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Structural and functional insight into the mechanism of an alkaline exonuclease from *Laribacter hongkongensis*

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ABSTRACT

Alkaline exonuclease and single-strand DNA (ssDNA) annealing proteins (SSAPs) are key components of DNA recombination and repair systems within many prokaryotes, bacteriophages and virus-like genetic elements. The recently sequenced β-proteobacterium *Laribacter hongkongensis* (strain HLHK9) encodes putative homologs of alkaline exonuclease (LHK-Exo) and SSAP (LHK-Bet) proteins on its 3.17 Mb genome. Here, we report the biophysical, biochemical and structural characterization of recombinant LHK-Exo protein. LHK-Exo digests linear double-stranded DNA molecules from their 5'-termini in a highly processive manner. Exonuclease activities are optimum at pH 8.2 and essentially require Mg2+ or Mn2+ ions. 5'-phosphorylated DNA substrates are preferred over dephosphorylated ones. The crystal structure of LHK-Exo was resolved to 1.9 Å, revealing a ‘doughnut-shaped’ toroidal trimeric arrangement with a central tapered channel, analogous to that of λ-exonuclease (Exo) from bacteriophage-λ. Active sites containing two bound Mg2+ ions on each of the three monomers were located in clefts exposed to this central channel. Crystal structures of LHK-Exo in complex with dAMP and ssDNA were determined to elucidate the structural basis for substrate recognition and binding. Through structure-guided mutational analysis, we discuss the roles played by various active site residues. A conserved two metal ion catalytic mechanism is proposed for this class of alkaline exonucleases.

INTRODUCTION

The accurate repair of broken or damaged double-stranded DNA (dsDNA) molecules by homologous recombination (HR) is a process common to all living organisms (1,2). The process of HR involves genetic exchange (crossover) between two dsDNA molecules via stretches of DNA that share significant sequence homology. In prokaryotes, it plays pivotal roles in chromosome repair, replication and segregation (3,4). HR also promotes horizontal gene transfer between prokaryotes and resident plasmids, bacteriophages or assorted mobile genetic elements (5). Consequently, many prokaryotes have acquired HR-promoting proteins of viral/phage origin in addition to their ‘endogenous’ DNA recombinational-repair machinery (6).

Two-component HR-promoting proteins of viral/phage origin, comprising an alkaline exonuclease and a single-stranded DNA (ssDNA) annealing protein [SSAP, also referred to as a synaptase (7)] are widely distributed throughout the prokaryotic world (6,8–10). In these systems, the alkaline exonuclease protein fully digests one strand, or partially digests both strands of the linearized dsDNA substrate; generating ssDNA molecules, or long ssDNA ‘tails’ several hundred nucleotides...
in length. These are ‘coated’ with the partnering SSAP protein, which promotes their annealing with complementary regions of ssDNA on the chromosome or episome (11–13). This most likely occurs at the replication fork (14–16). DNA repair and replication proteins process the genetic intermediates formed, resulting in the creation of fully double-stranded recombinant DNA molecules (6,7,14). The best studied of the bacterial-based, two-component exonuclease and SSAP systems are the ‘Red’ proteins Exo (λ-exonuclease) and Bet from bacteriophage-λ (7,17); RecE and RecT from the Escherichia coli Rac prophage (7,12,18,19); G34.1P and G35P from bacteriophage SPP1 (13,20,21); gp60 and gp61 from Mycobacteriophage Che9e (22,23); and SXT-Exo (s066) and SXT-Bet (s065) from the SXT genetic element of Vibrio cholerae (24–26). Over recent years, various partnering pairs of exonuclease and SSAP proteins have been utilized in ‘recombineering’ procedures to create genetic alterations within a variety of prokaryotic organisms; including E. coli, Salmonella, Pseudomonas and Mycobacteria (22,27–29). In these systems, the exonuclease and SSAP proteins mediate HR-type events between PCR-generated dsDNA ‘targeting cassettes’ (encoding the desired genetic change) and the bacterial chromosome or resident episome (14).

First isolated and characterized in the 1960s (30,31) (and references therein), the activities and biophysical properties of the λ-exonuclease protein have been extensively studied. Phylogenetically related to exonucleases from the Baculoviridae and Herpesviridae families, it is the prototypical member of the alkaline exonuclease family (8), which belongs to the restriction endonuclease-like PD-(D/E)XK superfamily (32). Its activities are optimal at pH 9.2–9.5, and are dependent upon Mg2+ concentrations; although Mn2+ can substitute with a 3- to 4-fold reduction in catalytic efficiency (31). λ-exonuclease binds to the termini of linear dsDNA molecules, and processively digests the 5’-ended strand to produce long 3’-ssDNA tails, with the concomitant liberation of 5’-mononucleotides. It has previously been estimated that an average of 3000 (33) or 18 000 (34) nucleotides are digested without dissociation of the enzyme from the undigested strand. It has a strong preference for dsDNA molecules containing 5’-phosphorylated (5’-PO4) ends, digesting them an estimated 2- to 100-fold more effectively than the corresponding dephosphorylated (5’-OH) substrates (35–37). Its average digestion rate has been estimated to be ~4 nt/s (35), 9 nt/s (38), 15–20 nt/s (39), 32 nt/s (34) or ~1000 nt/s (40), using a variety of bulk or single-molecule scale biophysical approaches.

The crystal structure of λ-exonuclease was reported in 1997 (41), revealing an unusual toroidal (doughnut-shaped) trimeric arrangement with a central tapered channel. This funnel-shaped channel had an inner diameter of ~30 Å at one end, which is wide enough to accommodate a dsDNA substrate. However, it was only ~15 Å wide at the narrow end, which is only capable of accommodating ssDNA. Each monomer contained a Mn2+ ion housed within a pocket lining the channel, which presumably equate to the active sites that normally accommodate catalytic Mg2+ ions. This led the authors to propose a mechanism whereby the dsDNA substrate enters at the wide opening, and the non-hydrolyzed 3’-strand exits at the opposite end.

The crystal structure of the catalytically active C-terminal domain of the functionally equivalent RecE exonuclease was recently reported (38). RecE formed an analogous toroidal tetrameric arrangement, with a central tapered channel that contained the catalytic residues. Although λ-exonuclease and RecE share little overall primary sequence similarity and have quite distinct multimeric arrangements, the central channels within these two proteins have a highly similar size and topology. This strongly suggests a conserved mode of action where the terminus of the dsDNA molecule enters at the wide end of the channel; the strands are separated, and the 5’-end is processively hydrolyzed as the protein encircles and translocates along the undigested 3’-strand (38,41).

As part of our ongoing efforts to characterize prokaryotic DNA recombination proteins, we identified two adjacent genes within a suspected prophage region on the genome of Laribacter hongkongensis HLHK9, which encode homologs of alkaline exonuclease (LHK-Exo, LHK_01497) and SSAP (LHK-Bet, LHK_01496) proteins (42). Laribacter hongkongensis is a recently identified species of β-proteobacteria that is a potential diarrheal pathogen (43,44). Here, we describe the functional and structural characterization of LHK-Exo, which shares 29% primary amino acid sequence identity with λ-exonuclease. The crystal structure of LHK-Exo was solved to 1.9 Å resolution, revealing a ‘doughnut-like’ toroidal structure, similar to that previously observed for λ-exonuclease (41). LHK-Exo functioned as a DNA exonuclease, digesting linear dsDNA with a clear preference for 5’-phosphorylated ends. Under the guidance of the structure of LHK-Exo in complex with DNA, a set of active site point mutants were constructed and functionally characterized to probe sequence–structure relationships. We compare the activities and properties of LHK-Exo with those of related nucleases; and discuss mechanisms for substrate recognition and binding, as well as a potential catalytic mechanism for this class of alkaline exonucleases.

