RecO Protein Initiates DNA Recombination and Strand Annealing through Two Alternative DNA Binding Mechanisms*

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**Background:** RecO anneals DNA and initiates homologous recombination.

**Results:** Binding of RecO to DNA/SSB is regulated by zinc during annealing and by RecR during recombination.

**Conclusion:** Alternative DNA repair reactions are supported by different DNA binding mechanisms of RecO.

**Significance:** Results explain how the same RMP supports multiple reactions during DNA repair and chromosome maintenance.

Recombination mediator proteins (RMPs) are important for genome stability in all organisms. Several RMPs support two alternative reactions: initiation of homologous recombination and DNA annealing. We examined mechanisms of RMPs in both reactions with *Mycobacterium smegmatis* RecO (MsRecO) and demonstrated that MsRecO interacts with ssDNA by two distinct mechanisms. Zinc stimulates MsRecO binding to ssDNA during annealing, whereas the recombination function is zinc-independent and is regulated by interaction with MsRecR. Thus, different structural motifs or conformations of MsRecO are responsible for interaction with ssDNA during annealing and recombination. Neither annealing nor recombinase loading depends on MsRecO interaction with the conserved C-terminal tail of single-stranded (ss) DNA-binding protein (SSB), which is known to bind *Escherichia coli* RecO. However, similarly to *E. coli* proteins, MsRecO and MsRecOR do not displace SSB from ssDNA, suggesting that RMPs form a complex with SSB-ssDNA even in the absence of binding to the major protein interaction motif. We propose that alternative conformations of such complexes define the mechanism by which RMPs initiate the repair of stalled replication and support two different functions during recombinational repair of DNA breaks.

Homologous recombination (HR) plays an important role in multiple aspects of chromosome maintenance and genome stability (1–5). The activity of RecA-like recombinases, which support HR, is regulated at several levels (6, 7). HR is initiated by recombinase binding to single-stranded (ss) DNA and formation of a so-called presynaptic complex (8). This step is inhibited by ssDNA binding proteins such as gp32, SSB, and replication protein A (RPA). Recombination mediator proteins (RMPs) overcome such an inhibitory effect and stimulate recombinase binding to ssDNA in response to DNA damage (9). Examples of RMPs include the phage UvsY protein, bacterial RecF, RecO, and RecR proteins (RecFOR) and eukaryotic Rad52, PALB2, and BRCA2 proteins (10–14). Mutations of BRCA2 and PALB2 are associated with cancer predisposition (15–17). RecO and RecR are implicated in drug resistance and host immune response evasion mechanisms in pathogens (18–20).

RecO, RecR, and RecF form an epistatic group involved in DNA repair and replication restart in *Escherichia coli* and other bacteria (19, 21–30). *E. coli* RecR (EcRecR) is required for EcRecO function in presynaptic complex formation in vitro, whereas EcRecF plays a regulatory role (31, 32). EcRecF stimulates efficiency of EcRecA binding to the double-stranded (ds)/ssDNA junction substrate in the presence of excess EcSSB (33, 34). RecO has a secondary recombinase-independent function of ssDNA annealing (SSA), as shown for homologs from several bacteria (35–38). As in the case of HR, SSB inhibits DNA annealing, and RecO overcomes an inhibitory effect of SSB. Neither EcRecR nor EcRecF is required for SSA by EcRecO, and EcRecR inhibits SSA activity of EcRecO (38). Eukaryotic RMPs, Rad52, and the BRCA2 homolog Brh2 also possess a similar SSA activity despite lacking sequence or structural homology with RecO and with each other (35–38). It is unclear why these proteins support two functions. One of the ideas with some experimental support in yeast suggests that SSA supports a second-end capture reaction during the post-strand invasion step of HR (39–43). How different functions are activated at different stages of HR is unknown. The function of SSA in *E. coli* is also not defined (44). Our recent genetic studies of *Mycobacterium smegmatis* RecO (MsRecO) revealed its critical role in HR and SSA pathways of dsDNA break repair (29, 45).
The mechanism by which RMPs facilitate recombinase binding to ssDNA protected by ssDNA-binding proteins is poorly understood. There are two steps in presynaptic filament formation: nucleation and extension. SSB and RPA inhibit the nucleation steps but can be beneficial for the extension (6). Rad52 and BRCA2 simultaneously bind both RPA and Rad51, thus bringing Rad51 to ssDNA proximity through protein–protein interactions (37, 46–50). Interaction with BRCA2 also alters the DNA binding properties of Rad51 (51). In contrast, EcRecOR lacks RecA binding motifs, and purified proteins do not form stable complexes with EcRecA. The interaction can potentially take place between DNA-bound proteins. For example, EcRecOR prevents dissociation of EcRecA from ssDNA (52), and EcRecFR limits the extension of EcRecA filament beyond ssDNA gap (53). However, the mechanism by which RecOR promotes initial binding of RecA to SSB-coated ssDNA is unknown.

