Complementation of One RecA Protein Point Mutation by Another

EVIDENCE FOR TRANS CATALYSIS OF ATP HYDROLYSIS*

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The RecA residues Lys248 and Glu96 are closely opposed across the RecA subunit-subunit interface in some recent models of the RecA nucleoprotein filament. The K248R and E96D single mutant proteins of the Escherichia coli RecA protein each bind to DNA and form nucleoprotein filaments but do not hydrolyze ATP or dATP. A mixture of K248R and E96D single mutant proteins restores dATP hydrolysis to 25% of the wild type rate, with maximum restoration seen when the proteins are present in a 1:1 ratio. The K248R/E96D double mutant RecA protein also hydrolyzes ATP and dATP at rates up to 10-fold higher than either single mutant, although at a reduced rate compared with the wild type protein. Thus, the K248R mutation partially complements the inactive E96D mutation and vice versa. The complementation is not sufficient to allow DNA strand exchange. The K248R and E96D mutations originate from opposite sides of the subunit-subunit interface. The functional complementation suggests that Lys248 plays a significant role in ATP hydrolysis in trans across the subunit-subunit interface in the RecA nucleoprotein filament. This could be part of a mechanism for the long range coordination of hydrolytic cycles between subunits within the RecA filament.

Homologous DNA recombination is a vital component of DNA metabolism, central to processes such as recombinational DNA repair of stalled replication forks and the exchange of genetic material during meiosis in eukaryotes and conjugation in prokaryotes. RecA protein is the central recombinase in Escherichia coli, and RecA homologues are present in nearly every organism. In E. coli, RecA participates not only in the restart of stalled replication forks but also the induction of the SOS response upon cellular DNA damage distress and translesion DNA synthesis via the error-prone DNA polymerase V (1–3).

RecA protein functions as a nucleoprotein filament. When bound to DNA, RecA promotes the hydrolysis of ATP or dATP. Hydrolysis occurs uniformly throughout the filament. ATP hydrolysis is important for some RecA processes including net disassembly of the nucleoprotein filament. The K248R and E96D single mutant proteins restore dATP hydrolysis to 25% of the wild type rate, with maximum restoration seen when the proteins are present in a 1:1 ratio. The K248R/E96D double mutant RecA protein also hydrolyzes ATP and dATP at rates up to 10-fold higher than either single mutant, although at a reduced rate compared with the wild type protein. Thus, the K248R mutation partially complements the inactive E96D mutation and vice versa. The complementation is not sufficient to allow DNA strand exchange. The K248R and E96D mutations originate from opposite sides of the subunit-subunit interface. The functional complementation suggests that Lys248 plays a significant role in ATP hydrolysis in trans across the subunit-subunit interface in the RecA nucleoprotein filament. This could be part of a mechanism for the long range coordination of hydrolytic cycles between subunits within the RecA filament.

waves of hydrolysis move sequentially through the filament with a separation of six subunits (9). RecA could potentially use the organized waves of ATP hydrolysis to act as a motor, driving completion of strand exchange beyond the barriers mentioned above, or to promote replication fork regression (10, 11). The detailed mechanism by which the ATP hydrolysis is coordinated in the RecA-dsDNA nucleoprotein filament is currently unknown.

RecA is a member of an ATPase family with characteristic single or multiple RecA-like folds, composed of a central β-sheet flanked by α-helices. The ATPase family includes helicases like PcrA, ABC transporters, and with relatively less structural similarity, the AAA+ family of ATPases including ClpA (12). These proteins all use the energy from ATP hydrolysis to move macromolecules or move along macromolecules, and hence, may be considered molecular motors. A sequence alignment of RecA and helicases including DnaB (13) highlights the similarities. The authors call attention to a new motif, the [KR]×[KR] motif. This motif is conserved among the DnaB, RecA, Sm, and KaiC families, although absent from the archaean and eukaryotic homologues of RecA.

