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Two Modes of Binding of Dinl to RecA Filament Provide a New Insight Into Regulation of SOS Response by Dinl Protein

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Abstract

The RecA protein plays a principal role in the bacterial SOS response to DNA damage. The induction of the SOS response is well understood and involves the cleavage of the LexA repressor catalyzed by the RecA nucleoprotein filament. In contrast, our understanding of the regulation and termination of the SOS response is much more limited. RecX and DinI are two major regulators of RecA's ability to promote LexA cleavage and a strand exchange reaction and are believed to modulate its activity in ongoing SOS events. DinI's function in the SOS response remains controversial since its interaction with the RecA filament is concentration-dependent and may result in either stabilization or depolymerization of the filament. The 17 C-terminal residues of RecA modulate the interaction between DinI and RecA. We demonstrate that DinI binds to the active RecA filament in two distinct structural modes. In the first mode DinI binds to the Cterminus of a RecA protomer. In the second mode DinI resides deeply in the groove of the RecA filament with its negatively charged C-terminal helix proximal to the L2 loop of RecA. The deletion of the 17 C-terminal residues of RecA favors the second mode of binding. We suggest that the negatively charged C-terminus of RecA prevents DinI from entering the groove and protects the RecA filament from depolymerization. Polymorphic binding of DinI to RecA filaments implies an even more complex role of DinI in the bacterial SOS response.

Introduction

RecA is a key player in maintaining the integrity of the bacterial genome. It is crucial for DNA repair via homologous recombination and required for the induction of the SOS response in bacteria^{1–3}. The RecA protein of *E. coli* contains 352 amino acid residues and consists of a large core domain and a smaller C-terminal domain (residues 271-352). The last 24 residues were found to be disordered in all crystal structures^{4–8}, except the *M. smegmatis* RecA complexed with dATP⁹. Most bacterial RecA proteins contain a high concentration of negatively charged residues at their C-terminus¹⁰.

Double-stranded DNA (dsDNA) breakages or stalled replication forks produce regions of single-stranded DNA (ssDNA). Polymerization of RecA monomers on ssDNA initiates

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transcription of more than 40 proteins involved in the SOS response in bacteria¹¹. Synthesis of these proteins is normally inhibited by the LexA repressor which can cleave itself¹² upon binding within the helical groove of the RecA-DNA filament¹³. Also, the RecA-DNA filament acts as an ATP-dependent recombinase and is capable of promoting DNA pairing and strand exchange¹⁴. The repair of the damaged dsDNA results in degradation of ssDNA and depolymerization of the RecA nucleoprotein filaments which in turn restores the pool of LexA repressor and shuts down the SOS system.

To prevent undesired DNA rearrangements, bacteria developed a sophisticated system to control the formation of the RecA-ssDNA filament. RecA activity is modulated by its C-terminus in a Mg²⁺-dependent fashion¹⁵. Deletion of the 25 C-terminal residues results in faster RecA nucleation¹⁶. The 17 C-terminal RecA residues prevent binding of RecA to dsDNA, favoring RecA polymerization on ssDNA, and modulate RecA's ability to displace SSB protein from ssDNA¹⁷. Recently crystal structures of the RecA-ssDNA/dsDNA filaments have been solved¹⁸, but these did not provide any structural information about the RecA C-terminus because this region was used as linker to construct a RecA polypeptide.

The other two well known modulators of RecA filaments are RecX and DinI. RecX protein is an intrinsic inhibitor of RecA activities, and overexpression of RecA in the absence of the *RecX* gene is toxic for bacteria^{19,20}. RecX inhibits the RecA-dependent strand exchange reaction and co-protease activity by means of slow depolymerization of RecA-DNA filaments^{21,22}. Similarly to LexA, RecX binds deep in the helical groove of RecA filaments¹³. The competition between LexA and RecX for binding within the RecA helical groove may contribute to inhibition of LexA cleavage by RecX. Removal of the 17 C-terminal RecA residues (RecA Δ C17) significantly alters the ability of RecX to inhibit the strand exchange reaction and DNA-dependent ATPase activity²³.

