (pH 7.5), 10% glycerol) and applied to a 1 ml phosphocellulose column that had been equilibrated with buffer P. The column was washed with 200 mM NaCl in buffer P and LigB was step-eluted with 600 mM NaCl in buffer P containing 1 mM EDTA. The phosphocellulose ligase preparation (0.6 mg of protein) was stored at –80°C. The protein concentrations were determined by using the Bio-Rad dye reagent with bovine serum albumin as the standard. The K126A mutant was produced and purified from a 100 ml bacterial culture using the same protocol.

RESULTS

Purification of E.coli LigB and demonstration of ligase activity

The ligB gene was cloned into a T7 RNA polymerase-based bacterial expression vector so as to fuse the 562 amino acid LigB protein to a 20 amino acid N-terminal leader peptide containing 10 tandem histidines. The expression plasmid was introduced into E.coli BL21(DE3), a strain that contains the T7 RNA polymerase gene under the control of a lacUV5 promoter. The recombinant His-tagged protein was purified from a soluble extract of IPTG-induced bacteria by nickel-agarose affinity chromatography and phosphocellulose cation exchange chromatography steps. SDS–PAGE analysis showed that the phosphocellulose preparation was highly enriched with respect to the 64 kDa LigB polypeptide (Fig. 1B).

We assayed the ability of the recombinant LigB protein to seal a duplex DNA substrate containing a single nick (Fig. 1C). NAD+ and magnesium were included in the assay mixtures. Activity was evinced by conversion of the 5′-32P-labeled 18mer substrate to 36mer product (Fig. 1C). Thus LigB is indeed a DNA ligase.

The initial step in DNA ligation involves formation of a covalent enzyme-adenylate intermediate. In order to assay adenyllyltransferase activity, we incubated the recombinant LigB protein with [32P-AMP]NAD+ and magnesium. This resulted in the formation of a 5′-labeled covalent nucleotidyl-protein adduct that comigrated with the LigB polypeptide during SDS–PAGE (Fig. 1D, WT). We conclude that LigB is active in covalent nucleotidyl transfer with NAD+ as the AMP donor.

Alanine mutation of the motif I lysine of LigB

The KxDG sequence (motif I) is the signature feature of the ligase/capping enzyme superfamily of nucleotidyl transferases that form a covalent lysyl-NMP intermediate (8). To gauge the contribution of the motif I lysine residue (Lys126) to the activity of LigB we replaced it with alanine. The mutant protein K126A was isolated from soluble lysates by nickel-affinity and phosphocellulose chromatography; its purity was comparable to that of wild-type LigB (Fig. 1B). K126A was inert in nick ligation (Fig. 1C) and in ligase-AMP formation with [32P-AMP]NAD+ (Fig. 1D). This result is consistent with Lys126 being the site of covalent AMP attachment, as it is in other poly(nucleotide) ligases, and it excludes the possibility that the observed nick-joining activity of the recombinant LigB protein might be attributable to contamination by the endogenous LigA.

Characterization of LigB

Initial experiments were performed to determine the optimal conditions for nick joining. LigB was active in Tris–HCl buffer from pH 6.5 to 9.0 (not shown). Nick joining by LigB depended on a divalent cation cofactor; this requirement was satisfied by either magnesium (1–10 mM) or manganese (1–10 mM) (data not shown). A comparison of various divalent cations at 5 mM concentration showed that neither calcium nor zinc supported LigB activity and that cobalt was about one-quarter as effective as magnesium (not shown).

The LigB dependence of nick joining is shown in Figure 3A. Activity in the absence of added NAD+ was attributed to pre-adenylated LigB in the enzyme preparation. The linear dependence of nucleotide-independent strand joining on input enzyme suggested that 14% of the enzyme molecules had AMP bound at the active site. Inclusion of NAD+ in the reaction mixture stimulated nick ligation 3-fold. Inclusion of ATP failed to stimulate the joining reaction above the level achieved in the absence of added nucleotide. Indeed, none of the eight standard rNTPs or dNTPs were able to stimulate LigB (not shown). NADP was also inactive (not shown). NAD+ titration experiments showed that nick-joining activity was equivalent from 7 to 60 µM NAD+ and that higher NAD+ concentrations (>100 µM) were inhibitory. At ≥0.5 mM NAD+ LigB activity was reduced to the basal level seen in the absence of added nucleotide. Inhibition
Figure 4. Reaction of LigB with nicked DNA-adenylate. A reaction mixture containing (per 20 µl) 50 mM Tris–HCl (pH 7.5), 10 mM (NH₄)₂SO₄, 5 mM DTT, 5 mM MgCl₂, 250 fmol of ³²P-labeled nicked DNA-adenylate and 2 pmol of LigB was incubated at 37°C. The reaction was initiated by the addition of LigB. Aliquots (20 µl) were withdrawn at the times specified and quenched immediately. The extent of ligation is plotted as a function of time. The nicked DNA-adenylate substrate used in the ligation reaction is illustrated at the top.

by 0.5–1 mM NAD⁺ was also observed for E.coli LigA (not shown).