The [KR]×[KR] motif is involved in the in trans catalysis of ATP hydrolysis for some RecA homologues including gp4 of bacteriophage T7 (13–16). The helicase domain of T7 gp4 constitutes a 5’ to 3’ hexamer ring helicase. The crystal structure of a gp4 fragment that retains hexamer formation and helicase activities (residues 241–566) (15) strays from 6-fold rotational symmetry. Indeed, only four of six subunits have bound nucleotide (ADPNP). Arg522 is the third residue in the [KR]×[KR] motif, and in the four subunits that bind ADPNP, Arg522 is close (3.5 Å) to the γ-phosphate of the bound nucleotide of the neighboring subunit. However, Arg522 is displaced in the two nucleotide-free subunits to more than 10 Å away from the equivalent position where the nucleotide would be bound. A conformational change such as this is suggested to couple hydrolysis and helicase activity. It is proposed that Arg522 of T7 gp4 senses hydrolysis of ATP, communicates the hydrolysis between subunits, and promotes conformational changes associated with ATP hydrolysis (14–16). In this manner, Arg522 would mimic an arginine finger, as is found in Ras and its GTPase-activating protein (GAP). In the Ras-GAP system, it is postulated that Arg789 of GAP similarly reaches across the GAP-Ras interface to stabilize the γ-phosphate of the Ras-bound ATP (17).

Another example of in trans catalysis is the RuvB protein (18–20). The RuvB protein is an ATPase that forms a hexameric ring and complexes with RuvA to promote branch migration that is dependent on ATP hydrolysis. Although this protein exhibits some structural similarity to RecA, it appears to lack the [KR]×[KR] motif. The crystal structure features a dimer of Thermus thermophilus RuvB, with each subunit having three domains (labeled N, M, and C). The nucleotide is bound between the N and M domains (19). The structure of each subunit is very similar, except for the area immediately around the bound nucle-
otide. One subunit appears to have a triphosphate bound, while the other appears to lack a γ-phosphate; the authors postulated that the structural differences between the subunits could reflect conformational changes dependent on ATP hydrolysis. The crystal structures of wild type and several point mutants of Thermatoga maritima RuvB also indicate ADP is bound between two domains of the RuvB subunit (18), suggesting to the authors that the state of the bound nucleotide guides conformational changes between RuvB subunits. More specifically, sensor 1, Walker A, and Walker B were proposed to sense the γ-phosphate and divalent cation, while sensor 2 would accommodate the sugar ring and diphosphate. A study of E. coli RuvB point mutants further extends the proposition that ATP hydrolysis causes conformational changes between subunits by indicating that Arg174 (corresponding to Arg170 of T. maritima RuvB) acts in trans across the subunit interface in the RuvB hexamer (20). The Arg174 residue of RuvB does not comprise the active site for ATP hydrolysis in one subunit, but Arg174 is near the catalytic site of the adjacent subunit in the hexamer. The Walker A motif mutant K68A and R174A mutant are both deficient in ATP hydrolysis, but they were found to complement each other when mixed together (20). Hence, Arg174 of RuvB appears to participate in trans in ATP hydrolysis.

In E. coli RecA protein, the residues in the [KR]×[KR] motif are Lys248 and Lys250 (13). These two residues are hypothesized to be important for ATP hydrolysis and potentially communicate ATP hydrolysis between adjacent subunits in the RecA nucleoprotein filament according to the similarities to homologous proteins described above. Furthermore, the promising importance of one or both of the Lys248 and Lys250 residues was reinforced upon examination of the structure of the RecA-dsDNA nucleoprotein filament (21).