Another member of the SOS regulon is the DinI gene which encodes a small protein containing 81 amino acid residues^{24,25}. Originally it was shown that the over-expression of DinI protein conferred severe UV sensitivity on wild-type cells and resulted in the inhibition of LexA and UmuD cleavage²⁶. Later it was suggested that under normal expression levels DinI would inhibit UmuD cleavage so as to limit SOS mutagenesis, while having little effect on LexA processing²⁶. The role of DinI as a down-regulator of the SOS response was also supported by the observation that the maximum binding of DinI to RecA filaments occurred at later stages of the SOS response²⁷. It was proposed that a negatively charged C-terminal helix in DinI could compete with ssDNA for binding to the L2 loop of RecA, which in turn would displace the ssDNA from the RecA nucleoprotein filament and inhibit the strand exchange reaction²⁷. At a stoichiometric ratio of DinI to RecA, DinI has a substantial stabilizing effect on the RecA filament²⁸. It also allows LexA cleavage and does not interfere with an on-going strand exchange reaction. The deletion of 17 C-terminal residues of RecA has a dramatic effect on the RecA-DinI interaction²⁸ – it alters DinI affinity for the RecA filament. In contrast to the wild type RecA, RecA Δ C17 has reduced strand exchange activity when DinI is present. We used electron microscopy and image analysis to locate the binding site of DinI on the RecA filament and to explore the structural consequences of the deletion of the 17 C-terminal residues of RecA on its ability to interact with DinI.

Results and discussion

The formation of the RecA filament requires ATP-binding, but ATP hydrolysis is not required for the co-protease activity²⁹ or for the catalysis of the strand exchange reaction³⁰. In the presence of ATP RecA forms disordered filaments due to protomer dissociation upon ATP hydrolysis. Such filaments are inappropriate for image analysis. We have shown previously that the long and ordered RecA filaments formed in the presence of non-

hydrolysable ATP analogs are in an active extended conformation³¹. In this study we used AMP-PNP to form RecA-DNA or RecA Δ C17-DNA filaments alone (Fig. 1a and b), or in the presence of DinI (Fig. 1c and d). We extracted 6,381 segments of RecA and 5,198 of RecAAC17 to generate 3D-reconstructions of those filaments (Fig 2a). Both sets converged to a pitch of ~92 Å consistent with the extended "active" form of the filament. Paradoxically, wild type RecA reconstruction possessed a weaker electron density in the Cterminal domain (CTD) region than the RecA Δ C17 reconstruction (Fig 2a, red arrowheads), but otherwise the two reconstructions were similar. This was consistent with our previous observations that the deletion of 18 C-terminal residues resulted in conformational changes in the CTD³². Recently crystal structures of RecA-ssDNA/dsDNA filaments have been solved¹⁸. Since our filaments were formed on dsDNA, we used the high resolution structure of the RecA-dsDNA filament (PDB 3CMT) to evaluate the reconstruction of the RecA Δ C17 filament. At a resolution of ~23 Å no perturbations of the original crystal structure were required to dock it into the electron density map (Fig 2b). This suggests that no large scale conformational changes to the RecA protomers within the extended RecA filament take place upon the deletion of the 17 terminal residues. The alignment of the RecA protomer from the RecA-dsDNA heteroduplex filament with the crystal structure of the M. smegmatis RecA⁹ places the missing 17 residues in the region spanning from the outer portion of the CTD to the channel in between two adjacent RecA protomers (Fig 2b, cyan spheres).

Incubation of the RecA or RecA Δ C17 filaments with DinI resulted in the disappearance of the helical pattern (Fig 1c,d) suggesting that DinI was bound to these filaments. We collected 5,891 segments of wt-RecA and 5,773 segments of RecA Δ C17 filaments decorated with DinI. The 3D-reconstructions of the two sets are shown in Fig 3b and c, respectively. The wt-RecA-DinI reconstruction (Fig 3b) showed two regions of density absent in the map of the pure RecA Δ C17 (Fig 3a) – the first site was an attachment to the CTD (Fig 3b, red arrow), while the second site was a continuous density located in the helical groove was observed in the RecA Δ C17-DinI reconstruction (Fig 3c, blue arrow), though no binding of DinI to RecA's CTD was detected in the overall reconstruction. It was unclear whether the two regions of binding of DinI found in the RecA-DinI reconstruction, or represented a mixture of two distinct modes.