The kinetics of nick joining were examined under conditions of LigB excess (400 nM LigB and 50 nM nicked DNA). The reaction rates and extents were virtually identical with or without added NAD⁺ (Fig. 3B). The reaction without NAD⁺ reflects a single turnover of nick joining by pre-formed LigB-AMP intermediate (present at ∼56 nM concentration in the reactions). An endpoint was attained in 2–5 min at which time 80% of the input DNA substrate was sealed. We calculated an apparent rate constant of 0.026 s⁻¹. LigB also catalyzed phosphodiester bond formation on a pre-adenylated DNA substrate (Fig. 4). Under conditions of enzyme excess (100 nM LigB and 12.5 nM nicked DNA-adenylate) LigB sealed 65% of the input DNA-adenylate in 10 min. The apparent rate constant for sealing of nicked DNA-adenylate was 0.008 s⁻¹. The step 1 reaction of LigB with 1 µM [³²P-AMP]NAD⁺ displayed pseudo-first order kinetics with an apparent rate constant of 0.022 s⁻¹ (not shown).

DISCUSSION

We have demonstrated that E.coli LigA encodes two NAD⁺-dependent DNA ligases, LigB lacks the tetracysteine Zn finger and the BRCT structural domains that are present in LigA and all other bacterial NAD⁺ ligases. In this respect LigB resembles the recently identified NAD⁺-dependent ligase of Ansmacta moorei entomopoxvirus, a eukaryotic poxvirus (22). The AmEPV ligase, its homolog from Melanoplus sanguinipes EPV, and E.coli LigB appear to comprise a new branch of the NAD⁺ ligase family.

A direct comparison of the biochemical properties of purified recombinant versions of the two E.coli DNA ligases using the same nicked DNA substrate and similar reaction conditions (19, the present study, and data not shown) indicates that LigA is much more active than LigB. The specific activity of LigB under steady-state conditions is only 1% that of LigA. The activity difference is attributable to the differential response to the NAD⁺ cofactor. Whereas LigA is stimulated ≥100-fold by NAD⁺ under steady-state conditions, we find that LigB is stimulated only 3-fold.

The rate of single-turnover nick joining by LigB-adenylate (0.026 s⁻¹) is much slower than that of LigA-adenylate (≥0.22 s⁻¹). Thus, it appears that LigB is impaired in one or more of the component steps of the ligase pathway subsequent to adenyla-
tion of the enzyme, either in the binding of ligase-adenylate to the nicked DNA, transfer of AMP from LigB to the 5’ PO₄ at the nick (step 2 chemistry), attack of the 3’ OH on the DNA-adenylate strand to form a phosphodiester (step 3 chemistry), or release of the AMP product. As we did not detect accumulation of DNA-adenylate during the single turn-
over reaction of LigB with nicked DNA, we presume that step 3 chemistry is not rate limiting. The apparent rate constants for step 3 under single turnover conditions were fairly similar for LigA (0.023 s⁻¹) and LigB (0.008 s⁻¹). As we have discussed previously (23), the isolated step 3 reaction can be para-
doxically slower than the overall ligase reaction (this is the case for LigA and LigB) and we have invoked rate-limiting conformational steps in the binding of nicked DNA-adenylate to the ligase active site that do not apply during the standard nick joining reaction. The rate of the ligase-adenylation for LigB (kobs ≥0.022 s⁻¹) was much slower than that for LigA (kobs, ≥0.4 s⁻¹).

Our comparison of LigB and LigA suggests that LigB is less active than LigA at multiple stages of the ligase joining pathway. Moreover, we find that whereas LigA can functionally substitute for the essential Cdc9 ligase in yeast, LigB cannot do so (19 and data not shown). The simple interpretation is that the intrinsic activity of LigB activity is too weak to seal the Okazaki fragments generated during DNA replication. Can the functional differences between LigA and LigB be attributed to the structural domains that are missing from LigB? In an attempt to answer this question, we performed a domain swap in which the entire C-terminal OB-fold, zinc finger, HHH and BRCT portion of LigA (residues 315–671) was fused to the N-terminal domain Ia/nucleotidyl transferase portion of LigB (residues 1–304). However, the LigB–LigA chimera was no more active in nick joining than LigB (not shown). A second domain swap in which just the BRCT domain of LigA (residues 586–671) was fused to the C-terminus of LigB also failed to elicit a gain-of-function compared with LigB (not shown). Thus, we are unable by this strategy to engineer a more active version of LigB.

It is conceivable that LigB activity in nick joining is stimulated by other bacterial proteins, just as eukaryotic DNA ligase IV is stimulated by its binding partner XRCC4 (24). It is also possible that the function of LigB activity is not geared toward Okazaki fragment sealing, but rather may be responsible for sealing Okazaki fragments generated during DNA replication. Can the functional differences between LigA and LigB be attributed to the structural domains that are missing from LigB? In an attempt to answer this question, we performed a domain swap in which the entire C-terminal OB-fold, zinc finger, HHH and BRCT portion of LigA (residues 315–671) was fused to the N-terminal domain Ia/nucleotidyl transferase portion of LigB (residues 1–304). However, the LigB–LigA chimera was no more active in nick joining than LigB (not shown). A second domain swap in which just the BRCT domain of LigA (residues 586–671) was fused to the C-terminus of LigB also failed to elicit a gain-of-function compared with LigB (not shown). Thus, we are unable by this strategy to engineer a more active version of LigB.

Finally, it is worth noting that LigB is not restricted to E.coli. The genome of Yersinia pestis, the causative agent of plague, appears to encode two NAD⁺-dependent ligases (25). One Yersinia ligase is a 670 amino acid homolog of E.coli LigA and the second is a 562 amino acid polypeptide with 43% side-chain identity to E.coli LigB. Just as in E.coli, the Yersinia ligB gene
is located adjacent to the gene for guanylate kinase. As more and more bacterial genomes are sequenced, it will be of interest to see how many species other than *E. coli* and *Yersinia* have multiple NAD+-dependent ligases.

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**REFERENCES**