EcRecO binds the C-terminal tail of EcSSB (EcSSB-Ct) (54). This interaction is critical for HR (27) and SSA in E. coli, but it is not conserved in other species. The SSB-Ct-binding site is significantly altered in the structure of Deinococcus radiodurans RecO (DrRecO) (55, 56), and MsRecO does not interact with SSB-Ct (29). Both Dr- and MsRecO possess a 4 Cys zinc finger motif (ZF), whereas there is only one cysteine in a structurally similar domain of EcRecO. A micromolar concentration of zinc in solution stimulates DNA binding and SSA of MsRecO (29).

In this study we have characterized the recombination function of MsRecO. Surprisingly, we found that zinc interaction with MsRecO is dispensable for the presynaptic complex formation. DNA binding of the zinc-depleted MsRecO is stimulated by MsRecR. Thus, MsRecO interacts with ssDNA via two alternative mechanisms in SSA and HR. Neither MsRecO nor MsRecOR completely displace SSB from ssDNA even in the absence of interaction with SSB-Ct. The existence of ssDNA/SSB/RMP complexes explains how two alternative reactions of the strand invasion and SSA can be supported by same RMP during consecutive steps of the recombinational repair.

**EXPERIMENTAL PROCEDURES**

**Chemicals**—Glycerol, HEPES, Mg(OAc)2, NaCl, polyethyleneimine, and Trizma base (Tris base) were purchased from Sigma, and ammonium sulfate was from Fisher. Tris(OAc)2 was purchased from Hampton Research Corp (Aliso Viejo, CA). nickel-nitritolactreatric acid resin was purchased from Molecular Cloning Laboratories (South San Francisco, CA), and 5-ml HiTrap columns were from GE Healthcare. Tris (2-carboxyethyl) phosphine hydrochloride (TCEP) was purchased from Gold Biotechnology, Inc. (St. Louis, MO). DNA oligonucleotides were purchased from Integrated DNA Technologies, Inc. (Coralville, IA).

**Protein Purification**—His-tagged E. coli and M. smegmatis RecO and RecR proteins were cloned and purified as described previously (29). Mycobacterial RecO was chelated with 1 mM EDTA, reduced with 1 mM DTT, and dialyzed extensively against storage buffer (40% (v/v) glycerol, 0.5 M NaCl, 20 mM HEPES pH 7.5, 1 mM TCEP). E. coli and M. smegmatis SSB proteins were expressed in E. coli BL21(DE3)pLys without fusion to affinity tags and purified by sequential polyethyleneimine and ammonium sulfate precipitation and heparin chromatography as described previously (29). The E. coli SSB-AC8 expression plasmid was a gift from Dr. M. Cox (University of Wisconsin, WI) and was purified by similar protocol. E. coli RecA expression plasmid was a gift from Michael Cox (University of Wisconsin, WI). EcRecA was inserted into pMC6SG7, expressed in BL21(DE3)pLys, and purified by ammonium sulfate precipitation, nickel-nitritolactreatric acid chromatography with cleavage of the N-terminal His6 tag by tobacco etch virus, and heparin chromatography. M. smegmatis RecA was amplified from genomic DNA and was inserted into pEcoli-Cterm 6xHN (Clontech) with a stop codon preceding the His6 C-terminal tag, expressed in BL21(DE3)pLys, and purified by sequential polyethyleneimine and ammonium sulfate precipitation and anion exchange chromatography.