The single particles approach to helical reconstruction allows us to sort segments of filaments by the occupancy and the mode of binding. We generated three model volumes (Fig 3d) – naked RecA filament, a RecA filament having additional density at its CTD (Fig 3d, red arrowhead), and a RecA filament with a continuous density in the helical groove (Fig 3d, cyan arrowhead). The three model volumes were projected and cross-correlated with the images of RecA-DinI and RecA Δ C17-DinI complexes. The frequency distribution for such a sorting is shown in Fig 3d as grey and black bars, respectively. The majority of the RecA-DinI images (~54%) had the highest correlation with the model having additional density attached to the CTD, while a quarter of segments were assigned to the "groove" class (Fig 3d, grey bars). The deletion of the 17 C-terminal residues did not alter the frequency of the groove mode, but significantly reduced the population of the "CTD" class (Fig 3d, black bars). Each class was iterated in the IHRSR procedure starting from a solid featureless cylinder until it converged to a stable solution. The resultant reconstructions are shown in Fig 3e. The poorly decorated RecA-DinI filaments yielded a reconstruction possessing weak traces of DinI (Fig 3e, black arrow), while the same class from the RecAAC17-DinI complex resulted in a map having continuous additional density in the helical groove (Fig 3e, cyan arrow). This suggested that the cooperativity of DinI binding to RecA was higher than that to RecA Δ C17. The CTD class reconstructions had a substantial additional mass attached to the CTD (Fig 3e, red arrows). The groove classes from both sets yielded maps

having substantial continuous density in the helical groove of the RecA filament (Fig 3e, blue arrows). Despite similar frequencies of the groove classes (Fig 3d), and because of the substantial additional density in the helical groove of the naked RecA Δ C17-DinI class reconstruction, our sorting suggested that the groove of the RecA Δ C17 filaments was more attractive to DinI than that of unmodified RecA filaments.

Since each class was reconstructed starting from a featureless solid cylinder, the details of the reconstructions reflected the real modes of interaction of DinI with the RecA filament rather than being model-biased solutions. We therefore used the reconstructions of the CTD and the groove classes of the RecA Δ C17-DinI complex to build atomic models of the two structural modes. The RecA protomer from the crystal structure of the RecA-dsDNA filament¹⁸ was used to model the position of RecA protomers in the RecA-DinI complex, while keeping the interface between the adjacent RecA protomers similar to that in the original crystal. Three consecutive RecA protomers docked into the density map are shown in Fig 4 as magenta, cyan, and orange ribbons. Docking of RecA into the two decorated maps allowed us to determine the density attributed to the DinI in each reconstruction. In both modes of binding the additional density perfectly matched the size and the shape of the DinI molecule (Fig 4, green ribbons). The models suggested that the two modes were mutually exclusive. That is, a DinI molecule could not be bound to the same RecA protomer in both modes without extensive steric clashes. If DinI were randomly bound to RecA filaments at either of the two sites, we would never be able to sort out the two discrete modes that we observe. Instead, our results are consistent with a cooperativity of binding so that within the lengths of the segments we are analyzing (~ 22 RecA subunits) there is predominantly one mode present. What could be responsible for such cooperativity? In the groove mode (Fig 4b) there appears to be an interaction between adjacent DinI molecules bound to RecA actin subunits, and such a DinI-DinI interaction could contribute to the cooperative nature of the binding. Since in the CTD-mode the two DinI molecules do not interact with each other, it is likely that in that particular mode the cooperativity in the interaction must reside mainly within RecA filament. That is, when DinI binds to one RecA protomer in CTD mode, structural changes in RecA filament greatly increase the likelihood that DinI will bind in this same manner to adjacent RecA subunits. This is consistent with many, many papers showing such internal cooperativity within RecA filaments^{33,34}.

In general, the docking of a high resolution structure of a protein into a 23 Å resolution map is quite ambiguous³⁵. Due to a substantial asymmetry of the RecA and DinI molecules when filtered to 23 Å, the orientation of both molecules in the resultant atomic models was unambiguous. In addition, the two models are supported by DinI mutagenesis studies which show that substitution of Arg 43 of DinI significantly reduces its ability to stimulate the RecA ATPase activity (Supplementary data). In both atomic models that residue is located in proximity to the RecA CTD (Fig 4a,b, red spheres).