An impediment to the study of the RecA protein has been the lack of a crystal structure of the RecA filament bound to DNA. The crystal structure of the RecA filament with ADP bound (22) and the crystal structure of the RecA filament (23) provided much insight. However, RecA is a DNA-dependent ATPase, and the apo-cystal represents the inactive form of the RecA filament (21). Recently a reconstruction of the RecA filament with dsDNA, based on complexes viewed by electron microscopy, was resolved (21). The crystal structure of the RecA filament was fitted within electron microscopic images of the RecA-dsDNA filament. To fit the crystal structure electron density into the electron microscopy images of the nucleoprotein filament, the RecA core was rotated and resulted in several significant structural changes. One difference between the apo-crystal structure and the electron microscopic reconstruction of the RecA-dsDNA nucleoprotein filament is location of the ATP binding site (Fig. 1). The nucleotide is bound away from the subunit-subunit interface within the filament in the apo-crystal structure. In contrast, the nucleotide is bound in between adjacent subunits in the RecA-dsDNA nucleoprotein filament reconstruction. The location of the nucleotide between two RecA subunits is supported by recent crystal structures of the archaeal RadA filament (24) and the eukaryotic Rad51 filament (25). VanLoo et al. (21) states there are several residues that do not contact the nucleotide in the apo-crystal structure but are near the nucleotide of the adjacent subunit in the electron microscopic reconstruction. The shifted residues include Lys248, Phe217, Arg222, Lys248, Lys250, and Pro254. The two residues Lys248 and Lys250 are of particular interest because, as noted above, they are the two residues that comprise the [KR]×[KR] motif in E. coli RecA protein.

In this study, we sought to examine the K248 residue as an introduction to studying potential roles for the [KR]×[KR] motif in E. coli RecA. The K248A mutation (26) has been previously examined, and although the K248A mutant protein appears to fold normally, the K248A mutant does not form a filament. As such, we chose to introduce a more subtle mutation: K248R. We find that the K248R mutant does bind DNA and forms a nucleoprotein filament, although it hydrolyzes neither ATP nor dATP at a readily measured rate. However, we discovered that a combination of the K248R mutant and another ATP hydrolysis-deficient mutant (27, 28), E96D, partially restores dATP hydrolysis. In the nucleoprotein filament, the Lys248 and Glu96 residues are located at the subunit-subunit interface near the bound nucleotide but are opposed across the interface. Direct or indirect coordination between these two residues could be a means of communicating ATP hydrolysis-mediated conformational changes between adjacent subunits in the RecA nucleoprotein filament.

**EXPERIMENTAL PROCEDURES**

*Enzymes and Biochemicals—*The E. coli SSB protein was purified as described previously (7). The E. coli wild type RecA protein was purified as described previously (29). The RecA K248R mutant protein was purified in the same manner as wild type. The RecA K248R/E96D double mutant protein was purified in the same manner as wild type with the following changes: the protein was eluted twice from the ammonium sulfate pellet in a solution of R Buffer (20 mM Tris·Cl buffer (10 mM KCl to R Buffer plus 100 mM KCl to R Buffer plus 500 mM KCl). The K248R/E96D mutant protein was eluted from the DEAE-Sepharose column with a linear gradient from R Buffer plus 100 mM KCl to R Buffer plus 500 mM KCl. The K248R/E96D mutant protein was eluted from the hydroxyapatite column with a linear gradient from 20 mM phosphate buffer (10 mM KH2PO4, 10 mM K2HPO4, 10% glycerol (v/v), 1 mM dithiothreitol) plus 150 mM ammonium sulfate. The K248R/E96D mutant protein was eluted from the diethylaminoethyl (DEAE)-Sepharose column with a linear gradient from R Buffer plus 100 mM KCl to R Buffer plus 500 mM KCl. The K248R/E96D mutant protein was eluted from the hydroxyapatite column with a linear gradient from 20 mM phosphate buffer (10 mM KH2PO4, 10 mM K2HPO4, 10% glycerol (v/v), 1 mM dithiothreitol) to 500 mM phosphate buffer (250 mM KH2PO4, 250 mM K2HPO4, 10% glycerol (v/v), 1 mM dithiothreitol). Peak fractions were identified by SDS-PAGE, pooled and dialyzed against R Buffer, and then loaded onto a PBE-94 column (Amersham Biosciences). The K248R/E96D mutant protein was eluted with a linear
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gradient from R Buffer to R Buffer plus 1 M KCl. The peak and nuclease-free fractions were pooled and concentrated with an Amicon Centrifoc concentrator (Millipore, Billerica, MA). The K248R/E96D mutant protein was dialyzed against R Buffer and flash-frozen in liquid nitrogen and finally stored at −80 °C. The E96D mutant RecA protein was purified similarly to the K248R/E96D mutant, excluding the PBE-94 column.