**DNA Binding**—Proteins were diluted into buffer A (50 mM NaCl, 25% (v/v) glycerol, 50 mM HEPES pH 7.5, 1 mM TCEP) or buffer B (100 mM NaCl, 25% (v/v) glycerol, 50 mM HEPES, pH 7.5, 1 mM TCEP). Binding of DNA was assayed by titrating 5 nM fluorescein 5'-labeled dT15 oligonucleotide by RecO proteins in buffers A and B in the absence or presence of 10 μM Zn(OAc)2 or 8 μM RecR. DNA binding was assayed using fluorescence anisotropy as previously described with the excitation/emission of 485/528 nm at room temperature using a BioTek Synergy 4 plate reader (55).

**Strand Exchange and ATP Hydrolysis**—The entire assay and series of incubations were conducted at 37 °C. The 90-mer oligonucleotide (GCC TCT AGT CGA GTC ATC AAT AGC AAA CCT TAT TCT TCC CAG TTA CAA GCA CTT AAG GTC TTG TTC GCA GAT GCC TTA GAG CTT ATT TGC) at 25 nM (molecule concentration) was incubated with 1 μM SSB for 15 min. 1 μM RecO and 2 or 8 μM RecR were incubated for 15 min. SSB-ssDNA and RecOR(R) were mixed and incubated for 15 min in Corning® 96-well Non-binding (NBSTM) clear bottom plates (product #3651). RecA (1 μM) and ATP (0.2 mM) were added to the SSB-ssDNA-RecOR mixture and incubated for 2 min. The target substrate dsDNA of 35 nucleotides long with Cy3-5’- and Cy5-3’-labeled dT15 series of incubations were conducted at 37 °C. The 90-mer oligonucleotide (GCC TCT AGT CGA GTC ATC AAT AGC AAA CCT TAT TCT TCC CAG TTA CAA GCA CTT AAG GTC TTG TTC GCA GAT GCC TTA GAG CTT ATT TGC) at 25 nM (molecule concentration) was incubated with 1 μM SSB for 15 min. 1 μM RecO and 2 or 8 μM RecR were incubated for 15 min. SSB-ssDNA and RecOR(R) were mixed and incubated for 2 min. The target substrate dsDNA of 35 nucleotides long with Cy3-5’- and Cy5-3’-labeled dT15 was added at 35 or 50 nM. The final volume of reaction was 100 μl. Cy3-5’-strand is complementary to the 3’-end of the 90-mer oligonucleotide. RecA loading onto DNA was assayed by measuring DNA-dependent ATP hydrolysis with a coupled ATP regeneration colorimetric assay consisting of ATP (0.2–0.25 mM), pyruvate kinase (2–2.25 units/ml), phosphoenolpyruvate (0.2–0.25 mM), lactate dehydrogenase (2–2.25 units/ml), and NADH (0.2–0.25 mM). Decrease of NADH absorbance at 340 nm was converted to ATP hydrolysis based on a previously published protocol (33, 52). Strand exchange was measured by FRET changes with excitation at 540/25 nm (with 25-nm bandwidth) and emission at 590/20 nm and 635/35 nm during 30 min. FRET was calculated using the equation, FRET = A/(A + D), where A and D are emissions from acceptor at 635 nm and donor at 590 nm. Strand exchange was normalized to 100%.
corresponding to maximum possible strand exchange product formation. FRET results were verified (not shown) by analyzing the strand exchange products on Native PAGE gels by incubating the samples in 5% glycerol (v/v), 1.2% SDS, 10 mM Tris-HCl, pH 8.0, 40 mM EDTA, 0.25 mg/ml Proteinase K at 42°C for 30 min and visualizing the products and substrates by excitation/emission at 532/580, 532/670, and 633/670 nm with a Typhoon imager (GE Healthcare). 90% of maximum FRET changes were observed during the first 15 min of reaction, and this time point was selected for bar graph plots to compare activities of RMPs at different conditions. The assay was performed in buffer C (5% (v/v) glycerol, 60 mM NaCl, 25 mM Tris(OAc) 2, pH 7.0, 5 mM Mg(OAc) 2, 1 mM TCEP, 3 mM potassium gluconate (KGlu)) or buffer D (5% (v/v) glycerol, 60 mM NaCl, 25 mM Tris(OAc) 2, 10 mM Mg(OAc) 2, 1 mM DTT, 3 mM KGlu, pH 7.0, in the case of M. smegmatis or 7.5 in case of E. coli).