We used the recent crystal structure of the *M. smegmatis* RecA⁹ to reveal the position of the 17 C-terminal residues in our atomic models (Fig 4, yellow spheres). Interestingly, in both modes the C-terminus was located far from the DinI molecule (Fig 4a,b). The removal of the 17 C-terminal residues of RecA did not alter the mode of binding of DinI to RecA; rather it only changed the frequency of the existing modes (Fig 3d), suggesting that the C-terminus is not involved in direct interaction with DinI. Why then does the deletion of these C-terminal residues alter the interaction between RecA and DinI? Most of the bacterial RecA proteins contain a high concentration of negatively charged residues at their C-terminus¹⁰. In all but one RecA crystal structure this region of RecA has been found to be disordered. The DinI molecule has an extended negatively charged ridge³⁶ in its C-terminal helix. If the C-terminus of RecA is located in the groove of the filament, electrostatic forces may preclude extensive binding of DinI in the helical groove and favor the CTD mode of binding.

Alternatively, the deletion of the C-terminus may cause conformational changes in RecA, and this could alter the frequency of the two modes. This is consistent with our observations that the deletion of the 17 C-terminal residues stabilizes the CTD of RecA (Fig 2a). In addition, the extreme C-terminal residues could be making a direct favorable contact with DinI increasing the frequency of this mode. A high resolution structure of the RecA-DinI complex is required to distinguish among these possibilities.

It has been shown that at low concentration DinI can stabilize the RecA filament without interfering with LexA cleavage and an ongoing strand exchange reaction²⁸. When DinI binds to RecA in the CTD mode, it bridges adjacent RecA protomers across the helical groove (Fig 4a, red arrows) contributing to the intrafilament contacts. Importantly, being bound in this mode leaves the groove available for LexA cleavage¹³ and for the invasion of dsDNA into the groove.

An extensive interaction between the adjacent DinI molecules in the groove mode may increase the affinity of DinI for the RecA filament. Because of the preferential binding of DinI to RecA Δ C17 in the groove mode, one would expect the RecA Δ C17-DinI complex to be more stable that the RecA-DinI one. Strikingly, the RecA Δ C17-DinI complex was shown to be 100 times more stable than the one formed with unmodified RecA²⁸.

When DNA in the atomic model of the RecA Δ C17 filament is positioned as in the crystal structure¹⁸, it is proximal to the L2 loop region of RecA (Fig 5b, red spheres) which has been implicated in DNA binding ^{37,38} and homologous pairing³⁹. The solution structure of DinI shows that six Asp/Glu residues in the C-terminal helix form an extended negatively charged ridge³⁶ (Fig 5c–f, cyan spheres), and this ridge was proposed to compete with ssDNA for binding to the RecA L2 region²⁷. In the CTD mode the negatively charged region of DinI is located at the outer surface of the RecA filament (Fig 5d, cyan spheres), and is far from the L2 loop of RecA (Fig 5d, red spheres). In the groove mode the negatively charged C-terminal helix forms a continuous helical path (Fig 5f, cyan spheres) flanking the L2 region of RecA (Fig 5f, red spheres). Our data show that when DinI is bound to RecA filament in the groove mode, it can compete with the DNA for binding to the L2 loop. This interaction may facilitate the dissociation of RecA subunits from the nucleoprotein filament.

RecX protein is a suppressor of the SOS response and inhibits LexA cleavage as well as the strand exchange reaction by means of depolymerization of RecA filaments^{21,23}. When the RecA filament is stabilized with AMP-PNP, RecX produces a continuous density that forms a block across the deep helical groove of the RecA filament¹³. Such a mode of binding intuitively explains how RecX can inhibit the LexA cleavage by simply blocking the penetration of LexA into the groove of the filament. It has been suggested recently that the positively charged concave surface of RecX can make an extensive contact with the DNA phosphate backbone⁴⁰. DinI binds to RecA Δ C17 filaments presumably in the groove mode, and its position on the RecA filament in that mode (Fig 4b) is very similar to that found for RecX protein¹³. Consistent with this picture, DinI inhibits the ability of RecA Δ C17 to catalyze the strand exchange reaction²⁸. RecX contains three small domains, each containing three helices. The protein appears to have been the result of a gene triplication, as the three domains show strong structural homology. The two helices in DinI align extremely well with two of the three helices in each of the RecX domains, and the fit of DinI to the central domain of RecX is shown (Fig 6). Whether this represents a convergent evolution or distant homology cannot be determined at this time.