The purified RecA protein (both mutants and wild type) and SSB protein concentrations were determined by absorbance at 280 nm, using extinction coefficients of ε260 = 2.23 × 10^4 M⁻¹ cm⁻¹ (30) and ε280 = 2.38 × 10^4 M⁻¹ cm⁻¹ (31), respectively. RecA protein and SSB preparations were free of detectable endo- and exonuclease activities on dsDNA and ssDNA. Unless otherwise noted, all reagents were purchased from Fisher. Phosphoendoproteinase (PEP), pyruvate kinase, lactate dehydrogenase, phosphocreatine, ATP, dATP, and NADH were purchased from Sigma. Creatine kinase and ATP·γS were purchased from Roche Applied Science. Restriction enzymes were purchased from New England Biolabs (Beverly, MA). Dithiothreitol was purchased from Research Organics (Cleveland, OH).

**DNA Substrates**—Poly(dT) was purchased from Amersham Biosciences, and the approximate average length is 229 nucleotides. Bacteriophage ΦX174 circular ssDNA and replicative form I supercoiled circular duplex DNA were purchased from New England Biolabs and Invitrogen, respectively. Full-length linear duplex DNA (dsDNA) was generated by digesting ΦX174 replicative form I DNA (5386 bp) with the XhoI endonuclease. Circular single-stranded DNA from bacteriophage M13mp8 (7229 nucleotides) was prepared as described (32). Concentrations of ssDNA and dsDNA were determined by absorbance using 108 and 151 μM A260⁻¹, respectively, as conversion factors. All DNA concentrations are given in micromolar nucleotide concentrations.

**Circular Dichroism**—Wild type RecA protein and two separate preparations of the RecA K248R/E96D double mutant protein were analyzed on a model Aviv 62A DS circular dichroism Spectrometer (Aviv, Lakewood, NJ) equipped with a temperature controller. For each protein, the solution included 0.25 mg/ml protein, 10 mM KH₂PO₄, and 10 mM K₂HPO₄. The solutions were measured at 4 °C and 37 °C. Graphs of wavelength (nm) versus molar ellipticity (degree cm² dmol⁻¹) were normalized with respect to the absolute minimum at 208 nm. The RecA K248R/E96D protein CD spectrum and wild type RecA protein CD spectrum were superimposed to identify any differing points on the graphs. Two separate preparations of the RecA K248R/E96D mutant protein were examined.

**Reaction Conditions**—All reactions were carried out at 37 °C in 25 mM Tris acetate buffer (80% cation, pH 7.5), 10 mM magnesium acetate, 5% (v/v) glycerol, 1 mM dithiothreitol, 3 mM potassium glutamate, 3 mM ATP or dATP, an ATP regenerating system (12 mM phosphocreatine and 10 units/ml creatine kinase), 6 mM poly(dT) linear ssDNA, and 8 μM total RecA. Reactions were incubated at 37 °C for 10 min before ATP (to 3 mM) was added to start the filament formation. The reaction was again incubated at 37 °C for 10 min to allow RecA to form a filament on the DNA. ATP·γS was then added to 3 mM, and the reaction was incubated at 37 °C for 3 min to stabilize the RecA·DNA filament. The reaction solution was then diluted 5- or 10-fold in buffer (200 mM ammonium acetate, 10 mM HEPS, pH 7.5, and 10% glycerol) and prepared for spreading onto grids. Digital images of the nucleoprotein filaments were taken at ×15,000 magnification. Filament lengths were measured using MetaMorphp analysis software (Molecular Devices). At least 50 filaments were measured for each RecA protein. Each filament was measured three times, and the average length was calculated. Filaments less than 30 nm long were difficult to distinguish from the background and were not included in the analysis.