**MATERIALS AND METHODS**

**Materials**

All buffer salts, metal salts, chemicals and biochemicals were purchased from Sigma-Aldrich or GE Biosciences. Picogreen reagent was from Invitrogen (cat.# P7589). Unmodified and phosphorothioate modified oligonucleotides were purchased from TechDragon Ltd. (Shatin, Hong Kong). Where applicable, oligonucleotides were 5’-phosphorylated using T4 polynucleotide kinase (NEB), according to the manufacturer’s instructions, then desalted on Auto Seq G50 columns (GE Biosciences). All restriction enzymes were purchased from New England Biolabs (NEB; Ipswich, MA, USA). pET28a and pET32a plasmid DNA were from Novagen; pUC18 was from Stratagene and pMAL-c2 was purchased...
from New England Biolabs. Restricted (linearized) dsDNA was routinely purified using a Qiaquick Gel Extraction kit (Qiagen GmbH). When applicable, linearized dsDNA was 5'-dephosphorylated using calf intestinal phosphatase (NEB) then gel purified. dsDNA was resolved by electrophoresis on agarose/Tris–acetate EDTA (TAE) gels and stained with ethidium bromide (Sigma). ssDNA was analyzed by denaturing polyacrylamide gel electrophoresis on 7 M urea–TBE polyacrylamide (12%, 37.5:1) gels, which were stained using SYBR Gold Nucleic Acid Gel Stain (Invitrogen). Gels were visualized by UV-transillumination on a ChemiDoc XRS molecular imaging system equipped with Quantity One v4.6.6 software (BioRad). A 1 kb Plus DNA Ladder (Invitrogen) was used as a marker for dsDNA, and Oligo Length Standards 20/100 Ladder (Integrated DNA Technologies Inc., USA) was used for ssDNA gels. Graphs were prepared using SigmaPlot 10.0 (Systat Software, Inc).

Methods

Cloning, expression and purification of LHK-Exo. The LHK-exo gene (LHK_01497; GenBank: NC_012559.1, 1455161..1455772) was PCR amplified from Laribacter hongkongensis HLHK9 genomic DNA (Expand Polymerase, Roche) using the LHkexofor1 (ATATCCAGAAACTCCAC) and LHKexorev2 (TAAATATGCTCAGSCSGSG) primers. The amplified gene product was ligated via NcoI/HindIII (restriction sites underlined) into pET28a (Novagen) to create gene (LHK_01497; GenBank: NC_012559.1, 1455161..1455772) was PCR amplified from Laribacter hongkongensis HLHK9 genomic DNA (Expand Polymerase, Roche) using the LHkexofor1 (ATATCCAGAAACTCCAC) and LHKexorev2 (TAAATATGCTCAGSCSGSG) primers. The amplified gene product was ligated via NcoI/HindIII (restriction sites underlined) into pET28a (Novagen) to create plasmid pEXNH1, which encodes a 13aa C-terminal hexa-histidine fusion (...KLAAALEHHHHH*).

Escherichia coli BL21 (DE3) containing pEXNH1 was grown in Luria-Bertani (LB) medium (2 l) containing 50 μg/ml kanamycin (37°C), to an OD600 of 0.6–0.8, induced with IPTG (isopropyl-β-D-thiogalactopyranoside, 0.2 mM), then incubated at 25°C for 10 h. Bacterial cells were harvested by centrifugation (6000 g, 30 min), washed with PBS buffer, then resuspended in PBS buffer containing protease inhibitors (Complete, Roche) and lysed by sonication (ice cooling). The supernatant obtained after centrifugation (15 000 g, 30 min) was filtered (0.45 μm syringe filter, Millipore) then applied to a Ni²⁺-charged 5 ml Hitrap Chelating FF column (GE Healthcare) pre-equilibrated with PBS buffer. The column was washed (PBS buffer containing 10 mM imidazole, 20 μl) and the protein was eluted from the column using a linear gradient of imidazole (25–500 mM) in PBS buffer. Fractions containing the purest protein were pooled and concentrated (Amicon Ultra, Millipore) and buffer was exchanged using a G-25 Sephadex desalting column (GE Healthcare) pre-equilibrated with buffer A (20 mM Tris–HCl pH 8.0, 1 mM DTT). Protein was then purified by anion exchange chromatography (Resource Q, GE Healthcare), eluting with a linear gradient of 0–1 M NaCl in buffer A. Fractions containing the purest recombinant LHK-Exo protein were pooled then subjected to gel filtration chromatography (Superdex-200 HR, GE Healthcare) using buffer B (20 mM Tris pH 8.0, 150 mM NaCl, 1 mM DTT). The purity of resultant LHK-Exo was greater than ~98% as determined by SDS–PAGE analysis. Single residue mutants of LHK-Exo were constructed using the QuikChange Site-Directed Mutagenesis kit (Stratagene) according to the manufacturer’s instructions; using plasmid pEXNH1 as the template and the sets of oligonucleotides listed in Supplementary Table S1. Mutant proteins were expressed and purified as described above.

Qualitative determination of LHK-Exo activities

Mode of digestion. BamHI-linearized pET28a (1.8 μg, 0.54 pmol) was incubated with LHK-Exo protein (30 μg, 0.41 nmol of trimers) in 200 μl of Tris–HCl (pH 8.0, 50 mM), 50 mM NaCl, 7.5 mM MgCl2 at 37°C. Aliquots (20 μl) were withdrawn at various time points (indicated in the text) and analyzed on a 1% agarose/TAE gel.

Polarity of digestion. The 712-bp dsDNA substrates (‘unmodified’) were synthesized by PCR using pET28a as template, with two different pairs of primers, one pair of T7 (TAATACGACTCACTATAGGG) and T7rev (GCTAGTTATGGTCTAGCGG) for amplification of ‘unmodified’ substrate, and another pair of the phosphorothioate modified primers T7forPT3 (TAATACGACTCATA5G5G) and T7revPT3 (GCTAGTTATGGTCTAGC5G5G) for ‘PT-modified’ substrate (where subscript ‘S’ denotes a nuclease-resistant phosphorothioate linkage between 2 nt). Both PCR-amplified substrates were 5'-phosphorylated (T4 polynucleotide kinase, NEB) and purified (QIAquick PCR purification kit, QIAGen) prior to use. The two dsDNA substrates (0.1 μg, 0.23 pmol) were separately incubated with 6 μg of lambda-Exo (74 pmol of trimers) or LHK-Exo (82 pmol of trimers) in Tris–HCl (25 mM, pH 8.0), 7.5 mM MgCl2, 1 mM DTT (total volume 40 μl) at 37°C. Aliquots (20 μl) were quenched (20 mM EDTA + 1% SDS) immediately and after 20 min, then analyzed on 1% agarose TAE gels.

Single strand DNA exonuclease activities. Phosphorylated [5'-PO4-(dT)50] or non-phosphorylated [5'-OH-(dT)50] forms of a 50-mer or oligothymidine (0.4 nmol) were incubated with LHK-Exo protein (4.5 μg, 61.4 pmol of trimers) in 25 mM Tris–HCl, pH 8.0; 7.5 mM MgCl2; 1 mM DTT at 37°C; total volume 80 μl. Aliquots (20 μl) were removed at 0, 0.5, 5 and 20 min, and immediately quenched (30% glycerol, 7 M urea–TBE; 80 μl), prior to analysis by denaturing polyacrylamide gel electrophoresis.