Our data show that the C-terminus of RecA is involved in modulation of the RecA-DinI interaction. In the groove mode DinI resides deeply in the groove of the RecA filament with its negatively charged C-terminal helix proximal to the L2 loop of RecA. This supports the

original suggestion that the negatively charged C-terminal helix of DinI mimics DNA and interferes with the activation of RecA²⁷. We suggest that the negatively charged C-terminus of RecA prevents DinI from entering the groove and protects the RecA filament from depolymerization, since the deletion of the 17 C-terminal residues of RecA favors the groove mode of binding. How can such a regulatory mechanism work in the cell lacking truncated RecA protein? The deletion of the 17 C-terminal residues in RecA *in vitro* may mimic conformational changes in RecA filament that otherwise are triggered by regulatory proteins such as RecO and RecR. The enhanced binding of DinI to RecA at late stages of the SOS response²⁷ strongly argues for the existence of one or more modulators of the DinI-RecA interaction, since DinI synthesis starts at early stages of the SOS response.

Materials and Methods

Cloning

To clone an *E. coli dinI* gene that encodes the R43A mutant protein, the wild-type *dinI* gene in pEAW334 was used as the template in a PCR with the upstream primer consisting of the *dinI* gene starting at the BsaAI restriction enzyme site at base 115 and including at least 10 bases after the changed bases. For R43A, the CGT starting at base 126 of the *dinI* gene was changed to GCT. The downstream primer consisted of an EcoRI restriction enzyme site and the last 19 bases of the *dinI* gene. The PCR product was digested with BsaAI and EcoRI and ligated to pEAW340 cut with the same restriction enzymes. The plasmid designated pEAW340 is the wild-type *dinI* gene inserted into pUC19 sites. The presence of the *dinI* mutation was confirmed by direct sequencing of the resulting plasmids. The mutant *dinI* gene was then subcloned into pET21A cut at the NdeI and EcoRI sites, and the *dinI* genes of the resulting plasmids were again directly sequenced. The plasmid with *dinI* R43A was designated pEAW533.

To create His-tagged versions of the mutant and wild type DinI proteins, primers (5'-CCTAAGCCATATGCGAATTGAAGTCACC-3' and 5'-

GGATATCCTCGAGTTCGCTGACAACCAGTC-3') were used to amplify mutant *dinI* genes from their respective plasmids (pEAWs 533 and 334) and append flanking NdeI and XhoI restriction sites. PCR products and pET-21b(+) (Novagen) were cut with NdeI and XhoI and were ligated. The resulting plasmids were sequenced to ensure the integrity of the *dinI* coding region. The above cloning procedure resulted in mutant *dinI* genes with six His codons at the 3' end of the *dinI* coding sequence followed by a stop codon.

DNA substrates

The poly-dT DNA was purchased from Amersham Biosciences. Its average length was 319 nucleotides. The concentration of poly-dT was reported in terms of μ M nucleotides (nt).

Proteins

Untagged wild type DinI pr otein was over-expressed in *E. coli* STL2669/pT7pol26 cells. Ten liters of culture were grown in LB to an OD₆₀₀ of 0.5, and DinI protein expression was induced with 0.4 mM isopropyl-1-thio- β -D-galactopyranoside. Approximately 15 to 20 g of cells were harvested by centrifugation following 3 hours of incubation at 37 °C. Cells were flash-frozen in liquid N₂, and stored at -80° C. Cells were thawed over-night on ice in buffer containing 250 mM Tris-Cl (80% cation, pH 7.8) and 25 % w/v sucrose. The cells were resuspended to 20 % w/v and lysed via sonication with a Fisher Scientific Sonic Dismembrator Model 500 at 60% output, 5 times, for 1 min segments with alternating 0.5 s on/off pulses. Lysed cells were centrifuged for 60 min at 16,000 rpm in a Beckman JLA-16.25 rotor at 4°C. The lysis supernatant was dialyzed against P buffer (10 mM KH₂PO₄, 10 mM K₂HPO₄, 0.1 mM EDTA, 10% w/v glycerol, 1 mM DTT) and run over a