DNA Annealing—MsRecO annealing activity was assayed as described previously (55). 35-mer 5'-Cy3-labeled (/5Cy3/GCA AAT AAG CTC TAA GCC ATC TGC GAA CAC GCT) and complementary 3'-Cy5 labeled (AGG TCT TGT TCG AAT AAG CTC TAA GCC ATC TGC GAA CAA GAC CT) MsRecO annealing activity was assayed as described previously (55). 35-mer 5'-Cy3-labeled (/5Cy3/GCA AAT AAG CTC TAA GCC ATC TGC GAA CAC GCT) and complementary 3'-Cy5 labeled (AGG TCT TGT TCG AAT AAG CTC TAA GCC ATC TGC GAA CAA GAC CT) MsRecO was added. The FRET was measured by excitation at 540/25 nm and emission at 590/20 and 635/35 nm for 25 min at room temperature. Annealing was normalized to 100%, corresponding to a FRET of 0.6 for dsDNA formation. Background interaction between labeled oligonucleotides and proteins was subtracted as previously described by measuring Cy3 and Cy5 emissions of single-labeled oligonucleotides in the absence of proteins (57).

Protein Pulldown with ssDNA Immobilized on Avidin Beads—5'-Biotin-labeled 65-nucleotide-long ssDNA (/5Biosg/GCA AAT AAG CTC TAA GCC ATC TGC GAA CAA GAC CT) and complementary 3'-Cy5 labeled (AGG TCT TGT TCG AAT AAG CTC TAA GCC ATC TGC GAA CAA GAC CT) ssDNA bound to Ms- or EcSSB under physiological conditions was saturated with MsSSB in the presence or absence of 1 μM Zn(OAc) 2. The MsSSB-ssDNA complexes were mixed, and 1 μM MsRecO was added. The FRET was measured by excitation at 540/25 nm and emission at 590/20 and 635/35 nm for 25 min at room temperature. Annealing was normalized to 100%, corresponding to a FRET of 0.6 for dsDNA formation. Background interaction between labeled oligonucleotides and proteins was subtracted as previously described by measuring Cy3 and Cy5 emissions of single-labeled oligonucleotides in the presence of proteins (57).

**RESULTS**

**MsRecO Binding to DNA Is Stimulated Independently by Zinc and RecR**—Previously we demonstrated that binding of MsRecO to DNA is stimulated by the presence of a micromolar amount of zinc in solution (29). The interaction of MsRecO with ssDNA in the presence of zinc was sufficient to anneal ssDNA bound to Ms- or EcSSB under physiological conditions despite a lack of interaction between MsRecO and C-terminal tail of MsSSB. To test the recombination function of MsRecO, we first assessed the interaction of MsRecO with ssDNA in the presence of MsRecR as a required partner of RecO in RecA loading reaction. Unexpectedly, we found that MsRecR stimulated DNA binding of zinc-depleted MsRecO similarly to that of zinc (Fig. 1A). The apparent dissociation constants at 50 mM NaCl were Kd(MsRecO) = 2.08 ± 0.11 μM and Kd(MsRecO:Zn) = 0.041 ± 0.03 μM in the absence or presence of 10 μM Zn(OAc) 2, correspondingly. In the presence of MsRecR, Kd(MsRecOR) = 0.028 ± 0.006 μM without zinc. At 100 mM NaCl, stimulation of DNA binding of MsRecO by MsRecR was stronger than in the case of E. coli proteins (Fig. 1B; Kd(MsRecOR) = 0.44 ± 0.06; Kd(EcRecOR) = 2.76 ± 0.27). Therefore, the interaction of MsRecO with ssDNA is regulated by two alternative mechanisms. In one case the zinc finger domain is critical for DNA binding and SSA by MsRecO alone. In the second case, the coordination of the ZF domain structure by metal is dispensable for ssDNA binding by MsRecO bound to MsRecR. To understand the role of the alternative DNA interaction mechanisms in the recombination function of MsRecOR, we cloned and purified MsSSB and MsRecA proteins and developed an assay to monitor the DNA binding and strand exchange activities of RecA.