Quantitative determination of DNA exonuclease activities. Quenched PicoGreen DNA fluorescence assays were performed essentially as described previously (21,26,45) with minor modifications. Typical assays (60 μl) contained LHK-Exo protein (8 ng, 0.11 pmol of trimers) and 30 ng linear pUC18 DNA (0.018 pmol) or EcoRV-linearized pMAL-c2 (0.0073 pmol) in nuclease assay buffer [Tris–HCl (25 mM, pH 8.0), 7.5 mM MgCl2, 1 mM DTT]. Assays were initiated by the addition of linear dsDNA substrate, incubated at 25°C for 20 min before being quenched by the addition of EDTA to 20 mM. The 250-fold diluted PicoGreen reagent

(Invitrogen) was added to quenched reaction mixtures and fluorescence levels (which are proportional to the quantities of dsDNA present) were measured at an excitation 485 nm/emission 535 nm (Perkin Elmer 1420 multi-label counter).

**Determination of optimal metal ion concentrations, pH and temperature for DNA exonuclease activities.** Quenched PicoGreen fluorescence assays were performed, quenched and analyzed as described above with minor modifications.

**Determination of optimal Mg\(^{2+}\) and Mn\(^{2+}\) ion concentrations.** Assays (60 μl) contained LHK-Exo protein (8 ng, 0.11 pmol of trimers), PstI-linearized pUC18 (30 ng, 0.018 pmol) in Tris–HCl (25 mM, pH 8.0), MgCl\(_2\) (1–15 mM) or MnCl\(_2\) (0.1–1.7 mM) at the concentrations indicated in the text; and were incubated at 25°C for 20 min.

**Determination of optimal pH.** Assays (60 μl) contained LHK-Exo protein (8 ng, 0.11 pmol of trimers), PstI-linearized pUC18 (30 ng, 0.018 pmol), 25 mM Tris–HCl, 7.5 mM MgCl\(_2\); adjusted to the appropriate pH value (pH 7.0–9.0); and were incubated at 25°C for 20 min.

**Determination of optimal temperature.** Assays (60 μl) contained LHK-Exo protein (40 ng, 0.55 pmol of trimers) and PstI-linearized pUC18 (30 ng, 0.018 pmol) in nuclease assay buffer. All solutions were pre-equilibrated at the temperature indicated in the text (34–54°C) using a thermostat-regulated water bath; an incubation time of 1 min was used. Four to six independent replicates of each experimental condition were performed. Data were fitted to hyperbolae using SigmaPlot 10.0 to calculate the \(k_{\text{cat}}\) during the initial phase or reaction (digestion rate stated in nucleotides per second; per trimer of LHK-Exo protein), reported as the mean ± standard deviation.

**Determination of digestion processivity.** LHK-Exo (6 μg, 82 pmol of trimers) protein and 60 ng (0.015 pmol) of linear dsDNA substrate (5'-PO\(_4\)-dDNA, EcoRV-linearized pMal-c2; or 5'-OH-dsDNA, 5'-dephosphorylated EcoRV-linearized pMal-c2) were equilibrated in 180 μl of Tris–HCl (25 mM, pH 8.0), 1 mM DTT, at 25°C for 5 min. A total of 90 μl of Tris–HCl (25 mM, pH 8.0), 1 mM DTT 15 mM MgCl\(_2\) was added to initiate the reaction; then exactly 15 s later, excess heparin (Sigma; 360 μg, 20 nmol) in 90 μl of Tris–HCl (25 mM, pH 8.0), 1 mM DTT, 7.5 mM MgCl\(_2\) was added to sequester unbound LHK-Exo (total reaction volume 360 μl). Assays were incubated at 25°C for 20 min, removing 60 μl aliquots at the times indicated in the text to determine dsDNA levels via fluorescent PicoGreen assays (as described above). Four to six independent replicates of each experimental condition were performed. Plotted data were fitted to hyperbolae using SigmaPlot 10.0 to obtain the digestion processivity from the extrapolated curves as previously described (26,36), which is reported as the mean ± standard deviation.

**Analysis of multimericity by analytical ultracentrifugation.** Sedimentation velocity analytical ultracentrifugation experiments were performed using a Beckman ProteomeLab™ XL-I analytical ultracentrifuge (60 000 rpm, 262,000 g, 293 K). LHK-Exo protein was prepared at a concentration of 1 mg/ml in 150 mM NaCl. Data were analyzed using the c(S) and c(M) models in the Sedfit program (46) to determine the sedimentation coefficient and apparent mass distributions, respectively.

**Crystallization and X-ray data collection.** The purified recombinant LHK-Exo protein was concentrated to 20 mg/ml in crystallization buffer (20 mM Tris–HCl, pH 8.0). Crystallization conditions were screened with the Hampton Research Screens kits. Crystallization was performed by the sitting-drop vapor-diffusion method at 293 K, with 1 μl protein solution mixed with 1 μl reservoir solution and equilibrated with 100 μl reservoir solution containing 0.1 M Tris–HCl pH 8.4 and 1.0 M potassium sodium tartrate tetrahydrate. Good quality crystals were obtained within 2 days, and were flash cooled and stored in liquid nitrogen for future data collection. The X-ray diffraction data were collected to 1.9 Å resolution at 100 K at beamline BL5A of the Photon Factory (Japan) using an ADSC-Q315 detector. Reservoir solution containing 20% glycerol was used as cryoprotectant. The diffraction data were processed with the HKL2000 program suite (47). The LHK-Exo crystals obtained belong to space group \(P_6_3\) with unit cell parameters of: \(a = b = 108.9\ \text{Å}, c = 47.6\ \text{Å}, \alpha = \beta = 90^\circ\) and \(\gamma = 120^\circ\).

To obtain substrate-bound complex structures, native LHK-Exo crystals were soaked at 293 K for 24 h in a solution containing dAMP or a 5'-phosphorylated pentamer of oligothymidine [5'-PO\(_4\)-(dT)\(_5\)] at a final concentration of 1 mM in 0.1 M Tris–HCl pH 8.4 and 1.0 M potassium sodium tartrate tetrahydrate. Immediately prior to data collection, crystals were soaked in cryoprotectant solution consisting of 20% glycerol and the reservoir solution. X-ray diffraction data were collected in-house at 100 K on a Rigaku R-AXIS IV++ detector using Cu K\(_\alpha\) radiation (\(\lambda = 1.5418\ \text{Å}\)) from a Rigaku MicroMax-007 rotating anode X-ray generator operating at 40 kV and 20 mA. All diffraction data were integrated, scaled and merged using the HKL2000 suite (47). A complete summary of the data collection statistics is given in Table 1.

**Structure determination and refinement.** The structure of LHK-Exo was solved by molecular replacement (MR) with the monomeric \(\lambda\)-exonuclease (PDB code: 1AVQ) employed as a search model, using the Phaser program (48) in the CCP4 suite (49). Good quality electron density maps enabled initial model building by ARP/WARP (50). Further model building and manual adjustments were facilitated using the Coot program (51) and structure refinement was performed by Refmac5 (52). The stereochemical quality of the structure model was monitored with the PROCHECK program (53).