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ceramic hydroxyapatite column. Wild type DinI protein was not retained by the hydroxyapatite and the wash fractions containing DinI were dialyzed against R buffer (20 mM Tris-Cl 80% cation, pH 7.8, 0.1 mM EDTA, 10% w/v glycerol, 1 mM dithiothreitol (DTT)). The DinI was then applied to a Q-sepharose anion exchange column and eluted with a linear gradient from R-buffer to R-buffer + 1 M KCl. Gradient fractions containing DinI protein were dialyzed back into R-buffer and run over a heparin sepharose column. Heparin sepharose flowthrough fractions containing DinI protein were concentrated on a 1 ml Q-sepharose column with a step gradient from R-buffer to R + 1 M KCl. The concentrated DinI was run over a Hi-Prep Sephacryl 16/60 S-100 size exclusion column in R + 200 mM KCl. Fractions containing pure wild type DinI protein were pooled and dialyzed into R-buffer for storage. The wild type DinI protein was free of detectable ssDNA nuclease activity.

To over-express the His-tagged DinI proteins, plasmids bearing the dinI genes were transformed into the E. coli STL327 (DE3) strain which was grown, harvested and frozen as described above for untagged wild type DinI. Five ml of lysis buffer (50 mM potassium phosphate, 300 mM NaCl, 10 mM imidazole, 10% w/v glycerol, pH 8.0) were added per gram of cell pellet to resuspend the pellet. Cells were thawed overnight, on ice, with gentle stirring. Protease inhibitors E-64 and Leupeptin at 0.5 μ g/ml, and Pepstatin at 1 μ M, were added to fully resuspended cells. The cells were lysed at 4°C for 2 hours with lysozyme at a final concentration of 4 mg/ml. Cells were sonicated and centrifuged as described above. The supernatant was incubated in batch with ~5 ml Ni-NTA slurry pre-equilibrated with lysis buffer for 1 h at 4°C. The slurry was poured into a solid support, then washed with 30 column volumes of wash buffer (50 mM potassium phosphate, 0.3 M NaCl 20 mM imidazole, 10% w/v glycerol, pH 8.0). Histidine-tagged DinI protein was eluted with 5 column volumes of elution buffer (same as wash buffer except with 500 mM imidazole). Wild type histidine-tagged DinI (WT His-DinI) was dialyzed against R buffer and eluted from a 1 ml Q-sepharose column with a linear gradient from R buffer to R-buffer + 1 M KCl. Fractions containing WT His-DinI were run over a Hi-Prep Sephacryl 16/60 S-100 size exclusion column and a ceramic hydroxyapatite column, and concentrated on a 1 ml Qsepharose column with a step gradient as described above. Eluted WT His-DinI protein was dialyzed against R buffer for storage. His-R43A DinI protein was purified as for the wild type His-tagged protein, except that it was not concentrated on the 1 ml Q-sepharose after the ceramic hydroxyapatite column. All purified proteins were flash frozen with liquid N_2 and stored at -80° C. Each was determined to be pure by SDS-PAGE and had no detectable ssDNA nuclease activity at 20 µM DinI after 2 hours of incubation at 37 °C. The theoretical extinction coefficient of 1.44×10^4 M⁻¹ cm⁻¹ at 280 nm for wild type DinI calculated using the ProtParam tool (from the ExPASy web site) was used to determine the concentration of all the DinI proteins.

Each DinI protein was diluted in R-buffer to a concentration between 0.1 and 0.2 mg/ml and was submitted to the Biophysics Instrumentation Facility at UW-Madison to check for proper folding by circular dichroism (CD). All data were collected in 0.1 cm cuvettes at 37 °C, with a 1 nm bandpass at 1 nm intervals with a 5 second averaging time at each point. An R-buffer blank was recorded and subtracted from each spectrum. Conversion to molar ellipiticity was carried out using DinI concentrations measured as described above and a mean residue weight of 110 Da. The CD spectra for the DinI proteins were normalized by arbitrarily setting the molar ellipiticity at 218 nm to $-1 \text{ deg cm}^2 \text{ decimol}^{-1}$.