A cytochrome c method was used to visualize poly(dT) linear ssDNA as a control. Samples were prepared as described previously (36), except the spreading solution was assembled by combining 32 μl of 1 M Na₂CO₃, 40 μl of 0.126 M Na₂EDTA, 400 μl of 37% HCHO, 292 μl of 1 M formamide, and 10 μl of 5 M KOH. The final DNA concentration was 0.004 μg/μl.

**RESULTS**

**Experimental Design**—We focused on the contribution of two residues at the subunit-subunit interface of the nucleoprotein filament to catalysis of ATP hydrolysis by the RecA protein. Using purified proteins, we examined the ATP hydrolysis activity of the K248R and E96D single mutants and the K248R/E96D double mutant RecA proteins. Mixtures of the K248R and E96D single mutant proteins were also examined. RecA-mediated ATP hydrolysis is DNA-dependent. Electron microscopy was used to help determine to what extent the observed rates of ATP hydrolysis, when reduced in the case of a mutant RecA protein, result directly from a catalytic impairment or indirectly from an inability to bind DNA and form nucleoprotein filaments. The effect of the reduced ATP hydrolysis rate on the ability of the mutants to catalyze the in vitro DNA three-strand exchange reaction was also examined. This allows us to uncover the contribution of nucleoprotein filament formation to the catalysis of ATP hydrolysis by the RecA protein and a poten-
Whether the Lys248 residue might act in nucleoprotein filament formation. From these initial observations, it was not clear whether the K248R mutant RecA protein. Reactions included 8 μM RecA protein and 6 μM poly(dT) linear ssDNA. The RecA protein was preincubated with the DNA for 10 min before dATP was added. Time 0 corresponds to the time of dATP addition. The average observed rates of dATP hydrolysis were 92.45 ± 1.91 and 0.57 ± 0.14 μM/min for wild type and K248R RecA protein, respectively.

The K248R RecA Mutant Does Not Hydrolyze ATP—The K248R mutant hydrolyzed ATP or dATP at a minimal rate relative to the wild type E. coli RecA protein (Fig. 2). The elimination of ATP and dATP hydrolysis indicated that the Lys248 residue is critical for catalysis. As described below, the K248R mutant was still able to form short nucleoprotein filaments on DNA, indicating that the mutation does not perturb the subunit-subunit interface enough to completely prevent filament formation. From these initial observations, it was not clear whether the Lys248 residue might act in cis or in trans to promote hydrolysis. A RecA nucleoprotein filament reconstruction suggests Lys248 acts in trans across the subunit-subunit interface.

A Combination of the E96D and K248R Mutations Partially Restores ATP Hydrolysis Activity—In an effort to partially restore ATP hydrolysis activity to the K248R filament, the K248R protein was mixed with the E96D mutant in different ratios. In such mixtures of the two single mutants, the resulting mixed filament would have four possible subunit-subunit interfaces, depicted in Fig. 3. Two adjacent K248R subunits would create an interface that mimics all interfaces of the K248R filament. Two adjacent E96D subunits would create a second interface, mimicking all interfaces of the E96D filament. Adjacent E96D and K248R subunits could align in two possible orientations: one mimics the E96D/E96D interface and the other mimics the wild type interface. We hypothesized that the wild type interface formed by the mixture of the E96D and K248R single mutants would hydrolyze dATP, thus partially rescuing catalysis of ATP hydrolysis. The E96D and K248R single mutants were mixed in varying ratios from 3:1 to 2:1 to 1:1. As shown in Fig. 4, the greatest rate of hydrolysis was observed for the 1:1 ratio of the two single mutants. The K248R and E96D subunits could be adjacent in the filament to mimic the E96D-E96D interface. The K248R and E96D subunits could be adjacent in the filament in two possible orientations: one would mimic the wild type RecA subunit-subunit interface, while the other would combine both the K248R and E96D mutations at the subunit-subunit interface. This last interface would mimic the subunit-subunit interface formed by the K248R/E96D double mutant RecA protein, depicted on the right.