The substrate-bound complex structures were also determined by MR with the native enzyme employed as
Ramachandran plot
Structure refinement statistics

Data set LHK-Exo LHK-Exo:dAMP LHK-Exo:ssDNA
Data collection statistics
Wavelength (Å) 1.0000 1.5418 1.5418
Space group P63 P63 P63
Cell dimensions, a/b/c (Å) 108.9/108.9/47.6 108.3/108.3/47.8 108.7/108.7/48.2
Resolution (Å) 50–1.9 50–2.6 50–2.8
Average I/σ (I) 32.6 (2.0) 43.7 (4.3) 35.5 (3.4)
Completeness (%) 95.1 (70.6) 99.5 (100.0) 99.9 (100.0)
Generously allowed (%) 0 0 0.6
Allowed (%) 5.3 8.2 5.9
Most favored (%) 94.7 91.8 93.5

Structure refinement statistics
Resolution (Å) 50–1.9 50–2.6 50–2.8
Average R-factor (Å) 37.6 65.5 69.9
Rmerge (%) 20.4/24.1 20.8/26.2 19.3/24.8
RMSD bond lengths (Å) 0.019 0.016 0.014
RMSD bond angles (°) 1.629 1.608 1.550

Values in parentheses correspond to the highest resolution shell.

RESULTS AND DISCUSSION
LHK-Exo is an alkaline exonuclease that digests dsDNA with strict 5’- to 3’-polarity

The LHK_01497 gene from the HLHK9 type strain of L. hongkongensis was cloned into the pET28a vector as a C-terminal hexahistidine fusion, and subsequently expressed to high levels in E. coli. Three-step purification by immobilized Ni²⁺ ion affinity, anion exchange and gel-filtration chromatography afforded multi-milligram quantities of LHK-Exo protein of high purity suitable for biochemical and structural characterization. Analogous to the phylogenetically related λ-exonuclease and SPP1-Chu (G34.1P) proteins, recombinant LHK-Exo strictly required Mg²⁺ or Mn²⁺ ions for the hydrolysis of linear dsDNA molecules (Supplementary Figure S1) (20,21,31). Analysis of quenched aliquots from a time-course incubation of LHK-Exo with linear dsDNA by agarose gel electrophoresis, indicated that LHK-Exo was functioning as an exonuclease that did not release significant amounts of partially digested products (Figure 1A). This is consistent with the activities of λ-exonuclease (30,54), SPP1-Chu (20,21), RecE (55) and SXT-Exo (26), that all function as highly processive dsDNA exonucleases.

Modified dsDNA substrates (712 bp in length), containing three consecutive ‘nuclease resistant’ phosphorothioate linkages near the 5’-termini of both strands (PT modified) were synthesized using a PCR-based strategy, to probe the polarity of exonucleolytic digestion (Figure 1B). Analogous dsDNA substrates with no backbone modifications (unmodified) were similarly synthesized as controls. By substituting phosphodiesters linkages with phosphorothioate linkages near the 5’-termini, the activities of a 5’- to 3’-exonuclease should be severely inhibited, whereas the activities of a 3’- to 5’-exonuclease should be relatively unaffected. We have previously used this approach to probe the in vivo DNA hydrolytic activities of the SXT-Exo protein (26). λ-exonuclease was included as a control, as this enzyme has previously been shown to hydrolyze DNA in the 5’- to 3’-direction. LHK-Exo degraded negligible amounts of the (5’-phosphorylated) ‘PT-modified’ dsDNA substrate (Figure 1B, lane 5), but fully digested the analogous ‘unmodified’ dsDNA substrate under the conditions tested (lane 3). λ-exonuclease exhibited analogous activities (lanes 6–9). These assays clearly demonstrated that LHK-Exo specifically digests dsDNA from the 5’-termini; i.e. it hydrolyzes dsDNA with strict 5’- to 3’-polarity.

We probed the ssDNA hydrolysis activities of LHK-Exo using 5’-phosphorylated and non-phosphorylated forms of a 50-mer of oligothymidine [5’-PO₄(dT)₅₀ and 5’-OH-(dT)₅₀ respectively] as model substrates. LHK-Exo effectively hydrolyzed both 5’-phosphorylated and non-phosphorylated ssDNA substrates (Figure 1C and D); although the 5’-phosphorylated oligonucleotide was digested ~5- to 10-fold more rapidly. As was the case for dsDNA substrates, Mg²⁺ ions were essentially required for activity (Supplementary Figure S2). Notably, the hydrolysis of 5’-OH-(dT)₅₀ was strongly inhibited at high salt concentrations (1 M KCl), while the degradation of the
5'-phosphorylated form was less affected (Supplementary Figure S2). This is consistent with 5'-phosphorylated DNA molecules binding to the LHK-Exo protein more effectively than the corresponding non-phosphorylated ones, and consequently being preferred as substrates (see below), as is the case for the λ-exonuclease, SXT-Exo and SPP1-Chu proteins (20,21,26,36).

The dsDNA exonuclease activities of LHK-Exo were investigated in a quantitative manner using the sensitive PicoGreen fluorescent DNA stain, as has previously been used in analogous studies (21,36,45). As may be seen in Figure 2, LHK-Exo activities were optimal in the presence of Mg$^{2+}$ or Mn$^{2+}$ ions at a concentration of 1 mM (Figure 2B). This is roughly similar to concentrations of Mg$^{2+}$ ions utilized by related exonucleases of viral/phage origin: λ-exonuclease [1–2.5 mM (31)]; BGLF5 from the Epstein-Barr virus [EBV; 20 mM (56); 5 mM (57)]; baculovirus alkaline exonuclease [4–8 mM (58)]; UL12 from Herpes simplex virus 1 [2–5 mM (59)]; V-TREX from Choristoneura fumiferana nucleopolyhedrovirus [5 mM (60)]. Equivalent levels of dsDNA digestion were observed (within experimental error) when Mg$^{2+}$ or Mn$^{2+}$ ions were present at their optimal concentrations; suggesting that LHK-Exo can accommodate and utilize either ion equally well. This is notably different to the metal ion selectivities of related
exonucleases; where activities have been reported to be 4- to 10-fold higher in the presence of optimal concentrations of Mg$^{2+}$ ions, compared with optimal Mn$^{2+}$ ion concentrations (31,57–59). dsDNA degradation levels were highest at pH 8.2 (Figure 2C), and at a temperature of 41°C (Figure 2D). The optimal pH value for LHK-Exo is comparable to those determined for related exonucleases: λ-exonuclease [pH 9.2–9.5 (31)]; SPP1-Chu [pH 9.2–9.5 (21)]; SXT-Exo [pH 8.2 (26)]; BGLF5 [pH 8.0 (56); pH 8.5–9.5 (57)]; baculovirus alkaline exonuclease [pH >9–10 (58)]; UL12 [pH 9–10 (59)]; V-TREX [pH 9.5 (60)]. When the pH and Mg$^{2+}$ ion concentrations were optimal, LHK-Exo hydrolyzed 5'-phosphorylated linear dsDNA at a rate of 7.0 ± 1.8 nt/s (per LHK-Exo monomer, per DNA end; at 25°C; Table 2). This rate of hydrolysis is comparable to that previously determined for λ-exonuclease under a variety conditions (34,35,38,39).