**MsRecOR Stimulates DNA Binding and Strand Exchange Activities of MsRecA**—Binding of the purified MsRecA to 90-mer ssDNA and the strand exchange activity with a donor 35-bp dsDNA substrate were simultaneously monitored as described in Fig. 2. ATP hydrolysis (Fig. 2, B and D) and strand exchange products were monitored as described in Fig. 2. ATP hydrolysis (Fig. 2, B and D) and strand exchange products were monitored as described in Fig. 2.
exchange activities of MsRecA alone without MsSSB (Fig. 2, C and E) were significantly weaker than those of EcRecA (Fig. 2, B–E, Table 1), in agreement with previously published data (58). The strand-exchange activity of MsRecA was observed to be particularly weak with the longer M13 DNA substrates (data not shown). SSB inhibited the DNA binding of RecA from both organisms. In both systems RecO alone did not support RecA function. MsRecOR complex stimulated the DNA binding and strand exchange activities of MsRecA in the presence of MsSSB and the strand exchange in the absence of MsSSB (Fig. 2). EcRecOR stimulated EcRecA function only in the presence of EcSSB, in agreement with previous reports (27). Thus, in the Gram-positive bacterium *M. smegmatis*, with a more complex array of DNA repair pathways in comparison to *E. coli*, activity of MsRecA is more dependent on function of RMPs.

**Stimulation of MsRecA Binding to DNA by MsRecOR Does Not Depend on MsRecO Interaction with Zinc**—MsRecO protein was extensively dialyzed against buffer with chelating and reducing agents and assayed for strand annealing and for recombination mediator functions. Under these conditions, MsRecOR supported the interaction of MsRecA with ssDNA and strand exchange, whereas MsRecO did not stimulate annealing (Fig. 3). The addition of zinc to the reaction buffer restored the strand annealing activity of MsRecO but not of MsRecOR. It partially inhibited MsRecA activity, probably due to zinc interference with RecA function. Nevertheless, MsRecOR did stimulate strand exchange reaction.

**MsRecR Inhibits the Annealing Activity of MsRecO**—EcRecR was shown to inhibit annealing activity of EcRecO (38). Likewise, we observed that MsRecR inhibited SSA by MsRecO even

**FIGURE 2. MsRecOR stimulates DNA binding and strand exchange activities of MsRecA.** A, design of the experiment. The 90-mer ssDNA was incubated with SSB followed by incubation with RecO or RecOR. RecA was added followed by the addition of donor dsDNA 35-mer composed of a 3'-Cy5- and a 5'-Cy3-labeled strands complementary to the 3'-end of the 90-mer ssDNA. Final concentrations are 90-mer ssDNA (25 nM), 35-bp dsDNA (35 nM), RecA (1 μM), SSB (1 μM), RecO (1 μM), and RecR (2 μM). DNA-dependent ATP hydrolysis by RecA was measured by a coupled ATP regeneration colorimetric assay with NADH. Decrease of NADH absorbance at 340 nm corresponds to ATP hydrolysis. Strand exchange was measured simultaneously by a decrease of Cy3/Cy5 FRET upon strand exchange with the complementary 3'-end of ssDNA 90-mer. The decreasing FRET was inverted and normalized to reflect the percentage of maximum strand exchange. B and C, real time measurements of ATP hydrolysis (B) and strand exchange (C) reactions are shown for EcRecA (1), MsRecA (2), EcRecA with EcSSB (3), MsRecA with MsSSB and MsRecOR (4), and DNA alone (5). D and E, bar graphs representing ATP hydrolysis (D) and strand-exchange efficiency (E) by *E. coli* (gray) and *M. smegmatis* (black) RecA after a 15-min reaction time in the presence of 90-mer ssDNA and 35-bp dsDNA with cognate SSB, RecO, and RecR in buffer C and ATP regeneration system.