The Mutant RecA Proteins Form Nucleoprotein Filaments—To confirm that the reduced rate of hydrolysis was not indirectly due to misfolding of the protein, circular dichroism spectra of the K248R/E96D double mutant were compared with that of the wild type RecA protein. Spectra of the mutant and wild type RecA proteins were identical, indicating that the K248R/E96D protein folded as well as wild type. Two different preparations of the K248R/E96D mutant yielded the same results (data not shown).

The DNA-independent rate of hydrolysis was examined. Wild type RecA, E96D, K248R, K248R/E96D, and the E96D+K248R mixture did not readily hydrolyze ATP or dATP in the absence of DNA (Fig. 5), indicating the observed nucleotide hydrolysis for wild type, K248R/E96D, and the E96D+K248R protein mixture was dependent on the presence of DNA.

We used electron microscopy to visualize wild type and mutant RecA proteins bound to DNA. As shown in Fig. 6, wild type RecA and the mutants formed filaments on poly(dT) at pH 7.5 with dATP. No fila-
with 30 – 45 min. No nicked circular product formation was observed for E96D, K248R, K248R/E96D, or the 1:1 mixture of E96D and K248R single mutants with ΦX174 DNA (data not shown). The partial restoration of catalysis of ATP and dATP hydrolysis by K248R/E96D and the 1:1 mixture of E96D and K248R were insufficient to promote complete strand exchange. It was previously known that nucleotide hydrolysis is required for DNA strand exchange for long DNA substrates (longer than 3 kbp), DNA strand exchange with four DNA strands, and strand exchange past barriers in the DNA such as an insertion of multiple DNA base pairs in one of the DNA substrates (4 – 8). Hence, the results support the notion that ATP hydrolysis must occur throughout the filament to complete nicked circular product formation with ΦX174 DNA molecules.

K248R/E96D Nucleoprotein Filament Is Unable to Displace SSB during Filament Extension—Nucleation of wild type RecA onto ssDNA is inhibited by SSB prebound to the DNA. Once RecA has nucleated onto the DNA, however, addition of SSB to the reaction stimulates extension of the RecA filament. SSB binds to and melts secondary structure in the DNA, thus enabling extension of the RecA filament. All experiments described above were completed with poly(dT) linear ssDNA that has a reduced propensity to form secondary structure, thus eliminating the need for SSB. With M13mp8 circular ssDNA, however, secondary structure in the DNA can form and adding SSB to the reaction after nucleation of wild type RecA then stimulates complete formation of the nucleoprotein filament and maximal ATP or dATP hydrolysis. In contrast, the K248R/E96D double mutant did not hydrolyze ATP or dATP with M13mp8 when SSB was added after a time sufficient to permit RecA nucleation, indicating a complete K248R/E96D filament did not form in the presence of SSB.

To examine the effect of SSB further, the dATP hydrolysis of wild type RecA and K248R/E96D were compared on M13mp8 with and without SSB. Reactions included RecA protein and M13mp8 circular ssDNA and were initiated by the addition of dATP. As shown in Fig. 7, both wild type RecA and K248R/E96D hydrolyzed dATP at a substantial but reduced rate compared with the maximal rate observed on poly(dT). Both wild type RecA and K248R/E96D were able to form significant nucleoprotein filaments on M13mp8 ssDNA, however, addition of SSB to the reaction stimulated extension of the nucleoprotein filament and maximal ATP or dATP hydrolysis. In contrast, the K248R/E96D double mutant did not hydrolyze ATP or dATP with M13mp8 when SSB was added after a time sufficient to permit RecA nucleation, indicating a complete K248R/E96D filament did not form in the presence of SSB.