LHK-Exo digests 5'-phosphorylated dsDNA in a highly processive manner

The digestion processivity of LHK-Exo was determined by a ‘heparin trap’ method (61), which has similarly been used to characterize the λ-exonuclease (36), SPP1-Chu (21) and SXT-Exo (26) alkaline exonucleases. In this approach, a large excess of the heteropolysaccharide ‘DNA-mimic’ heparin is added to assays containing exonuclease and dsDNA substrate shortly after their initiation. This sequesters all unbound enzyme, and also prevents any disassociated enzyme from re-binding to initiate additional digestion events. DNA levels are subsequently quantified using the fluorescent PicoGreen reagent, to determine the average number of nucleotides digested during a single DNA-binding event. Under

Table 2. Rate and processivity of double strand DNA digestion mediated by wild-type LHK-Exo and mutants

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<tr>
<th>Proteins</th>
<th>5'-PO$_4$ dsDNA</th>
<th>5'-OH dsDNA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$k_{cat}$ (nt/s)</td>
<td>processivity (nt)</td>
</tr>
<tr>
<td>WT</td>
<td>7.0 ± 1.8</td>
<td>2368 ± 72</td>
</tr>
<tr>
<td>R12A</td>
<td>1.0 ± 0.3</td>
<td>955 ± 38</td>
</tr>
<tr>
<td>E68A</td>
<td>2.1 ± 0.5</td>
<td>720 ± 32</td>
</tr>
<tr>
<td>K114A</td>
<td>1.0 ± 0.3</td>
<td>655 ± 45</td>
</tr>
<tr>
<td>Y134A</td>
<td>1.3 ± 0.4</td>
<td>812 ± 47</td>
</tr>
<tr>
<td>Y134F</td>
<td>4.0 ± 0.8</td>
<td>1140 ± 59</td>
</tr>
<tr>
<td>Q137A</td>
<td>1.8 ± 0.6</td>
<td>1122 ± 114</td>
</tr>
<tr>
<td>Q137E</td>
<td>0.3 ± 0.1</td>
<td>440 ± 52</td>
</tr>
</tbody>
</table>

Analogous sets of quenched PicoGreen fluorescence assays were performed to quantify the respective rates and processivities with which representative 5'-phosphorylated dsDNA (5'-PO$_4$-dsDNA) or 5'-dephosphorylated dsDNA (5'-OH-dsDNA) substrates were digested by wild-type LHK-Exo or the specified active-site point-mutant under standardized conditions. See ‘Materials and Methods’ section for details.
Figure 3. Processivity of double strand DNA digestion by wild-type LHK-Exo and Arg12Ala mutant. Time course analysis of the digestion of 5'-phosphorylated double strand DNA (5'-PO₄-dsDNA: EcoRV-linearized pMal-c2) and 5'-dephosphorylated double strand DNA (5'-OH-dsDNA: 5' dephosphorylated EcoRV-linearized pMal-c2) substrates by wild-type LHK-Exo and the Arg12Ala mutant form using a 'heparin trap' approach. A total of 6 μg (82 pmol of trimers) of LHK-Exo or Arg12Ala mutant protein was incubated at 25°C with 60 ng (0.015 pmol) of 5'-PO₄-dsDNA or 5'-OH-dsDNA in Tris–HCl (25 mM, pH 8.0), 1 mM DTT, 7.5 mM MgCl₂. After 15 s, excess heparin was added to sequester all unbound protein, and to prevent dissociated protein from re-binding. Aliquots were removed at various time points (0-20 min), and dsDNA levels were determined using fluorescent PicoGreen assays, to enable the extent of DNA digestion to be calculated. In one set of assays, heparin was added to the LHK-Exo prior to the addition of dsDNA substrate, to confirm the efficacy of the heparin trap method (filled black squares, green line). Graphs show the mean number of nucleotides digested from each terminus (±SD; y-axis) plotted against the time of analysis (in minutes; x-axis).

The purified recombinant LHK-Exo protein was crystallized using the sitting-drop vapor-diffusion method. Its crystal structure was solved to 1.9 Å resolution by molecular replacement using the structure of λ-exonuclease (PDB code 1AVQ) as a search model, with the crystallographic Rwork and Rfree converging to 20.4% and 24.1%, respectively (Table 1). The final structure includes one LHK-Exo monomer and 177 water molecules. LHK-Exo presents in the asymmetric unit as a monomer with residues 5–203 traced well in the model. Three expression vector-encoded residues located at the C-terminus of the protein were also traced in the structure. The four N-terminal residues, as well as residues 28–32 and 55–57 were unresolved, indicating that these regions may exist as flexible regions (discussed below).

The overall structure of LHK-Exo adopts an α/β 'pistol-like' fold including seven α-helices and four β-strands (Figure 4A). In addition to λ-exonuclease, a DALI search identified another four related structures, including: a phage-related exonuclease from Haemophilus somnus 129PT (HS_1420; PDB code 3K93; Z-score 17.6; r.m.s.d. 2.4 Å for 175 residues); the BGLF5 nuclease from Epstein-Barr virus (PDB code 2W4B; Z-score 11.4; r.m.s.d. 2.5 Å for 168 residues) (62); the Shut-off and Exonuclease (SOX) protein from Kaposi’s sarcoma-associated herpesvirus (PDB code 3FHD; Z-score 10.8; r.m.s.d. 2.4 Å for 160 residues) (63) and the E. coli RecE protein (PDB code 3H4R, Z-score 10.0, r.m.s.d. 3.4 Å for 152 residues) (38). The LHK-Exo, HS_1420, SOX and BGLF5 proteins belong to the λ-exonuclease family; while RecE shares more similarity with the RecB family (8). All possess DNAse and/or RNAse activities, with the exception of the HS_1420 protein, which has not yet been functionally characterized. This suggests that a conserved catalytic core domain exists in this family, despite members sharing fairly low levels of sequence identity and possessing quite different molecular weights. Superimposing the structures of these nucleases with that of LHK-Exo enabled the conserved nuclease 'core domain' to be clearly identified. This comprised a central arrangement of β-strands surrounded by α-helices (Supplementary Figure S3, panels A–F). Insertions within this central domain, as well as N-terminal and C-terminal extensions, exhibited significant diversity in shape and size.

LHK-Exo forms a toroidal trimer

The five related nuclease structures mentioned above have diverse oligomeric states: being monomeric (BGLF5 and SOX), trimeric (λ-exonuclease and HS_1420) and tetrameric (RecE). Consequently, analytical ultracentrifugation was performed to determine the oligomeric state of LHK-Exo. The sedimentation velocity showed a single...
4.4 S species corresponding to a mass of $79 \pm 4$ kDa (Supplementary Figure S4). The calculated mass for a trimer of LHK-Exo proteins is $\approx 74$ kDa, indicating that LHK-Exo forms a stable trimer in solution, analogous to $\lambda$-exonuclease. Although LHK-Exo presents in the asymmetric unit as a monomer, the trimeric conformation can be identified via a symmetric operation involving two neighboring molecules. Forming a ‘doughnut-like’ toroidal structure (Figure 4B), the constructed conformation of the LHK-Exo trimer is highly similar to that previously obtained for $\lambda$-exonuclease (41). A total solvent-accessible surface area of $992 \ \AA^2$ (9.1%) is buried at each monomer interface; formed mainly by $\alpha_3$, the $\alpha_3-\alpha_4$ loop and $\alpha_7$ in one molecule, and $\alpha_5$ and the $\beta_3-\beta_4$ loop in the neighboring molecule (Figure 4D).