**TABLE 1**

| ATP hydrolysis rate in $k_{cat}$ (min$^{-1}$) of *E. coli* and *M. smegmatis* RecA | *E. coli* | *M. smegmatis* |
|---------------------------------------------|-----------|----------------|
| DNA-dependent ATP hydrolysis rates of RecA proteins in the presence of SSB and RecOR as calculated at 15 min of the reaction time in experiment described in Fig. 2. | | |
| RecA | 15.9 ± 0.4 | 6.8 ± 0.2 |
| RecA, SSB | 1.1 ± 0.2 | 1.6 ± 0.1 |
| RecA, RecO | 0 | 0 |
| RecA, RecOR | 8.2 ± 0.6 | 2.9 ± 0.3 |
| RecA, SSB, RecO | 0 | 0 |
| RecA, SSB, RecOR | 1.6 ± 0.6 | 4.2 ± 0.4 |
in the presence of zinc under conditions favorable for annealing (Fig. 3). Eco- and MsRecOR complexes bind both ss- and dsDNA with comparative affinity to that of RecO alone (55), suggesting that the inhibitory effect is not due to incompatibility of the complex with the end product of annealing reaction. We hypothesize that the conformation of ssDNA bound to RecOR is incompatible with the strand annealing, and, consequently, that RecO and RecOR support two alternative conformations of ssDNA.

MsRecOR Function Independently of Interaction with SSB-Ct—The annealing activity of EcoRecO depends on the interaction with SSB-Ct only at elevated salt concentrations. However, the presence of SSB-Ct was essential for EcoRecOR function in EcoRecA loading experiments even at low salt concentrations (59). We tested whether the mediator function of MsRecOR is dependent on the interaction with SSB-Ct. MsRecOR function was assayed with EcoSSB as the stimulatory effect of MsRecOR on the activity of EcoRecA in the presence of EcoSSB was similar to that of MsRecA with MsSSB. Comparison of reactions in the presence of EcoSSB and the mutant lacking last eight amino acids (EcoSSB-)/H9004 did demonstrated that MsRecOR supported EcoRecA binding to ssDNA and strand exchange under all conditions, whereas EcoRecOR activity was inhibited by EcoSSB-)/H9004 (Fig. 4).

MsRecO and MsRecOR Do Not Displace MsSSB from ssDNA—Interaction of EcoRecO with EcoSSB led to the hypothesis that RecO does not completely dismiss SSB from ssDNA upon binding (22). Accordingly, we demonstrated that EcoSSB retained on dT70 immobilized on avidin beads after incubation with excess of EcoRecO or EcoRecOR (54). Interestingly, similar results were obtained at low salt concentration with SSB-)/H9004 construct, where the interaction with SSB-Ct is dispensable for annealing. Because MsRecO does not bind SSB-Ct, we addressed the same question using mycobacterial proteins. We assayed MsSSB displacement using a similar pulldown assay with 65-mer ssDNA immobilized on beads. MsSSB was incubated with beads, and the unbound protein was removed. Next, the beads were incubated with the excess of MsRecO or MsRecOR in the absence of unbound MsSSB. Similar to the
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**DISCUSSION**

Reco and RecOR Support Two Distinct Conformations of ssDNA during SSA and HR—The data presented in this work demonstrate that different structural elements of MsRecO are involved in ssDNA binding in two alternative DNA repair pathways. During annealing, ssDNA binding depends on the presence of zinc in solution likely due to structural requirements of

E. coli system, no significant dissociation of MsSSB was observed upon incubation with MsRecO, and comparable amounts of both MsRecO and MsSSB were retained on beads (Fig. 5A). In the case of MsRecOR, a partial dissociation of MsSSB from DNA was observed. However, the amount of MsSSB retained on the beads was also comparable with that of MsRecOR (Fig. 5B). Moreover, the addition of MsRecA resulted in the retention of all four proteins on ssDNA. The results together with previous analysis of E. coli proteins suggest that RecO(R) can form a complex with ssDNA and SSB regardless of interaction with SSB-Ct.

MsSSB was previously reported to interact with MsRecA and stimulate its function (58). In our experiments, MsSSB inhibited ATPase and strand-exchange activities of MsRecA. However, in the presence of MsRecO the titration of the reaction by MsSSB revealed a partial stimulatory effect of MsSSB (Fig. 6, A and B). Such an increase was not observed with E. coli proteins, in line with the previously reported results (Fig. 6, C and D) (59). These data further support a functional interaction between RecO(R) and SSB on ssDNA.