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with the other could be direct or indirect. However, the results suggest an in trans participation of Lys\textsuperscript{248} and perhaps other residues in the ATP hydrolytic cycle. The cross-interface connection may help to organize ATP hydrolytic cycle-induced conformational changes between adjacent subunits.

Lys\textsuperscript{248} of E. coli RecA is conserved among bacterial RecA homologues.
addition of SSB protein. The RecA protein was added to the reaction; the hydrolysis by the wild type RecA protein; the 

\[
\text{Hydrolysis} \quad \text{by} \quad \text{wild} \quad \text{type} \quad \text{RecA} \quad \text{protein}
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\text{SSB was added to the reaction; the hydrolysis} \quad \text{by} \quad \text{the wild} \quad \text{type} \quad \text{RecA} \quad \text{protein}
\]

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\text{Recovery of ATP Hydrolysis in RecA K248R/E96D}
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\text{FIGURE 7. The extending K248R/E96D mutant RecA protein filament is compromised and does not displace SSB. RecA-catalyzed dATP hydrolysis was monitored. Reactions included 8 \mu M RecA protein, 6 \mu M M13mp8 circular ssDNA, 0.6 \mu M SSB (when included), and dATP. The RecA protein was preincubated with the DNA for 10 min before dATP was added to initiate the reaction. Time 0 corresponds to the time of dATP addition. For both RecA proteins, there are three conditions: SSB was added initially with dATP (contiguous dashed line), SSB was added ~20 min after dATP addition (solid line continued as dashed after SSB addition indicated by an arrow), or SSB storage buffer was added ~20 min after dATP addition (continuous solid line). The top three curves represent hydrolysis by the wild type RecA protein; the bottom three curves represent hydrolysis by the K248R/E96D RecA protein. The solid curves represent dATP hydrolysis before SSB protein was added to the reaction; the dashed lines represent dATP hydrolysis after the addition of SSB protein.}
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Some structural studies indicate that Lys\textsuperscript{248} is too distant to contribute to an ATP hydrolysis catalytic site within a RecA protomer (22, 23). However, a recent electron microscopic reconstruction of the RecA-dsDNA nucleoprotein filament (21), coupled to the overall subunit-subunit interface seen in recently determined structures of the RecA homologues Rad51 (25) and RadA (24), suggests that Lys\textsuperscript{248}, along with several other residues, lies in proximity to the nucleotide binding site of the adjacent subunit. Hence, Lys\textsuperscript{248} may contribute to nucleotide hydrolysis in the neighboring subunit in trans. Further support of this notion is the discovery of the \([\text{KR}]\times[\text{KR}]\) motif in RecA and helicases including T7 gp4 that is postulated to be involved in in trans catalysis of ATP hydrolysis and communication of the ATP hydrolytic cycle between subunits or domains (13–16, 18–20). Lys\textsuperscript{248} is the first residue in the \([\text{KR}]\times[\text{KR}]\) motif of \textit{E. coli} RecA (13). The effects of a conservative mutation introduced at residue 248, K248R, indicate Lys\textsuperscript{248} is crucial for ATP hydrolysis but at least partly dispensable for DNA binding and filament formation. The Glu\textsuperscript{96} residue is part of the ATP binding site within a single RecA subunit, and the E96D mutant also does not readily hydrolyze ATP. A mixture of K248R and E96D restored ATP hydrolysis activity to the nucleoprotein filament. Complementation of E96D and K248R together with the location of these residues at the subunit-subunit interface in the nucleoprotein filament illustrates the importance of this interface to nucleotide hydrolysis and suggests an in trans catalysis mechanism of nucleotide hydrolysis whereby the adjacent RecA subunit contributes catalytically.