Hydrophobic interactions between the neighboring subunits appear to be the main contributors towards the overall stabilization of the trimeric conformation. Two salt bridges; Glu53$_A$-Arg158$_B$ and Lys126$_B$-Glu194$_A$ also appear to play important roles (where the A and B subscripts refer to residues present on adjacent monomers). Buried water molecules located at the subunit interfaces may also play a role in trimer assembly, via the formation of hydrogen bonds. Wat237 links the O$_{\text{d1}}$ atom of Asp162$_B$ and the carbonyl oxygen atom of Lys199 A; Wat319 links the two carbonyl oxygen atoms of Asp162$_B$ and Ala202 A; and Glu53$_A$ is connected to Thr118$_B$ through Wat226 and Wat231. The hydrophobic interactions are mostly concentrated toward the centers of the interfaces; whilst the two salt bridges and the hydrogen bonds...
bond network mediated by the ordered water molecules are positioned toward the peripheries of the interfaces.

A tapered central channel enables the passage of a single strand of DNA

A tapered channel ~20 Å in length passes through the centre of the protein trimer (the ‘doughnut hole’), whose diameter is ~35 Å at the wide end and ~20 Å at the narrow end (Figure 4C). This central channel appears to be more ‘open’ (i.e. slightly wider) at both ends than the corresponding channel present in the λ-exonuclease trimer, due to the absence of the two (untraced) flexible loops. The RecE tetramer forms a similar central channel with a length of ~40 Å, which is tapered from ~30 Å at the wide end to ~10 Å at the narrow end (38).

The striking similarities in structural organization shared between the λ-exonuclease trimer, the RecE tetramer and the LHK-Exo trimer strongly suggests that they share a conserved mode of action. It has previously been proposed for λ-exonuclease and RecE that the dsDNA substrate enters through the wide end of the channel, and the non-digested 3'-ended ssDNA exits through the narrow end (38,41). For the LHK-Exo trimer, the positive electrostatic potential is predominantly concentrated at both ends of the central channel. However, the central channel does not have a strong overall positive electrostatic potential, and the active site possesses a strong negative charged surface (Figure 4C). Highly similar patterns of electrostatic potential distribution are also found within the central channels of the λ-exonuclease and RecE multimers. This suggests that for this group of exonucleases, the binding of the 5'-ended DNA strand (to be digested) may be primarily mediated by specific electrostatic and hydrogen bonding interactions. It also indicates that the ingress of the dsDNA substrate and the passage of the undigested strand out of the channel may be promoted primarily by electrostatic interactions.

In the LHK-Exo trimer structure, the two untraced flexible loop regions are positioned at both ends of the central channel. Residues 28–32 (TKSGY) that form the untraced loop between α2 and α3, extend out from the rim of the wide end (entrance) of the channel. Untraced residues 55–57 (RFS) lie between α3 and α4 on the opposite face of the toroid, lining the narrow channel ‘exit’. In the λ-exonuclease and RecE proteins, the two loops that correspond to these two stretches of untraced residues have been proposed to form important contacts with the incoming dsDNA substrate (residues 28–32), as well as the exiting strand of undigested DNA (residues 55–57) (38,41). From a functional perspective, it seems likely that these two loops are flexible in order to accommodate the translocation of the DNA molecule through the centre of the toroid, as the termini of the 5'-strand are processively hydrolyzed. However, sequence alignments reveal that both of these stretches of untraced residues show very low levels of homology with related exonucleases (Supplementary Figure S5); with the loop between α2 and α3 (residues 28–32) being absent in some homologs. The Arg55 residue, which is located in the small untraced loop proximal to the active site, may project toward the centre of the channel exit; analogous to Lys76 in λ-exonuclease and Arg858 in RecE (38,41). Correspondingly, this residue may form important binding contacts with the undigested 3'-strand as it translocates through the central channel of the enzyme.

Coordination of active-site metal ions

Active-site metal ions were not detected in the native crystal structures of either the λ-exonuclease or RecE proteins, and their positions were only identified after Mn<sup>2+</sup> ions were soaked into the crystals (38,41). Notably, in the native structure of LHK-Exo, two bound magnesium ions were identified in a negatively charged pocket within the active site (Figure 5A). One magnesium (II) ion (Mg<sup>2+</sup>) is coordinated to the Oe2 atom of Glu112 (2.3 Å); the Oe2 atom of Asp102
(2.4 Å); the carbonyl oxygen atom of Ile113 (2.5 Å); and to two water molecules, Wat383 (2.4 Å) and Wat384 (2.3 Å). The Mg\textsuperscript{2+}-binding site is essentially identical to the corresponding metal ion-binding site in λ-exonuclease, with the exception that two water molecules also contribute to the coordination geometry in our structure. It may also be noted that Mn\textsuperscript{2+} ions are chelated by histidine residues located in analogous positions in the active sites of both the RecE (His652) (38) and RecB (His956) exonucleases (64). The second, putatively more weakly bound magnesium ion (Mg\textsubscript{2n}), is located 3.7 Å away from Mg\textsubscript{2m}. A corresponding metal ion is absent in the λ-exonuclease structure. Mg\textsubscript{2n} is coordinated by the Oe1 atom of Glu112 (2.7 Å); the carbonyl oxygen atom of Ser100 (2.7 Å); the Oδ1 atom of Asp102 (3.3 Å); and two water molecules, Wat387 (2.7 Å) and Wat388 (3.2 Å). In contrast to the octahedral coordination geometry typically observed in Mg\textsuperscript{2+}-bound enzymes, both Mg\textsuperscript{2+} ions in the LHK-Exo structure have a partial octahedral coordination, leaving positions available for potential interactions with DNA substrates or water molecules during the catalytic process (see below).

Based on the sequence analysis of exonuclease homologs, it has previously been proposed that the highly conserved Glu68 residue (Glu85 in λ-exonuclease) is a potential ligand for Mg\textsuperscript{2+} ions (8). In our structure, Glu68 seems slightly too far from the Mg\textsuperscript{2m} ion (4.1 Å) to act as a direct ligand; however, a water molecule (Wat384) forms a bridge between Glu68 and the Mg\textsuperscript{2m} ion. To probe the possible involvement of this residue, the Glu68Ala mutant was constructed, expressed and biochemically characterized. This mutant protein hydrolyzed linear 5’-phosphorylated dsDNA at a rate of 2.1 ± 0.5 nt/s (pH 8.0, 7.5 mM MgCl\textsubscript{2}, 25°C), which was 70% lower than that of the wild-type LHK-Exo enzyme (7.0 ± 1.8 nt/s). However, the hydrolytic activities of the Glu68Ala mutant increased 1.6-fold (3.4 nt/s) when the MgCl\textsubscript{2} concentrations were increased to 20 mM, while those of the wild-type enzyme decreased ~35% to 4.7 nt/s (data not shown). This indicated that the Glu68Ala mutant had a reduced affinity for Mg\textsuperscript{2+} ions, which could be partially compensated for by increasing their concentration in solution. The coordination and functional roles of these two Mg\textsuperscript{2+} ions are discussed further below.

### Recognition of 5’-phosphorylated DNA termini

It has previously been shown that λ-exonuclease digests dsDNA substrates that have a phosphate group at their 5’-termini (5’-PO\textsubscript{4}) significantly more efficiently than analogous (dephosphorylated) substrates lacking this group (5’-OH) (30,35). Subramanian et al. (36) subsequently proposed that a phosphate ion observed in a pocket located 7 Å away from the Mg\textsuperscript{2+} ion in the λ-exonuclease crystal structure most likely occupied the position of the 5’-phosphorylated terminus of the DNA strand to be digested (41). They identified the Arg28, Ser35, Ser117 and Gln137 residues as forming key electrostatic interactions with this phosphate ion; which are very highly conserved amongst λ-exonuclease family members (Supplementary Figure S5). Mutation of the Arg28 residue of λ-exonuclease significantly reduced its digestion rate and processivity, and essentially abolished its ability to discriminate between 5’-phosphorylated and non-phosphorylated dsDNA substrates; demonstrating that it played a key role in binding the 5’-phosphate group. Contrastingly, an analogous residue is absent in the RecE structure (38), and correspondingly the phosphorylation status of the 5’-strand does not appear to significantly affect the rate of dsDNA hydrolysis (55).