FIGURE 5. MsRecOR and MsRecA share ssDNA with MsSSB. A, SDS-PAGE analysis showing proteins pulled down on avidin-immobilized ssDNA. Lane 1, initial mixture of MsSSB (10 μM) with avidin-bound 65-mer ssDNA (load); lane 2, unbound MsSSB in supernatant (SN) after incubation with ssDNA beads; lane 3, proteins in solution after the third round of beads washing (W); lane 4, MsSSB retained on beads after the third wash (Bds); lane 5, ssDNA beads pre-bound to MsSSB mixed with MsRecO (10 μM); lane 6, unbound protein in initial mixture; lane 7, unbound protein after the third wash; lane 8, proteins remaining on beads after three washes. B, similarly prepared ssDNA immobilized on beads and bound to MsSSB were incubated with MsRecO (10 μM) and MsRecR (40 μM) (lanes 1–4) and, consequently, with MsRecA (5 μM) and ATPγS (1 mM) (lanes 5–8) in buffer C. Lanes 9 and 10 contain negative controls for beads without DNA incubated with MsSSB (SSB) or MsRecOR (OR), respectively, after extensive wash.

FIGURE 6. MsSSB partially promotes MsRecA activity in the presence of MsRecOR. A and B, shown are bar graphs representing values of ATP hydrolysis (A) and strand-exchange efficiency (B) by MsRecA (1 μM) at 15 min with ssDNA preincubated with different amounts of MsSSB in buffer C in the absence (gray) or presence (black) of MsRecO (1 μM) and MsRecR (2 μM). C and D, similar bar graphs showing ATP hydrolysis (C) and strand-exchange efficiency (D) by EcRecA (1 μM) upon titration with EcSSB in buffer C in the absence (gray) and presence (black) of EcRecO (1 μM) and EcRecR (2 μM).

the ZF domain. The depletion of zinc by chelating agents results in an annealing-inactive protein with significantly lower affinity to ssDNA. However, DNA binding is restored in the presence of MsRecR, which by itself has a very weak affinity to DNA in the millimolar range of concentration.

RecR has a highly conserved amino acid sequence (60, 61). Despite its essential role in RecA binding, RecR does not form stable complexes with RecA. A weak DNA-dependent interaction between RecR and RecA is still a possibility. Alternatively, we hypothesize that the specific conformation of ssDNA bound to RecOR is favorable for interaction with RecA. Interestingly, the excess of RecR in solution does not inhibit RecA function. An inhibitory property of RecR in DNA annealing further supports the importance of specific ssDNA conformation. For example, the RecOR complex interacts with comparable affinities with both ds- and ssDNA. Thus, the inhibition can be explained by a conformation of ssDNA incompatible with annealing. Therefore, we suggest that RecO can support two alternative conformations of ssDNA; one favorable for annealing and another, in the presence of RecR, favorable for RecA binding to ssDNA. Likewise, DNA conformational changes supported by gp32/UvsY proteins were suggested to play a major functional role in homologous recombination (62).

*Mycobacterial RMPs Co-localizes with SSB on ssDNA—At physiological conditions, interaction with SSB-Ct is critical for function of EcRecO and EcRecR, which do not displace EcSSB from ssDNA during the initial steps of SSA and HR. MsRecO does not bind MsSSB-Ct. However, we observed a co-localization of MsRecO and MsRecOR with MsSSB on ssDNA. A partial displacement of MsSSB cannot be ruled out by this experiment. Nevertheless, the amount of the protein retained on beads-immobilized ssDNA is comparable with that of MsRecO or MsRecOR. Moreover, MsSSB slightly facilitates MsRecOR-mediated strand-exchange activity of MsRecA in a limited con-
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![Diagram of RecO, RecR, and RecA interactions](image)

**FIGURE 7. Hypothetical model of strand exchange and annealing reactions mediated by RecO(R).** 1. RecO binds SSB-ssDNA and promotes conformational changes of ssDNA favorable for recruitment of RecA to ssDNA at the site of stalled replication or (2A) to the recessed ssDNA end at dsDNA break (DSB). Alternately, (2B) RecO forms intermediate complexes with SSB-ssDNA and facilitates SSA when complementary ssDNA is available during a second-end capture reaction or (3) during annealing of repetitive ends. Reversibility of each intermediate step eliminates the inhibitory effect of RMPs on other functions of SSB-ssDNA complexes and permits selective activation of each pathway only under kinetically favorable conditions.