Both the K248R and E96D single mutants of RecA protein will bind to ssDNA and form filaments. The K248R mutant has a reduced capacity to tandemly and noncovalently link short filaments to create longer ones, indicating a somewhat compromised subunit-subunit interface. Nevertheless, the capacity of both mutant proteins to bind to poly(dT) suggests that their inability to hydrolyze ATP or dATP is primarily due to a catalytic defect in ATP hydrolysis and not indirectly from an inability to form a filament on DNA. Despite K248R and E96D both being deficient in ATP and dATP hydrolysis activity, partial dATP hydrolysis activity is restored (to 25% of the wild type RecA hydrolysis rate) when these two mutants are mixed at a 1:1 ratio. Perhaps more surprisingly, the K248R/E96D double mutant protein promotes dATP hydrolysis at a rate that is more than 10-fold greater than either of the single mutants alone when bound to poly(dT). The restored hydrolysis is due at least in part to the formation of some wild type subunit-subunit interfaces that hydrolyze dATP or ATP when the K248R and E96D single mutants are mixed. However, the observation that the K248R/E96D double mutant hydrolyzes ATP and dATP indicates that the subunit-subunit interface formed when the K248R and E96D mutations are opposed to each other across the interface is also at least partially enabled for hydrolysis. The Glu\textsuperscript{96} residue may activate the water molecule for nucleophilic attack on ATP hydrolysis (22). An aspartate at this location would necessarily affect this reaction. The presence of an arginine at position 248 may perturb the structure at the interface to bring the γ-phosphate of the ATP molecule and the mutant aspartate residue at position 96 in closer proximity to the nucleophilic water molecule. Conceivably the K248R mutation, when present with the normal Glu\textsuperscript{96} across the interface, could destabilize the interface and prevent ATP hydrolysis. We do not know if the distribution of the two single mutant proteins in the mixed filament is random but would argue that it is unlikely that the distribution of mutant proteins would leave the wild type subunit-subunit interfaces underrepresented. If 25% of the interfaces formed mimic the wild type interface to hydrolyze dATP (Fig. 3), and 25% of interfaces mimic the K248R/E96D interface to hydrolyze dATP as well, then the wild type interfaces in the mixed filaments must be hydrolyzing dATP at less than 100% of the wild type efficiency to arrive at the observed rates. The results thus suggest that longer range coordination of ATP hydrolytic cycles occurs in the filaments; adjacent subunit-subunit interfaces affect each other. This conclusion suggests the importance of contiguous hydrolysis-enabled interfaces in the RecA nucleoprotein filament.

An extending K248R/E96D filament on ssDNA does not displace SSB, whereas wild type RecA is able to displace SSB and extend a filament. The K248R mutation must introduce instability at the subunit-subunit interface, despite the cooperation with E96D. The less robust interaction at the subunit-subunit interface impedes filament formation against a barrier, such as SSB.

These results indicate that each of the two single mutants, K248R and E96D, has a distinct catalytic defect and partial restoration of nucleotide hydrolysis results from a complementary interaction of the K248R and E96D residues between adjacent subunits in trans. The results suggest that the Lys\textsuperscript{248} residue has an intermolecular catalytic role in nucleotide hydrolysis in the RecA nucleoprotein filament. Coordination of ATP hydrolysis between adjacent subunits as observed recently (9) may manifest through Lys\textsuperscript{248}. The Lys\textsuperscript{248} residue of \textit{E. coli} RecA presents another example of the \([\text{KR}]\times[\text{KR}]\) motif acting across a subunit or domain interface to catalyze ATP hydrolysis and potentially direct conformational changes dependent on the nucleotide state. This is a developing mechanism for motor ATPase proteins with RecA-like protein folds.

The lack of a \([\text{KR}]\times[\text{KR}]\) motif in the archaeal and eukaryotic members of the RecA family is intriguing. RecA promotes a number of DNA transactions, all requiring ATP hydrolysis, which the RadA, Rad51, and Dmc1 proteins are not able to promote. The \([\text{KR}]\times[\text{KR}]\) motif may provide an important clue to help understand the molecular basis of this distinction.

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References