In the LHK-Exo structure, the putative phosphate-binding pocket incorporates the Arg12, Thr17, Ser19, Ser100 and Gln137 residues. To confirm the proposed involvement of these residues in binding the 5’-phosphorylated DNA terminus, we determined the structure of LHK-Exo in complex with dAMP to a resolution of 2.6 Å. The 5’-phosphate group of this deoxy-mononucleotide fits comfortably into the binding pocket, coordinated tightly by Arg12, Thr17, Ser19 and Ser100 (Figure 5B and C; Supplementary Figure S6). Compared with λ-exonuclease, the phosphate-binding site in LHK-Exo is located 2 Å further away from the active site. Consequently, Thr17 rather than Gln137 appears to be involved in stabilizing the phosphate group. An equivalent residue is present in λ-exonuclease (Thr33), SXT-Exo (Thr22) and PLU2936 (Thr34), but is noticeably absent in other homologous proteins. Furthermore, the adenine base moiety of the nucleotide does not appear to form any specific interactions with LHK-Exo. It interacts loosely with Trp8 and Phe9, which may supply a flexible binding environment capable of accommodating the four different nucleobase moieties present in DNA substrates.

Notably, Arg12 undergoes a significant conformational change upon dAMP binding (Figure 5B); adopting an orientation consistent with that of Arg28, which binds the phosphate ion in the λ-exonuclease structure. To determine its putative involvement in binding the terminal nucleotide residue of the DNA chain to be hydrolyzed, the exonuclease activities of the Arg12Ala mutant were investigated. The mutant protein digested 5’-phosphorylated linear (5’-PO\textsubscript{4}) dsDNA substrates at a rate of 1.0 ± 0.3 nt/s, which is 7-fold lower than that of wild-type LHK-Exo. However, it digested the analogous 5’-dephosphorylated (5’-OH) dsDNA substrates at an essentially identical rate (1.1 ± 0.3 nt/s) (Table 2). Consistent with the proposed role of Arg28 in λ-exonuclease (36), these results indicate that the corresponding arginine residue in LHK-Exo plays a pivotal role in discriminating between 5’-phosphorylated and non-phosphorylated forms of DNA.

### Structure of the LHK-Exo protein complexed with a ssDNA oligomer reveals a binding channel for the hydrolyzed DNA strand

The λ-exonuclease and RecE exonuclease proteins have the ability to distinguish and separate the two strands of the DNA duplex, and to specifically digest the 5’-ended strand in a highly processive manner (30,36,54,55). In order to better understand the mechanistic basis for 5’-stand recognition and binding, we soaked crystals
of LHK-Exo with buffer containing a synthetic 5-mer of oligothymidine that had been enzymatically phosphorylated at the 5'-termini [5'-PO_4-(dT)_5]. We were subsequently able to solve the structure of LHK-Exo-oligonucleotide complex to 2.8 Å. We could clearly identify four complete nucleotide residues (T1–T4) in each monomer (Figure 6A and Supplementary Figure S6). The ssDNA chain fits comfortably into a channel formed by α4, α6 and the α5–β2 loop (Figure 6B). The T1 residue was located in the same site as the dAMP 5'-mononucleotide was (see above); coordinated tightly by Arg12, Thr17, Ser19 and Ser100. The 5'-phosphate group of dAMP (Figure 6A). However, the 3'-OH of the T1 deoxyribose ring was located 1.5 Å closer to the active site, and was hydrogen bonded to Mg_2+ (3.0 Å). The fact that the dAMP and T1 oligonucleotide residues occupied the same binding site in the LHK-Exo crystal structure, suggests that this is the most stable nucleotide binding
The scissile phosphate between the T1 and T2 nucleotides was stabilized by the two Mg
ions via the O1 atom. In our experiments, the O1 atom of the phosphate moiety between T2 and T3 formed a 2.9 Å hydrogen bond with the main chain amide nitrogen of Cys115. The O1 atom of the phosphate moiety between T3 and T4 was stabilized by hydrogen bonds to the hydroxyl group of Thr120 (2.6 Å) and the main chain amide nitrogen of Asn117 (3.1 Å). Notably, no specific interactions appeared to be formed between the base moieties of the oligonucleotide and the LHK-Exo protein. This would be consistent with the exonuclease protein exhibiting little or no preference for dsDNA substrates with differing nucleotide compositions. We could not identify the T5 residue at the 3′-termini of the oligonucleotide in the complex structure, suggesting that this nucleotide does not form stable binding interactions with the enzyme.

Our findings indicate that LHK-Exo recognizes and tightly binds at least four consecutive nucleotides at the 5′-termini of the DNA strand to be digested (T1–T4), via the sugar–phosphate backbone. This stretch of residues at the 5′-termini is most probably separated from the complementary residues of the (undigested) DNA strand. However, it is difficult to extrapolate our results to estimate the degree to which the B-form DNA duplex is unwound by enzyme, or precisely how many base pairs are separated. Notably, our X-ray data is consistent with the findings of Dahlroth et al. (63), who modeled an oligonucleotide into the active site of the SOX exonuclease crystal structure. They predicted that SOX was also likely to accommodate 4 nt.

It should be noted that the soaking experiments did not lead to ssDNA molecule cleavage in our complex structure, which was similarly observed for the EcoRV endonuclease (65). While LHK-Exo possessed notable ssDNA exonuclease activities in the presence of 7.5 mM Mg
ions (in solution phase at pH 8.0, 37°C), there was negligible activity when Mg
ions were omitted (Supplementary Figure S2). The crystal soaking experiments were conducted at high salt concentrations, at 20°C, in the absence of free Mg
ions, which may have facilitated the retention of an intact tetranucleotide molecule within the active site. In addition, some quaternary structure changes may occur during the catalysis process that are in conflict with the crystal packing forces, which may be responsible for the prevention of ssDNA cleavage. It may also be the case that the position of the tetranucleotide molecule within the active site corresponds to a protected region of the substrate DNA as it is processed by LHK-Exo, analogous to a proposed ~11-bp protected region of DNA in λ-exonuclease (41), although further experiments are needed to confirm this.

A similar ssDNA-binding channel exists in λ-exonuclease and the HS_1420 phage-related exonuclease from Haemophilus somnus 129PT, indicating that they may share a common binding mechanism for the 5′-DNA strand (Figure 6C and D). However, the corresponding ssDNA-binding channel of the λ-exonuclease trimer appears to be slightly more ‘closed’ at the 5′-phosphate recognition site, due to the presence of an extra α-helix and 310-helix located at its N-terminus (Figure 6B–D).