ATP hydrolysis assays to monitor the effect of the Dinl mutant proteins on RecA

A spectrophotometric, coupled-enzyme assay was used to measure RecA-catalyzed ATP hydrolysis^{41,42}. The ADP produced by hydrolysis was converted back to ATP by the enzyme pyruvate kinase and the substrate phosphoenolpyruvate. The resultant pyruvate was

reduced to lactate by lactate dehydrogenase using NADH as a reducing agent. The conversion of NADH to NAD⁺ was monitored as a decrease in absorbance at 380 nm on a Varian Cary 300 instrument equipped with a temperature controller and 12 position cell changer. The cell pathlength was 0.5 cm. The concentration of oxidized NADH, and thus the concentration of hydrolyzed ATP, was calculated from the NADH extinction coefficient $\epsilon_{380} = 1.21 \text{ mM}^{-1} \text{ cm}^{-1}$.

Each ATP hydrolysis assay contained a reaction mixture of 25 mM Tris.OAc (88% cation, pH 7.5), 10 mM magnesium acetate, 4 mM potassium glutamate, 5 % (w/v) glycerol, 1 mM DTT, 3.5 mM phosphoenolpyruvate, 10 U/ml pyruvate kinase, 10 U/ml lactate dehydrogenase, 1.5 mM NADH, 3 μ M poly-dT, and 3 mM ATP. DinI proteins, or compensating storage buffer, were incubated with the above reaction mixture at 37 °C for 10 min. The DinI concentration was varied from 0 to 20 μ M. Wild type RecA protein was then added to a final concentration of 1 μ M. RecA-catalyzed ATP hydrolysis was monitored at 37 °C until the NADH was exhausted. The k_{cat} for RecA-catalyzed ATP hydrolysis divided by the concentration of RecA binding sites on the poly-dT.

Electron microscopy and image analysis

The RecA-dsDNA-AMPPNP, and RecA C17-dsDNA-AMPPNP complexes were incubated in 25 mM triethanolamine-HCl (Fisher) buffer (pH 7.2) at 37° C for 10 min, with [RecA] or $[\text{RecA C17}]=2 \,\mu\text{m}$, RecA to calf thymus double-stranded DNA (Sigma) ratio of 40:1 (w/w), AMP-PNP (Sigma) 1.5 mM, magnesium acetate (Sigma) 2.5 mM. The DinI complexes were formed in the same manner except that 20 uM DinI was present. Samples were applied to carbon-coated grids and stained with 2% (w/v) uranyl acetate (w/v). Images were recorded on film using a Tecnai 12 electron microscope operating at 80 keV with a nominal magnification of ×30,000. Negatives were scanned with a Nikon Coolscan 8000 densitometer, at a raster of 4.2 Å per pixel. SPIDER⁴³ was used for image processing. Segments of filaments (80×80 pixels) of the RecA (n=5.300) or RecA $\Delta 17$ (n=5.301) formed on double-stranded DNA (dsDNA) in the presence of AMP-PNP were extracted from the micrographs and reconstructed using the IHRSR method⁴⁴. Both sets converged to the same helical parameters and yielded a twist of 58.5° and axial rise of 15 Å. Segments (80 \times 80 pixels) of RecA-dsDNA-AMP-PNP (n=5,891) or RecA Δ 17-dsDNA-AMP-PNP (n=5,773) decorated with DinI protein were extracted from micrographs, and overall reconstructions were calculated (Fig. 3b and c, respectively). Both sets converged to the same helical symmetry of $59^{\circ}/15$ Å with the IHRSR method. The central protomer from the RecA-dsDNA crystal structure (PDB 3CMU) was used to generate a model RecA filament. The DinI solution structure (PDB 1GHH) was used to build up two models - with the DinI attached to the CTD (Fig. 3d, red arrowhead), or located deeply in the groove of the RecA filament (Fig. 3D, cyan arrowhead). The model of the RecA filament along with the two decorated models were used as references in cross-correlation sorting. The three reference volumes were scaled to 4.2 Å per pixel and projected into 80×80 -pixel images with an azimuthal rotational increment of 4°, generating 270 reference projections (3 × 90). RecAdsDNA-AMP-PNP-DinI and RecAA17-dsDNA-AMP-PNP-DinI segments were crosscorrelated with the 270 reference projections. Reconstructions were independently generated for each class (Fig. 3E) starting from a featureless solid cylinder. The two naked classes (n=1,375 and 3,683, respectively) both converged to the symmetry of $59^{\circ}/15$ Å. The two CTD classes (n=3,094 and 766, respectively) yielded symmetries of 58.7°/14.7 Å and 58.4°/ 14.4 Å, while the groove classes (n=1,422 and 1,324, respectively) converged to 58.8°/14.6 Å and $58.5^{\circ}/14.8$ Å helical parameters. The resolution of the reconstructions was judged to be ~23 Å using the FSC = 0.5 criterion.