The mechanism of DNA sharing by SSB and RMPs remains to be investigated. 65-nucleotide-long ssDNA is wrapped around SSB tetramer in a so-called (SSB)$_{65}$ mode (68–71) with some nucleotides forming extensive contacts with the protein, whereas others are more solvent-exposed. RecO and RecOR can interact initially with such less tightly bound regions. Alternatively, RecO(R) can alter the DNA binding mode of SSB from (SSB)$_{65}$ to (SSB)$_{135}$ where half of ssDNA wrapped around SSB tetramer can become available for RecO(R) binding (72). In either case, RecO and RecOR alter the conformation of ssDNA bound to SSB to promote annealing or RecA recruitment.

The Role of SSB-RMP-ssDNA Complexes in DNA Repair—RecA binding to ssDNA occurs at an early stage of DNA repair as RecA-ssDNA complexes initiate such events as the SOS response in bacteria. Thus, RMPs, which stimulate RecA binding to ssDNA, can function as initial sensors of DNA damage or stalled replication. The existence of SSB-RMP-ssDNA complexes observed in our studies, in studies of UvsY/gp32 (73), and of Rad52-RPA (47, 74) can explain how RMP initiates recombinase loading in response to DNA damage. We speculate that SSA and HR reactions proceed through several reversible steps of RecO(R) interaction with SSB-ssDNA as depicted in Fig. 7. Such a mechanism will have two important features. First, a sequential formation of reversible complexes can lead to relatively slow kinetics, e.g. compared to that of replication. Second, the completion of each reaction depends on the availability of an appropriate substrate, e.g. complementary ssDNA in case of SSA. Otherwise, the complexes can dissociate to free SSB for interaction with other proteins. These features can explain how such RMP-SSB complexes will load RecA on ssDNA upon replication stalling but will not interfere during replication (34, 75–77). Accordingly, the lethality of a *LivrD/*Rep E. coli mutant due to slow replication and lack of anti-recombinase activity is bypassed by deletion of either of the RecFOR proteins (78, 79).

In the case of dsDNA break repair, the proposed model will explain why RMPs initially support only presynaptic complex formation on a resected DNA end but not SSA due to the lack of the complementary ssDNA substrate. In the post-strand invasion step, upon generation of complementary ssDNA, RMPs can support the second-end capture reaction as was previously hypothesized (66, 80). The existence of reversible Rad51-DNA complexes and Rad52-RPA complexes was previously proposed to control recombination initiation in eukaryotes (81).

Regulation of RecO Function in Mycobacteria—MsRecO retains weak SSA activity for a limited time immediately after purification under reducing conditions without zinc in solvent (data not shown). Estimation of zinc binding affinity by MsRecO revealed an apparent dissociation constant of 0.1 μM (29). Such moderate to low affinity, as compared with other zinc finger domains (82), can serve as a regulatory mechanism of SSA activity of MsRecO in mycobacteria. Zinc is an important cofactor in all organisms, and mycobacteria possess several mechanisms that regulate the intracellular zinc concentration (83, 84). An overall concentration of zinc in bacteria is estimated at 0.2 mM, although the amount of unbound zinc is in the femtomolar concentration range (85). Therefore, zinc binding by MsRecO depends on competition for zinc with other zinc binding molecules in the cell. The ZF domain of MsRecO can also be sensitive to oxidation as it contains four cysteines. Therefore, the annealing function of mycobacterial RecO can be inhibited upon zinc depletion or oxidative conditions, whereas a recombination mediator function should be supported under such conditions.

The structural role of the ZF domain in RecO remains to be investigated. Despite low homology, structures of Dr- and EcRecO are similar including the folding of ZF domain (54, 56). MsRecO revealed an apparent dissociation constant of 0.1 μM (29). Such moderate to low affinity, as compared with other zinc finger domains (82), can serve as a regulatory mechanism of SSA activity of MsRecO in mycobacteria. Zinc is an important cofactor in all organisms, and mycobacteria possess several mechanisms that regulate the intracellular zinc concentration (83, 84). An overall concentration of zinc in bacteria is estimated at 0.2 mM, although the amount of unbound zinc is in the femtomolar concentration range (85). Therefore, zinc binding by MsRecO depends on competition for zinc with other zinc binding molecules in the cell. The ZF domain of MsRecO can also be sensitive to oxidation as it contains four cysteines. Therefore, the annealing function of mycobacterial RecO can be inhibited upon zinc depletion or oxidative conditions, whereas a recombination mediator function should be supported under such conditions.
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