We further noted an additional binding pattern in the structure of the oligonucleotide–protein complex. The three central phosphodiester groups of the DNA chain (T1–T2, T2–T3 and T3–T4) are only coordinated to the protein via their O1 atoms, leaving their O2 atoms oriented toward the same inner-surface of the channel (Supplementary Figure S7). This surface is composed of Lys114, Tyr134 and Gln137; three residues that are highly conserved among λ-exonuclease family members (Supplementary Figure S5). The distances between the three O2 atoms and the respective polar side-chain groups of these residues are maintained within 4.5 Å. We speculated these three residues may play important roles in binding and orienting the 5′-DNA strand during its step-wise hydrolysis. To probe the role of these universally conserved residues, we prepared the Lys114Ala, Tyr134Ala, Tyr134Phe, Gln137Ala and Gln137Glu mutants and characterized their activities. The Lys114Ala and Tyr134Ala mutants digested 5′-phosphorylated dsDNA with rates of 1.0 and 1.7 nt/s, respectively; which was roughly 15% and 25% that of the wild-type enzyme (Table 2). However, the digestion rate of the Tyr134Phe mutant was 4.0 nt/s, which was ~60% that of wild-type LHK-Exo protein. The processivity of digestion mediated by these three mutants were all roughly similar (~1100 nt hydrolyzed per binding event), which is ~50% that of the wild-type enzyme. This data indicate that neither of these individual residues plays vital roles in DNA hydrolysis or translocation. However, the fact that the Tyr134Phe mutant was significantly more active than the Tyr134Ala mutant suggests that the presence of the benzyl side-chain moiety (compared to a methyl group) is beneficial for the overall protein architecture, or at least plays an ancillary role in substrate binding or catalysis. The dsDNA digestion rate of the Gln137Ala mutant was 4-fold lower than the wild-type (1.8 ± 0.6 nt/s), while the Gln137Glu mutant was 23-fold less active. This ~6-fold difference in catalytic activities between these two mutant forms most likely results from considerable electrostatic repulsion between the negatively charged pendant carboxylate group and the phosphate of the DNA substrate. Notably, it has been proposed that the corresponding glutamine residue in RecE (Gln781), is hydrogen bonded to Lys761; thereby modulating the orientation or catalytic activity of this highly conserved lysine residue (see below) (38).

Proposed two metal ion catalytic mechanism for LHK-Exo

Although the crystal structure of λ-exonuclease was solved more than a decade ago, its precise catalytic mechanism remains to be established. Our crystal structure of the LHK-Exo protein in complex with a ssDNA molecule and Mg
ions gives a highly informative snapshot of the active site of this exonuclease enzyme. In particular, the relative positioning of the two Mg
ions suggests that
this protein may utilize a two-metal ion catalytic mechanism for DNA hydrolysis (Figure 7A). This has previously been proposed to occur in a variety of polymerases (66, 67) and nucleases (68–72), but within diverse catalytic scenarios. The two bound Mg2+ ions are located approximately 4 Å away from each other, parallel to the scissile bond. Mg2+ A is tightly coordinated by Glu112, Asp102, Ile113 and the scissile phosphate; but Mg2+ B is more weakly bound by Glu112, Ser100, Asp102 and the scissile P-O. In the presence of DNA, the two Mg2+ ions are jointly coordinated by the side-chain carboxylate groups of two residues that are highly conserved within this protein family: Asp102 and Glu112 (Supplementary Figure S5), as well as the O1 atom of the scissile phosphate. It must be noted however, that neither of these magnesium ions have a fully occupied octahedral coordination in our structure. It is possible that water molecules crucially involved in the catalytic process may be missing due to the modest resolution of our data set (discussed below). As described previously, Mg2+ A may contribute to the deprotonation and subsequent stabilization of the nucleophilic water molecule. Mg2+ B could stabilize the scissile P-O bond and facilitate the departure of the ribose 3'-hydroxyl moiety. Lys114 is postulated to act as a conserved general base catalyst. (B) Architecture of the active site of the BglII restriction endonuclease with two complexed Ca2+ ions (PDB code: 1DMU). Adenosine and thymidine nucleotides of the bound DNA molecule are shown in pale green-colored stick form. Active site residues are shown in gray-colored stick form and the bound Ca2+ ions are shown as gray spheres. Lys144 has been proposed to act as a conserved general base catalyst, in a 'two-metal' catalytic mechanism. The two water molecules that putatively act as the nucleophile and general acid are colored red. The blue arrow indicates the scissile P-O bond.

In the proposed two metal ion catalytic mechanism, the highly conserved Lys114 (equivalent to Lys131 in λ-exonuclease, Lys761 in RecE, Lys144 in BglII) may function as a conserved general base for the activation of the nucleophilic water molecule, which has previously been suggested (38, 41). Mg2+ A and the (protonated) ε-amino group of Lys114 may stabilize the negatively charged pentacoordinate phosphate intermediate. Substituting Lys114 for an alanine resulted in a ~85% loss in activity compared with wild-type LHK-Exo (1.0 ± 0.3 nt/s compared to 7.0 ± 1.8 nt/s; Table 2). Its digestion processivity was similarly reduced by ~75% (655 ± 45 nt compared to 2368 ± 72 nt digested per binding event). This indicates that while this highly conserved lysine residue may play an important role in facilitating phosphodiester bond hydrolysis, its individual functions are not essential.
bound structure of LHK-Exo due to the modest resolution and slightly smeared electron density. Furthermore, we are not able to identify the precise mechanism by which LHK-Exo promotes the formation and stabilization of the pentacoordinate phosphate intermediate. However, the analogous composition and orientation of the active site residues and metal ions within the BglI and LHK-Exo crystal structures supports the operation of an equivalent two-metal ion catalytic mechanism. A more precise identification and positioning of the crucial residues and water molecules within the LHK-Exo active site will require higher resolution structures in complex with ssDNA or even (partially) dsDNA model substrates.

The biophysical, biochemical and structural data presented here suggest that a considerable number of residues within the central channel of the LHK-Exo trimer are actively involved in binding, orienting and translocating the DNA substrate; as well as playing a direct catalytic role in the iterative hydrolysis of nucleotides from the terminus of the 5'-strand. Our results are consistent with the mutational analysis of the RecE exonuclease, where the precise roles of the individual active site residues proved similarly difficult to unambiguously establish, and there appears to be a significant amount of catalytic redundancy (38). While many key mechanistic points remain to be established, the structures of the mononucleotide and oligonucleotide-bound complexes of LHK-Exo described here are the first such structures reported for a member of this notable class of exonucleases. They provide structural evidence consistent with the operation of a two metal ion catalytic mechanism for this protein, and offer significant additional insight into the catalytic mechanisms of related nucleases of phage/viral origin.

**ADDENDUM**

While revising our manuscript, we became aware of an ‘early edition’ article reporting two DNA-bound complexes of λ-exonuclease (75). The authors used 12-mer dsDNA (Form 1) or a 14-mer:12-mer DNA duplex (containing a 2nt 5'-overhang; Form 2) for co-crystallization with the wild-type and catalytically inactive K131A mutant forms of λ-exonuclease, respectively. Interestingly, the position of the ssDNA in our LHK-Exo complex structure overlaps precisely with the 5'-end of the 14-mer DNA strand in the Form 2 structure, including the 2-nt overhang. The λ-exonuclease:dsDNA structures reported by Zhang and colleagues are highly consistent with our LHK-Exo:ssDNA structure reported here. Taken together, our studies reveal that these two exonuclease proteins share an equivalent arrangement of metal cofactor and substrate bonding interactions, and employ a functionally identical two metal ion catalytic mechanism.

**ACCESSION NUMBERS**

The coordinates of the crystal structures of LHK-Exo, and the LHK-Exo:dAMP and LHK-Exo:ssDNA complexes have been deposited in the RCSB Protein Data Bank with the accession codes 3SYY, 3SZ4 and 3SZ5, respectively.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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