UCSF Chimera software⁴⁵ was used to fit the high resolution structures of RecA and DinI into the experimental density maps. Atomic coordinates from crystal structures were converted to density maps, and these were filtered to the resolution of the experimental map and docked manually.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Fig. 1.

Electron micrographs of RecA-DNA filaments (a) and RecA Δ C17-DNA filaments (b) show strong periodic striations that arise from the one-start right handed RecA filament helix. These striations disappear when DinI is added to wt-RecA (c) or RecA Δ C17 filaments (d).



Fig. 2.

3D-reconstructions of RecA and the RecA Δ C17 filaments (a) are similar in the core region, but differ in the C-terminal domain (CTD) region (red arrowheads). The crystal structure of the RecA-dsDNA filament¹⁸ fits without perturbation into the RecA Δ C17 reconstruction (b). The RecA protomers are shown as blue ribbons, while the dsDNA molecule is in red. The position of the 24 C-terminal residues found ordered in the crystal structures of the *M*. *smegmatis* RecA⁹ are shown as cyan spheres.



Fig. 3.

Comparisonof the 3D-reconstructions of pure RecA Δ C17 filament (a), RecA-DinI (b) and RecA Δ C17-DinI (c) complexes reveals additional density bound to the C-terminal domain of RecA (red arrow) along with the continuous density residing in the helical groove (blue arrows). Cross-correlation sorting using model volumes (d, yellow surfaces) revealed differences in the frequency of the two modes of DinI binding to RecA (d, grey and black bars). 3D-reconstructions of each of the six classes shown in (d) are shown in (e) as solid surfaces. While the reconstructions of the CTD (red arrows) and the groove (blue arrows) modes from the RecA-DinI and RecA Δ C17-DinI complexes are similar, the reconstructions of the naked classes differ. Poorly decorated segments from RecA-DinI filaments have few traces of the DinI bound (black arrow), while such filaments extracted from the RecA Δ C17-DinI complex possess continuous density in the helical groove (cyan arrow).



Fig. 4.

3D-reconstructions of the CTD (a) and the groove (b) modes are shown with three consecutive RecA protomers docked into the electron density maps (magenta, cyan, and orange ribbons). Three DinI molecules in each mode are shown as green ribbons. The 17 C-terminal residues present in the crystal structure of the *M. smegmatis* RecA⁹ are shown as yellow spheres. In the CTD mode the DinI protein bridges adjacent RecA protomers across the helical strand (red arrows). Mutation in R43 (red spheres) significantly diminishes the ability of DinI to stimulate ATP hydrolysis by RecA.



Fig. 5.

In the atomic model of the pure RecA filament (a, blue ribbons) the dsDNA molecule (b, black ribbons) is proximal to the L2-loop of the RecA (b, red spheres). When DinI binds to RecA in the CTD mode (c,d, green ribbons) the six Asp/Glu residues in the C-terminal helix that form an extended negatively charged ridge³⁶ (c,d, cyan spheres) are distal from the L2-loop of RecA (d, red spheres). In the atomic model of the groove mode the negatively charged C-terminal helix forms a continuous helical pattern (e,f,, cyan spheres) flanking the L2 region of RecA (f, red spheres). The position of the single RecA protomer (blue ribbons in b,d,f) is marked with red asterisks in the filament models (a,c,e).



Fig. 6.

The RecX protein (red) consists of three domains, where each domain contains three α -helices⁴⁰. The structural similarity of each of the three domains has suggested a gene triplication. The DinI protein (yellow) is a single domain, containing two α -helices and a single β -sheet³⁶. Two of the three RecX helices superimpose quite well on the two helices in DinI, raising the possibility of homology for this RecA-binding domain. At the bottom DinI helix containing residues 17-31 is superimposed on the RecX helix containing residues 81-90, while at the top the DinI helix containing residues 57-73 is superimposed on the RecX helix containing residues 63-76.